

William Llewellyn's, ANABOLICS, E-Book Edition

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WILLIAM LLEWELLYN'S

ANABOLICS

E-Book Edition

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Avodart® (dutasteride) Proscar® (finasteride)

TESTOSTERONE STIMULATING DRUGS HCG (human chorionic gonadotropin)

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GLOSSARY

Part I

Anabolic Overview

An Introduction to Testosterone

Anabolic steroids are a class of medications that contain a synthetically manufactured form of the hormone testosterone, or a related compound that is derived from (or similar in structure and action to) this hormone. In order to fully grasp how anabolic steroids work, it is, therefore, important to understand the basic functioning of testosterone.

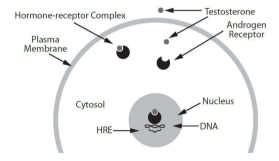
Testostence is the primary make sex hormone. It is manufactured by the Leydrig's cells in the tests at averying amounts throughout a person's life span. The effects of this hormone become most evident during the time of puberty, when an increased output of testosterore will elicit dramatic physiological changes in the male body. This includes the onset of secondary male chanderstimis used as a deependen dvice, hody and ficial hair growth, increased of output by the sobuccous glunds, development of secural organs, system will not finction properly fit insteament by the sobuccous glunds, and the sobuccous glunds, development of security and such as the fittee of the sobuccous glunds, development of security of the sobuccous system will not fittee properly fit insteament between an originalizant. All such effects are considered the masculinizing or "androgenic" properties of this hormone.

Increased testosterone production will also cause growth promoting or "ambdoic" changes in the body, including an enhanced rate of protein synthesis (kading to muscle accumulation). Testosterone is the reason males carry more mascle mass that wormer, as the two sees have vasily contrasting amounts of their hormone. More specifically, the adult male body will manufacture between 2.5 and 11 mpc rady "while females only produce about 1/4mg. The dominant sex hormone for women is earogen, which has a significanity different effect on the body. Among other things, a liver accumulate less muscle insue, have a aborter stature, and become more apt to hone vaclening with age (stateprotoxis).

The actual mechanism in which testosterone elicits these changes is somewhat complex. When free in the blood stream, the testosterone molecule is available to interact with various cells in the body. This includes skeletal muscle cells, as well as skin, scalp, kidney, bone, central nervous system, and prostate tissues. Testosterone binds with a cellular target in order to exert its activity, and will, therefore, effect only those body cells that posses the proper hormone receptor site (specifically the androgen receptor). This process can be likened to a lock and key system, with each receptor (lock) only being activated by a particular type of hormone (key). During this interaction, the testosterone molecule will become bound to the intracellular receptor site (located in the cytosol, not on the membrane surface), forming a new "receptor complex." This complex (hormone + receptor site) will then migrate to the cell's nucleus, where it will attach to a specific section of the cell's DNA, referred to as the hormone response element. This will activate the transcription of specific genes, which in the case of a skeletal muscle cell will ultimately cause (among other things) an increase in the synthesis of the two primary contractile proteins, actin and myosin (muscular growth), Carbohydrate storage in muscle tissue may be increased due to androgen action as well

Once this messaging process is completed, the complex will be released, and the receptor and hornnee will disasciature. Both are then free to migrate back into the cytosol for further activity. The testosterone molecule is also free to diffuse back into circulation to interact with other cells. The entire receptor cycle, including hornnees binding, receptor-hornnee complex, migration, gene transcription and subsequent return to cytosol is a slow process, taking buves, not minutes, to complex. For example, is at discussion. a single nijection of nandrolone, it is measured to be 4 to 6 hours before free androgen receptors migrate back to the cytosol after activation. It is also suggested that this cycle includes the splitting and formation of new androgen receptors once returned to cytosol, a possible explanation for the many observations that androgens are integral in the formation of their own recentor sites.²

In the kidneys, this same processs works to allow androgens to augment erythropoises (red blood cell production).³ It is this effect that leads to an increase in red blood cell productions, and possibly increased oxygen transport capacity, during anabolis/minorgenis stored therapy. Many abletess mistakeny assume that oxymetholene and bolenone are unique in this ability, due to specific uses or mentions of this effect in drag literature. In fact, stranulation of erythropoissis occurs with nearly all anaboli/androgenic strends, as this effect is simply tid with activation of the androgen receptor in kidney cells. The only real exceptions might be compounds such as dihydrotestosterone and some of its derivatives.⁴ which are rapidly broken, seen "anabolic_indrogenic discostintion" section), and therefore display low activity in these tissues.



CELLULAR DIAGRAM: Testosterone freely diffuses through the plasma membrane and binds with an intracellular androgen receptor. The hormone-receptor complex then enters the cell nucleus to bind with a specific segment of DNA (the Hormone Response Element), activating the transcription of specific genes.

Adipose (fat) tissues are also androgen responsive, and here these hormones

support the lipolytic (fit mobilizing) capacity of cells.³ This may be accomplished by an androgen-tied regulation of beta-adrenergic receptor concentrations or general cellular activity (through adenylate cycless).⁶ We also note that the level of androgens in the body will closely correlate (inversely) with level of stored body fit. As the level of androgens bornomes drops, typically the deposition of body fat will increase.⁷ Likewise as we enhance the androgen level, body fat may be depleted at a more active rate. The ratio of androgen to estrogen actions is in fact most important, as settogen plays a counter role by activity to increase the storage of body fat in

many sites of action.⁸ Likewise, if once wished to base fat during steroid use, estrogen levels should be kept low. This is clearly evidenced by the fact that non-aromatizing actions have a always been favored by bodybuilders holding to increase the look of definition and muscularity while aromatizing compounds are typically relegated to builting phases of training due to their tendency to increase body fat storage. Aromatization is discussed in more deall in a following section (see: <u>Editoreal Aromatizion</u>).

As mentioned, testosterone also elicits androgenic activity, which occurs by its activating recentors in what are considered to be androgen responsive tissues (often through prior conversion to dihydrotestosterone. See: DHT Conversion) This includes the sebaceous glands, which are responsible for the secretion of oils in the skin. As the androgen level rises, so does the release of oils. As oil output increases, so does the chance for nores becoming clogged (we can see why acre is such a common side effect of steroid use). The production of body and facial hair is also linked to androgen recentor activation in skin and scaln tissues. This becomes most noticeable as boys mature into nuberty, a period when testosterone levels rise ranidly, and androgen activity begins to stimulate the growth of hair on the body and face. Some time later in life and with the contribution of a genetic predisposition androgen activity in the scalp may also help to initiate male-pattern hair loss. It is a misconcention that dihydrotestosterone is an isolated culprit in the promotion of hair loss, however, as in actuality it is the general activation of the androgen receptor that is to blame (see: DHT Conversion). The functioning of sex glands and libido are also tied to the activity of androgens. as are numerous other regions of the central nervous/neuromuscular system. 1. Role of androgens in growth and development of the fetus, child, and adolescent, Rosenfield R L. Adv Pediatr. 19 (1972) 172-213

² Metabolism of Anabolic Androgenic Steroids, Victor A. Rogozkin, CRC Press 1991

3 Androgens and Erythropoeisis, J Clin Pharmacol, Feb-Mar 1974 p94-101

⁴ Effects of various modes of androgen substitution therapy on erythropoiesis. Jockenhovel F, Vogel E, Reinhardt W, Reinwein D. Eur J Med Res 1997 Jul 282(77):293–8

5. Testosterone increases lipolysis and the number of beta-adrenoceptors in male rat adipocytes. Xu XF, De Pergola G, Bjorntorp P. Endocrinology 1991 Jan;128(1):379-82

⁶ The effects of androgens on the regulation of lipolysis in adipose precursor cells. Endocrinol 126 (1990) 1229-34

⁷ Viscenal fat accumulation in men is positively associated with insulin, glucose, and C-peptide levels, but negatively with testostereone levels. Seidell JC, Bjorntorp L, Sjostrom L, et al.Metabolism 39 (1990) 897-901

⁸ Effects of testosterone and estrogens on deboil and trochanter adipocytes in two cases of transsounism Vague J, Meignen J.M. and Negrin J.F. Horm. Metabol. Res. 16 (1984) 380-381

Direct and Indirect Anabolic Effects

experimented with for many decades now, there is still some debate today as to exactly how steroids affect muscles mass. At this point in time, the primary mode of anabolic action with all anabolic/androgenic steroids is understood to be direct activation of the cellular androgen receptor and increases in protein synthesis. As follows, if we are able to increase our androgen level from an acternal source by supplementing testorence or a similar anabolic steroid, we can greatly enhance the rate in which protein is retained by the muscles. This is clearly the primary causes for muscle growth with all anabolic/androgenic steroids. As our hormone levels increase, so does androgen receptor activation, and ultimately the rate of protein synthesis.

But other indirect mechanisms could possibly affect muscle growth outside of the normally understood androgan action on protein synthesis. An indirect mechanism is one that is not brought about by activation of the androgen receptor, but the affect androgens might have on other hormones, or even the release of locally acting hormonics or growth promoters inside cells (parhaps mediated by other membrane board receptors). We must remember also that muscle muss disposition involves not only protein synthesis, but also meed to lock at antiogensis interview with these factors as well to gat a complete picture. Concerning the first possibility, we note that studies with testorteron suggest that this hormone does not increase tissue annino acid.

transport.⁹ This fact probably explains the profound synergy bodybuilders have noted in recent years with insulin, a hormone that strongly increases transport of mutrients into muscle cells. But regranding protein breakdown, we do see a second important pathway in which androgens might affect muscle growth.

Anti-Glucocorticoid Effect of Testosterone

Testosterone (and synthetic anabolic/androgenic steroids) may help to increase mass and strength by having an anti-atable feet on muscle cells. Considered one of the most important indirect mechanisms of androgen action, these hornones are shown to affect the actions of andher type of steroid hornone in the body, glucocorticoids (cortisol is the primary representative of histograph)¹⁶ Glucocorticoids (cortisol is the primary representative of histograph)¹⁶ Glucocorticoids hornones actually have the cact opposite effect on the muscle cell than androgens, namely sending an order to relase stored protent. This process is referred to as catabolism, and represents a breaking down of muscle kissue. Muscle growth is achieved the degenerative effects of costists. Uwith interne training and a proper dire, the body will typically store more protein than it removes, but this underlying batte is advays constant.

When administering nabelies steroids, however, a much higher androgen level can place glucocorticiods at a nobble disadvantage. With their effect reduced, fewer cells will be given a message to release protein, and more will be accumulated in the long mr. The primarily mechanism believed to bring this effect out is androgen displacement of glucocorticoids bound to the glucocorticoid receptor. In fact, nivirus tudies have supported this notion by demonstrating that testosterone has a very high affinity for this receptor, ¹¹ and further suggesting that some of its anabolic activity is directly mediated through this action. ¹² It is also suggested that androgens may indirectly interfere with DNA binding to the glucocorticoid response element. ¹³ Albough the exact underlying mechanism is still in debute, what is clear is that steroid administration inhibits protein breakdown, even in the fasted state, which seems clearly indicative of effect.

Testosterone and Creatine

In addition to protein synthesis, a rise in androgen levels should also enhance the synthesis of creatine in skeletal muscle tissues.¹⁴ Creatine, as creatine nhosphate (CP) plays a crucial role in the manufacture of ATP (adenosine triphosphate) which is a main store of energy for the muscles. As the muscle cells are stimulated to contract. ATP molecules are broken down into ADP (adenosine diphosphate), which releases energy. The cells will then undergo a process using creatine phosphate to rapidly restore ADP to its original structure in order to replenish ATP concentrations. During neriods of intense activity however this process will not be fast enough to compensate and ATP levels will become lowered. This will cause the muscles to become fatigued and less able to effort a strenuous contraction. With increased levels of CP available to the cells. ATP is replenished at an enhanced rate and the muscle is both stronger and more enduring. This effect will account for some nortion of the early strength increases seen during steroid therapy. Although nerhans not technically considered an anabolic effect as tissue hypertrophy is not a direct result androgen support of creatine synthesis is certainly still looked at as a positive and growth-supporting result in the mind of the bodybuilder

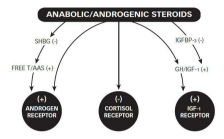
Testosterone and IGF-1

It has also been suggested that there is an indirect mechanism of testosterone action on muscle mass mediated by Insulin-Like Growth Factor. To be more

specific, studies note a clear link between androgens and itsue releases of 15 and responsiveness (n, his anable) hormone. For example, it has been demonstrated that increases in IGF-1 receptor concentrations in skeletal muscle are noted when eldely more angeiven replacement doeses of testosterone. 16 In essence, the cells are becoming primed for the actions of IGF-1, by testostrone. Alternately we see marked decreases in IGF-1 receptor protein levels with androgen deficiency in young men. It also appears that androgens are necessary for the local production and function of IGF-1 in sketcal muscle cells, independent of evaluating growth hormone, and IGF-1 levels. 16 Since we do know for certain that IGF-1 is at least a minor anablek hormone in muscle tissue, is seems reasonable to conclude with steroid theraw. Some Verlag, involved in the muscle growth norder

Direct and Indirect Steroids?

In looking over the proposed indirect effects of tetosterone, and pondering the effectiveness of the synthetic analytokicandogenis stretchis, we must resist the temptation to believe we can categorize stretoids as those which directly, and those which indirectly, promote muscle growth. The belief that there are two dichotomous groups or classes of steroids ignores the fact that all commercial stretoids promote not only muscle growth. The belief that there are two dichotomous groups or classes of steroids ignores the fact that all commercial stretoids promote not only muscle growth has the an and questiclear that all activate the cellular androgen receptor. I believe the theory behind direct and indirect stored descritations originated when some noted the low receptor binding affinity of seemingly strong anabole steroids like asymchohem and mehandrostronome.³ If they had poorly, yet work well, something else must be at work. This type of thaking fails to recognize other factors in the potency of these compounds, such as their long hadfirves, estrogenic activity, and weak interaction with restrictive binding proteins (see: <u>Free vs. Shound Tectoreme</u>). Which there may possibly defifterences in the other vs. Bound Tectorement. Which there may possibly defifterences in the way varous compounds count toter grown marcenty, such mat auvantages might even be found with certain synchright drug combustions, the primary mode of action with all of these compounds is the androgen receptor. The notion that steroid X and Y must never be stacked together because they both compete for the same receptor when stimulating growth, while X and Z should be combined because they work via different mechanisms, should likewise not be taken too seriously. Such classifications are based on speculation only, and upon reasonable investigation are query invalid.



MECHANISM OF ACTION DIAGRAM: The mechanism of anabolic action due to the administration of anabolic/androgenic steroids. AAS causes not only direct stimulation of the androgen receptor, but also supports muscles of free androgens, increasing androgen receptor density, inhibiting conterior al candrogens, increasing androgen-receptor density, inhibiting conterior al candrogens, mercanism (Erl/FeI-1, and supportssing IGF-1 binding protection).

⁹. Testosterone injection stimulates net protein synthesis but not tissue anino acid transport, Fernando A, Tipton K, Doyle D et al. Am J. Physiol (Endocrinology and Metabolism) 38:E864-71,1998.

¹⁰ Ghaorticoid antagonism by evercise and androgenic-anabolic steroids. Hickson RC, Czerwinski SM, Fakluto MT, Young AP. Med Sci Sports Exerc 22 (1990) 331-40

¹¹. Binding of glucorticoid antagonists to androgen and glucorticoid hormone receptors in rat skeletal muscle. Danhaire PA, Rousseau GG. J Steroid Biochem Mol Biol 24 (1986) 481-71

¹² Evidence for a sex-dependent anabolic response to androgenic steroids mediated by muscle glucorticoid receptors in the rat. Danhaive PA, Rousseau GG. J. Steroid Biochem Mol Biol. 29 (1988) 575-81

¹³ Glucorticoid antagonism by exercise and androgenic-anabolic steroids. Hickson RC, Czerwinski SM, Falduto MT, Young AP. Med Sci Sports Exerc 22 (1990) 331-40

¹⁴ The source of excess creatine following methyl testosterone. Samuels L. T., Sellers D. M., McCaulay C. J. J. Clin. Endocrinol. Metab. 6 (1946) 655-63

¹⁵. Ontogeny of growth hormone, insulin-like growth factor, estradiol and cortisol in the growing lamb: effect of testosterone. Amold AM, Peralta JM, Tonney ML J Endocrinol 150 (1996) 331-9 12 Jun;130(6):3677-38. 31, 2001. ^{16.} Testosterone administration to elderly men increases skeletal muscle strength and protein synthesis. Am J Physiol 269 (1995) E820-6

¹⁷. Testosterone deficiency in young merr marked alterations in whole body protein kinetics, strength, and adiposity. Mausas N, Hayes V, Welch S et al. J Clin Endocrin Metab 83 (1998) 1886-92

¹⁸ Endocrinology 114(6):2100-06 1984 June, "Relative Binding Affinity of Anabolic-Androgenic Steroids...", Saartok T; Dahlberg E; Gustafsson JA

Free vs. Bound Testosterone

A very small amount of testosterone actually exists in a free state, where interaction with cellular recentors is possible. The majority will be bound to the proteins SHBG (sex hormone binding globulin, also referred to as sex steroid binding globulin and testosterone-estradiol binding globulin) and albumin which temporarily prevent the hormone from exerting activity Steroid hormones actually bind much more avidly to SHBG than albumin (with approximately 1 000 times greater affinity) however albumin is present in a level 1 000 times greater than SHBG. Therefore, the activity of both binding proteins in the body is relatively equal. The distribution of testosterone in men is typically 45% of testosterone bound to SHBG and about 53% bound to albumin. The remaining 2% of the average blood concentration exists in a free unbound state. In women, the percentage of free testosterone is lower measured to be approximately 1% A hinding protein called ABP (androgen hinding protein) also helps to mediate androgen activity in the reproductive system although since it is found exclusively in these tissues, it is not relevant to muscle growth

The level of free testosterore available in the blood is likewise an important factor mediating its activity, as only a small percentage is really active at any given time. It must also be noted that as we also testosterone to form new anabolic/androgenic storisky, we also pryclaidly alter the affinity in which the steroid will bind to plasma proteins. This is an important consideration, as the higher percentage we have of free hormone, the more active the compound should be on a milligram the first more and the variance can be substantial between different compounds.

For example, Provinos® (1-methyl dihydrotestosterone) binds with SHBG many times more avidly than testosterone,¹⁹ while mibolerone (7,17 dmethyl-nandrolone) and bolasterone (7,17 dmethyl- testosterone) show virtually no affinity for this protein at all (clearly the reason these steroids are such potent androgens).

The level of SHBG present in the body is also variable, and can be altered by a number of factors. The most prominent scerns to be the concentration of estrogen and thyroid hormones present in the blood. We generally see a reduction in the amount of this plasma binding protein as estrogen and thyroid content decreases, and a rise in SHBG as they increase. A heightened androgen level due to the administration of anabolic/androgenik stervids has also been shown to kower levels of this protein considerably. This is clearly supported by a 1989 German study, which noted a strong tendency for MHer only 3 days of administering a daily doso cl 2 rangek tody were analy 50% in normal subjects. Smiller results have been obtained with the use of nigretable testostrone enanthate; however, milligram for milligram, the effect of stanozobid was much preader lower and the predist control for the stanozobid was much predist on. The form of administration much predistors. The form of administration much predistors and the predistor data predistors.

have been important in reaching this level of response. Although the injectable was not tried in the Gramma study, we can refer to others omparing the effect of oral vs. transdermal estrogen.²¹ These show a much gratter response in SHBG levels when the drug is given orally. This is perhaps explained by the fact that SHBG is produced in the liver. Therefore, we cannot assume that injectable Winstrol® (or nijectable steroids in general) will displuy the same level of ponency in this regard.

Lowering the level of plasma binding proteins is also not the only mechanism that allows for an increased level of rise testosterone. Servois that display a bigh affinity for these proteins may also increase the level of free testosterone for the servois service of the service service of the additional compound, more will be thit in an unbound state. A number of strotick including dihydrotestosterone, Provison⁸, and Oral-Turniabol (chlonodehydromethyltestosterone) display a strong tendency for this effect. If the level of free-testosterone can be altered by the use of different anabolic-indrogenic strotisk, the possibility also exists that one strotid can increase the potency of another through these same mechanisms. For example, Provison⁸ is a poor anabolic, but is externed by ligh affinity for SHBG might make it useful by allowing the displacement of other strotisk that are more active in these tissues.

We must not let this discussion lead us into thinking that binding proteins serve no valuable function. In fact they play a vial role in the transport and functioning of endogenous androgens. Binding proteins act to protect the steroid against rapid metabolism, ensure a more stable blood hormone concentration, and facilitate an even distribution of hormone to various body organs. The recent discovery of a specific receptor for Set Hormone Binding Clabulin (SHBG-R) based on the membrane surface of steroid responsive solely hormone transport. However, it remains clare that manipulating the tendency of a hormone to exist in an unbound state is an effective way to alter drug potency.

 Endocrinology 114(6):2100-06 1984 June, "Relative Binding Affinity of Anabolic-Androgenic Steroids...", Saartok T; Dahlberg E; Gustafsson JA

²⁰ Sex Hormone-Binding Globulin Response to the Anabolic Steroid Stanozolol: Evidence for Its Suitability as a Biological Androgen Sensitivity Test. J Clin Endocrinol Metab 68:1195,1989

21. Twenty two weeks of transfermal estradiol increases sex hormone-binding globulin in surgical menopausal women. Eur J Obstet Gynecol Reprod Biol 73: 149-52,1997

Estrogen Aromatization

Testosterone is the primary substrate used in the male body for the synthesis of estrogen (estration), the principal female sex hormone. Although the presence of estrogen may scene quiet unusual in men, it is structurally very similar to testosterone. With a sight alteration by the enzyme aronatase, estrogen is produced in the male body. Atomatuse activity occurs in various regions of the male body, including adipose, ²² liver, ²³ gonadal, ²⁴ central nervous system; ²⁵ and skeletal muscle²⁶ tissues. In the context of the average healthy male, the amount of estrogen produced is generally not very significant to one's body disposition. If an efficient of cholesterol values (See Side Effects: <u>Cardiovascular System</u>). However, in larger anomatis in does have potential to cause many unwanted (Ffects:

including water retention, female breast tissue development (gyneconastiu), and body fna taccomulation. For these reasons, many focus on minimizing the build-up or activity of estrogen in the body with aromatase inhibitors such as Arimidex and Cytadren, or anti-strogens such as a Comid or Novbay, particularly at times when gynecomastia is a worry or the athlete is attempting to increase muscle definitions.

We must, however, not be led into thinking that strongen serves no benefit. It is statulity a desirable hormone in many regards. Althetes have known for years that estrogenies steroids are the best mass builders, but it is only recently that we are finally coming to understand the underlying nechanisms why. It appears that reasons go beyond the simple size, weight, and strength increases that one would attribute to estrogen-related water retention, while is municist through increases is ghoese utilization, growth hormose scretcion and andopen recentor northformion.

Glucose Utilization and Estrogen

Entogen may play a very important role in the promotion of an anabolic state by affecting glocose utilization in muscle tissue. This occurs via an altering of the level of available glucose 6-phosphate dehydrogenase, an enzyme directly tied to the use of glucose for muscle tissue growth and trencymenton.²⁷ 25 More specifically, GGPD is a viail part of the pentose phosphate pathway, which is integral in determining the run nuclei acids and plasks are to be synthesized in cells for tissue repair. During the period of regeneration after which is believed to prepare at methods acids and plasks are to be which is believed to prepare at methods and the other encourwhen needed. Samprisngly, we find that estrogen is directly tied to the level of GGPD that is to be made available to cells in this recovery window.

The link between estrogen and G6PD was established in a study demonstrating levels of this dehydrogenase enzyme to rise after administration of testosterone propionate. The investigation further showed that the aromatization of testosterone to estradiol was directly responsible for this increase, and not the androgenic action of this steroid 29 The nonaromatizable steroids dihydrotestosterone and fluoxymesterone were tested alongside testosterone propionate, but failed to duplicate the effect of testosterone. Furthermore, the positive effect of testosterone propionate was blocked when the aromatase inhibitor 4-hydroxyandrostenedione (formestane) was added, while 17-beta estradiol administration alone caused a similar increase in G6PD to tesosterone propionate. The inactive estrogen isomer alpha estradiol, which is unable to bind the estrogen receptor, failed to do anything. Further tests using testosterone propionate and the antiandrogen flutamide showed that this drug also did nothing to block the positive action of testosterone, establishing it as an effect independent of the androgen receptor.

Estrogen and GH/IGF-1

Estogan may also play an important role in the production of growth hormone and IG-11. IG-1 (instituti-like growth factory) is an anabole hormone released in the liver and various perpheral tissues via the stimulus of growth hormone (See Drug Profiles: <u>Circuth Hormone</u>). IG-11 is responsible for the anabolic activity of growth hormone such as increased nitrogen returnitorytoric synthesis and cell hyperplass in profileration.) One of the first studies to bring this issue to our attention looked at the effects of the amis-strogen timoriden is IG-11 levels, demonstrating it to have a suppressive effect.³⁰ A second, perhaps more noteworthy, study took place in 1993, which looked at the effects of testosterone replacement therapy on GH and IGF-1 levels alone, and compared them to the effects of testosterone

combined again with tamoxifen, ³¹ When tamoxifen was given, GH and IGF-1 levels were nothely suppressed, while both values were elvated with the administration of testosterone enambate alone. Another study has shown 300 mg of testosterone enambate weekly to cause a sight IGF-1 increase in normal men. Here the 300 mg of testosterone ester caused an elvation of esteroid field to produce the same dasage of nandrokne decanote; however, this steroid field to produce the same increase. This result, by quite inst²¹²⁰ when scretching hyperbolic the same dasage of nandrokne decanote; however, this scretching hyperbolic the same dasage of nandrokne decanote; however, this scretching hyperbolic the same dasage were actually hyperbolic scretching hyperbolic the same dambatering on males with delety paberty, while dihydrotestostrone (non-aromatizable) scens to suppress GH and IGE-1 acressing ³³

Estrogen and the Androgen Receptor

It has also been demonstrated that estrogen can increase the concentration of androgen receptors in certain tissues. This was shown in studies with rats. which looked at the effects of estrogen on cellular androgen receptors in animals that underwent orchiectomy (removal of testes, often done to diminish endogenous androgen production). According to the study, administration of estrogen resulted in a striking 480% increase in methyltrienolone (a potent oral androgen often used to reference receptor binding in studies) binding in the levator ani muscle 34 The suggested explanation is that estrogen must either be directly stimulating androgen receptor production or perhaps diminishing the rate of receptor breakdown Although the growth of the levator ani muscle is commonly used as a reference for the anabolic activity of steroid compounds, it is admittedly a sex organ muscle, and different from skeletal muscle tissue in that it possesses a much higher concentration of androgen receptors. This study, however, did look at the effect of estrogen in fast-twitch skeletal muscle tissues (tibialis anterior and extensor digitorum longus) as well, but did not note the same increase as the levator ani. Although discouraging at first glance, the fact that estrogen can increase androgen receptor binding in any tissue remains an extremely significant finding, especially in light of the fact that we now know androgens to have some positive effects on muscle growth that are mediated outside of muscle tissue.

Estrogen and Fatigue

"Steroid praigue" is a common catchphrase these days, and refers to another important function of estrogen in both the male and fermal body, namely is ability to promote wakefulness and a mentally alert state. Given the common availability of potent third-generation aromatase inhibitors, bodybuilders today are (at times) noticing more extreme estrogen suppression than they had in the past. Othen associated with this suppression is fingle: Under state, conditions, the athlete, though on a productive cycle of drugs, may not be able to maximize this or her gains due to an inability to train at full vigar. This effect is sometimes also dubbed "steroid lettargy." The reason is that estrogen plays an important supporting role in the activity of serotomi. Secrotomi is one of the body's principle neurotransmitters, vital to mental alertness and the sleepvikae cycle.³⁵ Min furgineer with the incurvations neurotransmitter is also associated with chronic furgine suppression funges any second as the ist to future specificable. Theorem suppression in menopause has also been associated with fatigee.³⁹ as has the clinical use of newer (more potent) aromatase inhibitors like anastrozok.⁴⁰ letrozok effectiventes, effe

Anti-Estrogens and the Athlete

So what does this all mean to the bodybuikler booking to gain optimal size? Bascially 1 think te calls for a cautious approach to the use of estrogen maintenance drugs if mass is the key objective (things change, of course, if we are taking about cating). Obviously, anti-storgosmy should be used if there is a clear need for them due to the onset of estrogenic side effects, or at the very least, the drugs being administered should be substituted for nonestrogenic compounds. Gynceomastia is certainly an unvanted problem for the steroid user, as are noticeable far times gains. Build is the solution of most optimate of the sterogenic strength in the solution of musice mass, or keeping you energetic. An individual confident they will notice, or are not prove to getting, estrogenic side effects, my therefore want to hold off using estrogen maintenance drugs so as to achieve the maximum possible gains in tissue mass.

²² Aromatization of androgens by muscle and adipose tissue in vivo. Longcope C, Pratt JH, Schneider SH, Fineberg SE. J Clin Endocrinol Metab 1978 Jarv46(1):146-52

²³ The aromatization of androstenedione by human adipose and liver tissue. J Steroid Biochem, 1980 Dec;13(12):1427-31.

²⁴ Aromatase expression in the human male. Brodie A, Inkster S, Yue W. Mol Cell Endocrinol 2001 Jun 10;178(1-2):23-8

²⁵ A review of brain aromatase cytochrome P450. Lephart ED. Brain Res Brain Res Rev 1996 Jun;22(1):1-26

²⁶ Aromatization by skeletal muscle. Matsumine H, Hirato K, Yanaihara T, Tamada T, Yoshida M. J Clin Endocrinol Metab 1986 Sep;63(3):717-20

^{27.} Pentose Cycle Activity in Muscle from Fetal, Neonatal and Infant Rhesus Monkeys. Arch Biochem Biophys 117:275-81 1966

28 The pentose phosphate pathway in regenerating skeletal muscle. Biochem J 170: 17 1978

²⁹. Aromatization of androgens to estrogens mediates increased activity of glucose 6phosphate dehydrogensee in rat levator ani muscle. Endocrinol 106(2):441–431 (1980)

^{30.} Influence of tamoxifen, aminoghtethinide and goserelin on human plasma IGF-1 levels in breast cancer patients. J steroid Biochem Mol Bio 41:541- 3,1992

^{31.} Activation of the somatotropic axis by testosterone in adult males: Evidence for the role of aromatization. J Clin. Endocrinol Metab 76:1407-12 1993

³² Testosterone administration increases insulin-like growth factor-I levels in normal men. J Clin Endocrinol Metab 77(3):776-9 1993

³³ Androgen-stimulated pubertal growththe effects of testosterone and dihydrotestosterone on growth hormone and insulin-like growth factor-1 in the treatment of short stature and delayed puberty. J Clin Endocrinol Metab 76(4)996-1001 1993

³⁴ Modulation of the cytosolic androgen receptor in striated muscle by sex steroids. Endocrinology. 1984 Sep;115(3):862-6.

^{35.} Effect of estrogen-serotonin interactions on mood and cognition. Zenab Amin et al. Behav Cogn Neurosci Reviews 4(1) 2005:43-58

³⁶ Serotonin and the skeep/wake cycle: special emphasis on miscodialysis studies. Chiara M Portas et al. Progress in Neurology 60(200) 13-35.

 Reduction of serotonin transporters of patients with chronic fatigue syndrome. Numerous 2004 Dec 2:15(17):0571.4 Neuroreport 2004 Lieu 3;13(17):2371-4

³⁸ Association between serotonin transporter gene polymorphism and chronic fatigue syndrome. Narita M et al. Biochem Biophys Res Commun 2003 Nov 14;311(2)264-6

³⁹ Premenstrual Syndrome. Dickerson LM et al. Am Fam Physician 2003 Apr 15;67(8):1743-52

 Phase II trial of anastrozole in women with asymptomatic mullerian cancer. Gynecol Oncol. 2003 Dec;91(3):596-602.

 Letrozole. A review of its use in postmenopausal women with advanced breast cancer. Druss. 1998 Dec;56(6):1125-40. Review.

42. Exemestane: a review of its clinical efficacy and safety. Breast. 2001 Jur;10(3):198-208.

^{43.} A study of fadrozole, a new aromatase inhibitor, in postmenopausal women with advanced metastatic breast cancer. J Clin Oncol. 1992 Jar(10(1):111-6.

DHT Conversion

As we see from our discussion with estrogen, in considering the physiological effects of any steroid, we must look at all of its active metabolits, and not just the initial compound. This includes not only estrogenic products, but androgenic metabolites as well. With this in mid, it is important to note that the potency of testosterone is considerably increased in many androgen responsive tissues when it converts to didydatectosterone. More commonly referred to by the three-letter abbreviation DHT, this hormone is, in fact, measured to be approximately three to four times storage than testosterone. It is the most potent steroid found naturally in the human body, and important to discussi if we are to understand the full activity of testosterone, as well as other anabolic/androgenic steroids that undergo a similar conversion.

Testoatenone is converted to dihydrotestoatenone upon interaction with the 5alpha reductases enzyme. More specifically, this cruyper nerrows the (4-5double-board of testoatenone by the addition of two hydrogen atoms to its structure (there the name di-hydrot testosterone). The removal of this board is important, as in this case it creates a steroid that binds to the androgen receptor much more availy than its parent steroid. Schaha reductase is present in high amounts in fusieus of the prostate, akin, scalp, liver, and various regions of the central nervous system, and as such expressents a mechanism for the body to increase the potency of testosterone specifically testosterone way and than its way to the receptor without being converted to dihydrotestosterone, making DHT by far the active form of andocen here.

DHT and Androgenic Side Effects

In some regards this local potentiation of testosterone's activity may be unvectore, as higher androgenic activity in certant insisses may produce a number of undesirable side effects. Acne, for example, is often triggered by didyetoetosterone activity in the selacous signids, and the local formation of dihydrotestosterone in the scalp is typically blumed for triggering male pattern hair loss. You should know that it is a terrible microcreption among bodybuilders that dihydrotestosterone is an isolated culpir when it comes to anables and andmogenic, through the same culbur androgen receptor. Dihydrotestosterone of this receptor than most, and can be formed locally in certain androgen-sensitive tissues. All steroids can cause androgenic side effects in direct relation to their affinity for this receptor, and DHT has no known unique ability in this regard.

Benefits of DHT

While a bot of attention is being pail to the negative side effects of the androgen didydoctosterence, you should know that there are some known benefits to the strong androgenic activity brought about by this hormone as well. For example, DHT pays an important role in the organization and functioning of the central nervous system. Many neural cells contain active androgen receptors, and it is thought that there may even be a specific importance of dihydrotestosterone in this areas of the body. Studies have shown DHT to have a profoundly greater impact in these cells compared to testosterone. More specifically, animal models demonstrated that both testosterone and DHT would result in increased at the twenty-one hour profileration in neural cells three and seven hours after being administered, however only DHT was able to sustain this increase at the twenty-one hour to sustain the increase at the twenty-one hours.

mark.⁴⁴ Although some might contend that this difference is simply due to DHT forming a more stable and lassing complex with the androgen receptor, others suggest that DHT and testosterone might even be affecting neural cells differently. such that the diffyortestosterone-receptor complex and testosterone-receptor complex might be activating the transcription of different target genes.

The strong interaction between the central nervous system and skeletal mascks, collectively referred to as the neuromuscular system, is of kysimportance to the athlete. There appears to be filte doubt that the ability of the body to adapt to training, and to activate nerve endings in musck itsus, is relatin on the interactions of the neuromuscular system. Inhibiting the formation of DHT during a testotteence cycle may therefore inadvertenity interfere with strength and muscle mass gains. This would explain why bodyhuiders commonly report a drop in stervid pettery, when they add the 5-alphar calcutase inhibitor finasteride to a testostrone cycle. Many complain is added, which would not make sense if testostrone and androgar receptor a divitation in muscle tissa were solely responsible of grouts (Clarity more is involved, and we cannot look at dhydrotestosterone singly as a side-effect hormone.

⁴⁴. Neural androgen receptor regulation: effects of androgen and antiandrogen. Lu S, Simon NG, Wang Y, Hu S. J Neurobiol 1999 Dec;41(4):505-12

Brief History of Anabolic/Androgenic Steroids

While it had been clear for many centuries that the testicles were crucial for the male body to properly develop, it was not unil modern times that an understanding of testosterone began to form. The first solid scientific experiments in this rate, which eventually led to the discovery and replication. During this century a number of animal experiments were published, most of which involved the removal and/or impaintation of resistorial material from's a subject. Although very crude in design by today's standards, these studies ertainly kink from androgen injusticans. These were standards either the studies of hormones). By the turn of the century, scientists were able to produce the first experimental androgen injusticans. These were standard either through testosterone from animal testics. Again, the methods were rough but the final results proved to be very englishting. Chemiss finally synthesized the structure of testosterone in the mid-1990's, sparking a new wave of interest in this hornoose. With the medical community paying a tremendous amount of attention to this achievement, the possible therapeutic uses for a readily available synthetic testosterone quickly became an externely popular focus. Many believed the applications for this type of a medication would be externed prior tachering with uses ranging from the maintenance of an androgen deficiency, to that of a good health and wellbeing treatment for the sixkly or elderly. During the infancy of such experimentation, many believed they had crossed paths with a true "fountain of youth."

Dihydrotestosterone and nandrolone, two other naturally occurring steroids. were also isolated and synthesized in the early years of steroid development. To make things even more interesting, scientists soon realized that the androgenic, estrogenic, and anabolic activity of steroid hormones could be adjusted by altering their molecular structure. The goal of many researchers thereafter became to manufacture a steroid with extremely strong anabolic activity, but will display little or no androgenic/estrogenic properties. This could be very beneficial, because side effects will often become pronounced when steroid hormones are administered in supraphysiological amounts. A "pure" anabolic would theoretically allow the patient to receive only the beneficial effects of androgens (lean muscle mass gain, increased energy and recuperation, etc.), regardless of the dosage. Some early success with the creation of new structures convinced many scientists that they were on the right track. Unfortunately none of this progress led researchers their ultimate goal. By the mid-1950's, well over one thousand testosterone, nandrolone, and dihydrotestosterone analogues had been produced, but none proved to be purely anabolic compounds.

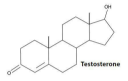
The failure to reach this goal was primarily due to an initial flawed understanding of testosterone's action. Scientists had noticed high levels of DHT in certain tissues, and believed this indicated an unusual receptor affinity for this hormone. This led to the belief that the human body had two different androgen receptors. According to this theory, one receptor site would respond only to testosterone (eliciting the beneficial anabolic effects). while the other is activated specifically by the metabolite. dihydrotestosterone. With this understanding, eliminating the conversion of testosterone to DHT was thought capable of solving the problem of androgenic side effects, as these receptors would have little or none of this hormone available for binding. More recently, however, scientists have come to understand that only one type of androgen receptor exists in the human body. It is also accepted that no anabolic/androgenic steroid can possibly be synthesized that would participate only with receptors in tissues related to anabolism DHT which was once thought not to bind to the same receptor as testosterone, is now known to do so at approximately three to four times the affinity of its parent, and the unusual recovery of DHT from androgen responsive tissues is now attributed to the distribution characteristics of the 5a-reductase enzyme.

Synthetic AAS Development

In order to develop products that would be effective therapeutically, chemists needed to solve a number of problems with using natural steroid hormones for treatment. For example, oral dosing was a problem, as our basic steroids testosterone, nandrokone, and dihydrotestosterone are ineffective when administered this way. The here would efficiently hread, down their structure before reaching circulation, so some from of alteration was required in order for a tablet or capsule to be produced. Our natural steroid hormones also have very short half-levs in the body, so when administered by higection, an extremely frequent and uncomfortable dosing schedule is required if a steady blood level is to be achieved. Therefore, extending stead activity was a major goal for many chemists during the carly years of synthetic AAS development. Scientistics also forcused on the nagging problems of possible excess entrogenic buildap in the blood, particularly with setsosterone, which can become very uncomfortable for patients underspoint therapy.

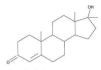
Methylated Compounds and Oral Dosing

Chemists realized that by replacing the hydrogen atom at the steroid's 17th alpha position with a carbon atom (a process referred to as alkylation), its structure would be notably resistant to breakdown by the liver. The carbon atom is typically added in the form of a methyl group (CH3), although we see oral steroids with an added ethyl



(C2H5) grouping as well. A steroid with this alteration is commonly described as a C1-17 alpha highed oral, although the terms of methyliated or ethylated oral steroid are also used. The alkyl group cannot be removed metabolically, and therefore inhibits reduction of the steroid to is insaviro 17-kotosteoid form by occupying one of the necessary carbon bonds. Before long, pharmacetrial companies had utilized this advance (and others) to manufacture an array of effective card steroids facilitated (and others) to Answer. The principle davaback to these compounds is that they phace notable amount of stress on the liver, which in some instances can lead to actual damage to this comm.

Because the alkyl group cannot be removed, it mediates the action of the steroid in the body. Methylestoterone, for example, is no simply an oral equivalent of testoterone, as the added alkylation changes the activity of this steroid considerably. One major change we see is an increased tendency for the storoid to produce estrogenic side effects, despite the fact that a actually lowers: the ability of the hormone to increat with aromates 4^{45} Apparently with 17-alkylation present on a steroid, aromatization (when possible) produces a more active form of extogen (typically 17-Japha-methyl to 17alpha-chyl estradiol). These estrogens are more biologically active than steriadiol due to the longer half-lift and weaker tendency to bind with semu proteins. In some instances, 17alpha-alkylation will also enhance the ability of the initial steroid compound to bind with and activate the estrogen or progesteroure receptor. An emanacement or exorogene properties is also obvious when we look at methandrostenolone, which is an alkylated form of boldenone (Equipoise®), and Nilevar, which is an alkylated form of the mild anabolic nandrolone. Dianabol is clearly more estrogenic than Equipoise®, a drug not noted for producing strong side



Methyltestosterone (+ 17alpha methyl)

effects of this nature. The same holds true for the comparison of Nilevar to Deca-Durabolin, a compound that we also know to be extremely mild in this regard.

C17 alpha alkylation also typically lowers the affinity in which the steroid binds to the androgen receptor, as is noted with the weak relative binding affinity of such popular agents as Dianabol and Winstrol (stanozob). However, since the alteration also greatly prolongs the half-life of a steroid, as well as increases the tendency for it to exist in a unbound state, it creates a more potent anabolicandrogenic agent in both cases. This explains why Dianabol and stanozobal are notably effective in relatively lower weekly dones (often 140 mg weekly will produce notable growth) compared to injectables such as testosterone and nandrokene, which often need to reach dones of 300-400 mg weekly for a similar level of effect.

Non-Alkylated Orals

In an attempt to solve the mentioned problems with liver toxicity we see with c17-alpha alkylated compounds, a number of other orals with different chemical alterations (such as Primobolan®, Proviron®, AndriolE, and Anabolicum Vister) were created Primobolan® and Proviron® are alkylated at the one position (methyl), a trait which also slows ketosteroid reduction. Andriol® uses a 17beta carboxylic acid ester (used with injectable compounds, discussed below), however, here the oil-dissolved steroid is sealed in a capsule and is intended for oral administration. This is supposed to promote steroid absorption through intestinal lymphatic ducts, bypassing the first pass through the liver. In addition to 1 methylation, Primobolan® also utilizes a 17 beta ester (acetate) to further protect against reduction to inactive form (here there is no lymphatic system absorption). Anabolicum Vister uses 17beta enol ether linkage to protect the steroid, which is very similar to esterification as the ether breaks off to release a steroid base (boldenone in this case). While all of these types of compounds do not place the same stress on the liver, they are also much less resistant to breakdown than 17 alkylated orals, and are ultimately less active milligram for milligram.

Esters and Injectable Compounds

You may notice that many injectable steroids will list long chemical names like testotoreor cylonate and testostrome en anthate, instead of just testostrome. In these cases, the cypionate and examble hydroxyl group of the testostrome molecule, which increase the active life span of the steroid preparation. Such alterations will relate the steroid's provide of water solubility and increase its oil solubility. Once an esterified compound has been injected, it will form a deposit in the macked insue (deposit) from which it will knowl at such as the steroid steroid steroid steroid steroid steroid steroid steroid compound. Which may also the steroid's teroid compound has been injected, attribute the steroid steroid steroid steroid steroid steroid steroid steroid compound. The steroid steroid steroid steroid steroid steroid steroid compounds will be for the case of the full donage to be related the parent hormoon will be free to exert is activity (while the ester is present the steroid is incrit).

There are a wide number of esters, which can provide varying release times, used in medicine toddy. To compare, an ester Rie deconnet can extend the release of active parent drug into the blood stream for three to four weeks, while it may only be extended for a few days with an accuted or proponent than if using a water-based (artagdh) testosterone, which is much more comfortable for the patient. We must remember when calculating dosagas, that the ester is figured into the steroid's measured weight. 100 mg of testosterone ensuring, there in the steroid's measured weight, 20m gof testosterone ensuring, the steroid's measured weight. 20m gof testosterone instances, an ester may accound for roughly 40% or around 10% to 55%. Below relax the the typical measure is somewhere around 10% to 55%. Below relax the three typical measure is somewhere around 10% to 55%. Below relax the three typical measure is somewhere around 10% to 55%. Below relax the total set of the source is somewhere around compared.

| 100 mg of steroid as: | Approximate Free Equivalent: | |
|-----------------------------|------------------------------|--|
| Trenbolone acetate | 87 mg | |
| Testosterone propionate | 83 mg | |
| Testosterone enanthate | 72 mg | |
| Testosterone cyplonate | 70 mg | |
| Testosterone undecanoate | 63 mg | |
| Nandrolone phenylpropionate | 67 mg | |
| Nandrolone decanoate | 64 mg | |

It is also important to stress the fact that exters do not alter the activity of the parent stretoid an any ways. They work only to slow its relaxe. It is quite common to hear people speak about the properties of different exters, almost as if they can magnedial plater a stretoid effectiveness. This is really nonsense. In the second stretoin the stretoin stretoing and the stretoing and interpret of the stretoing of the stretoing and the stretoing and about the stretoing and the stretoing and the stretoing and the always considered Statianton a very poor buy in the face of cheaper 250 mg enaminate amputes. Your musice clean see only its storetore, ultimately there is no difference. Reports of varying levels of musice gain, androgenie side effects, water retention, etc. are only sisses of timing. Faster releases the store streto will produce estrogen building faster amply because there they when we state that. Durabolish is a midler machine faster for the store we have as state that. Durabolish is a midler machine faster stretoing. compared to Deca. It is simply easier to control the blood level with a faster acting drug. Were virilization symptoms to become apparent, hormone levels will drop much faster once we stop administration. This should not be confused with the notion that the nandrolone in Durabolin8 acts differently in the body than that released from a shot of Deca-Durabolin8.

It is also worth noting that while the ester is typically hydrolyzed in general circulation, some will be hydrolyzed at the injection site where the steroid depot first contacts blood. This will cause a slightly higher concentration of both free steroid and ester in the muscle where the drug had been administered. On the plus side, this may equate to slightly better growth in this muscle, as more hormone is made available to nearby cells. Many bodybuilders have come to swear by the use of injection sites such as the deltoids, biceps, and triceps, truly believing better growth can be achieved if the steroid is injected directly into these muscles. The negative to this is that the ester itself may be irritating to the tissues at the site of injection once it is broken free. In some instances it can be so caustic that the muscle itself will become swollen and sore due to the presence of the ester, and the user may even suffer a lowgrade fever as the body fights off the irritant (the onset of such symptoms typically occurs 24-72 hours after injection). This effect is more common with small chain esters such as propionate and acetate, and can actually make a popular steroid such as Sustanon (which contains testosterone propionate) off-limits for some users who experience too much discomfort to justify using the drug. Longer chain esters such as decanoate and cypionate are typically much less irritating at the site of injection, and therefore are preferred by sensitive individuals.

Anabolic/Androgenic Dissociation

Although never complete, scientists had some success in their quest to separate the androgenic and anabolic properties of testosterone. A number of synthetic anabolic steroids had been developed as a result, with many being notably weaker and stronger than our base androgen. In order to first assess the anabolic and androgenic potential of each newly developed steroid. scientists had generally used rats as a model. To judge androgenic potency the typical procedure involved the post-administration measure (% growth) of the seminal vesicles and ventral prostate. These two tissues will often respond unequally to a given steroid, however, so an average of the two figures is used. Anabolic activity was most commonly determined by measuring the growth of the levator ani, a sex organ (not skeletal) muscle. This tissue may not be the most ideal one to use though, as it contains more androgen receptor than most skeletal muscles (the AR is still less abundant here than in target tissues such as the ventral prostate).47 48 In integrating both measures, the anabolic index is used, which relates the ratio of anabolic to androgenic response for a given steroid. An anabolic index greater than one indicates a higher tendency for anabolic effect, and therefore classifies the drug as an anabolic steroid. A measure lower than one in turn assesses the steroid as androgenic. There is some variance between experimental results and the actual real world experiences with humans, but (with a few exceptions) designations based on the anabolic index are generally accepted. Below are discussed a few factors that greatly affect anabolic/androgenic dissociation

Nandrolone and 19-norandrogens

The section of this book dealing with DHT conversion is important, because it helps us understand the anabolic steroid nandrolone and many of its derivatives. Nandrolone is identical to testosterone except it lacks a carbon atom in the 19th position, hence its other given name 19-anortesolsterone. Nandhovine is very interesting because in offers the greatest ruits of anabole to androgenic effect of the three natural steroids (see <u>Spinderic ASS</u> (<u>Funderic</u>). This is because it is metholized into a less potent structure (dhydroandrolone) in androgen target tissues with high concentrations of the S-alphar relactase enzyme, which is the exact opposed of what happens with testosterone. Apparently the removal of the c4-5 double bond, which normally increases the androgen receptor building capability of testosterone, causes an unusual lowering of this ability with nandrolone. Instead of becoming three to four times more potent, it becomes sevent limes weaker. This is a very desirable trait if you want to target anabole effects over androgenic. This characteristic also carries vero to mos synthetic storoids derived from nandrolone, making this an attractive base steroid to use in the synthesis of new, primarily nanbole, storoids.

5-alpha Irreducible Steroids

When we look at the other mild anabolic steroids Primobolan®, Winstrol®, and Anavar, none of which are derived from nandrolone, we see another interesting commonality. These steroids are DHT derivatives that are unaffected by 5alpha-reductase, and therefore become neither weaker nor stronger in androgen responsive target tissues with high concentrations of this enzyme. In essence, they have a very balanced effect between muscle and androgen tissues, making them outwardly less androgenic than testosterone. This is why these steroids are technically classified as anabolics, and are undeniably less troublesome than many other steroids in terms of promoting androgenic side effects. However, if we wanted to look for the absolute least androgenic steroid, the title would still go to nandrolone (or perhaps one of its derivatives). Female bodybuilders should likewise take note that despite the recommendations of others, steroids like Anavar, Winstrol and Primo are not the least risky steroids to use. This is of great importance, as male sex hormones can produce many undesirable and permanent side effects when incorrectly taken by females (See: Side Effects, Virilization),

3-alpha Hydroxysteroid Dehydrogenase

The 3-alpha hydroxysteroid dehydrogenase enzyme is also important, because it can work to reduce the anabole potency of certain stroxids considerably. As follows, not all potent binders of the androgen receptor are, as a rule, great muscle-building drugs, and this enzyme is an important factor. Dihydrotextosterose is a clear example. Just as the body converts testosterone to DHT as a way to potentiate its action in certain instance (sikn scalp, possible, etc.), it also has ways of countering the strong activity of DHT, in other issues where it is unneeded. This is accompliable by the mpda reduction of DHT to its materies active metabolities, mmedy win interaction with B-alpha bydroxystenid dehydrogenase enzyme. This enzyme is present in high concentrations in certain tissues, including sclekial muscle, and DHT is much more open to alteration by it than other stretostic

that posses a c+5 double-bond (like testoterono)⁴⁰. This causes disploctoststress to be an externely poor anabolic, despite the fact that is actually coholism a much higher affinity for the coholism androgen receptor than most other stretchis. Were it able to reach the coholism androgen receptor than most other stretchism. Were it ables to reach the coholism androgen receptor without fract being metaboliced by 3a-HSD, it certainly would be a formiable muscle-building stretchism (burden the stretchism) and the socommerciality protocols were never foreorized to an androgen build muss. This main is also shared by the currently popular eral androgen build muss. This main is how shared by the currently popular eral androgen burdent muscle is in anomene into an other one of DWT of neuroistic. riovanie, which is, in essence, just an oral form of Drif (1-menyi dihydrotestosterone to be specific) and known to be an extremely poor tissue builder.

Anabolics and Potency

One must remember that being classified as an anabolic just means that the steroid is more inclined to produce muscle growth than androgenic side effects. Since both effects are mediated through the same receptor, and growth is not produced by androgen receptor activation in muscle tissue alone (other CNS tissues, for example, are integral to this process as well). we find that a reduction in the androgenic activity of a compound will often coincide with a similar lowering of its muscle-building effectiveness. If we are just looking at overall muscle growth, androgenic steroids (usually potent due to their displaying a high affinity to bind with the androgen receptor in all tissues) are typically much more productive muscle-builders than anabolics. which usually bind with lower affinity in many tissues. In fact, with all of the analogues produced throughout the years, the base androgen testosterone is still considered to be one of the most effective bulking agents. The user must simply endure more side effects when acquiring his or her new muscle with this type of drug. Individuals wishing to avoid the stronger steroids will. therefore, make a trade-off, accepting less overall muscle gain in order to run a more comfortable cycle.

RBA Assay:

Another way of evaluating the potential ratio of anabolic to androgenic activity is the practice of comparing the relative binding affinity (RBA) of various steroids for the androgen receptor in rat skeletal muscle versus prostate. When we look at the detailed study published in 1984, we see some recognizable (and expected) trends. Aside from dihydrotestosterone and Proviron® (mesterolone), which undergo rapid enzymatic reduction in muscle tissue to inactive metabolites, the remaining anabolic/androgenic steroids seem to bind with near equal affinity to receptors in both tissues. They seem to be relatively "balanced" in effect. This study also discusses the unique activity of testosterone and nandrolone compounds, which are good substrates for the 5a-reductase enzyme found in androgen target tissues (such as the prostate), and seem to provide the most notable variance between anabolic and androgenic effect in humans due to this local metabolism. When it comes to real-world use in humans, anabolic steroids do not always behave in 100% uniformity with their anabolic and androgenic profiles as determined by animal models, so all such figures need to be taken with a small grain of calt

| Compound | Human SHBG | Rabbit Muscle | Rat Muscle | Rat Prostate | Ratio M vs. P |
|---------------------|------------|---------------|------------|--------------|---------------|
| methyltrienolone | <.01 | 1 | 1 | 1 | 1 |
| dihydrotestosterone | 1 | .07 | <.01 | .46 | .03 |
| mesterolone | 4.4 | .21 | .08 | .25 | .32 |
| testosterone | .19 | .07 | .23 | .15 | 1.53 |
| nandrolone | .01 | .20 | .24 | .60 | .4 |
| methyltestosterone | .05 | .1 | .11 | .13 | .85 |
| methenolone | .03 | .09 | .24 | .14 | 1.67 |
| stanozolol | .01 | .03 | .02 | .03 | .6 |
| methandrostenolone | .02 | .02 | .02 | .03 | .75 |
| fluoxymesterone | <.01 | .02 | .01 | .02 | .77 |
| oxymetholone | <.01 | <.01 | <.01 | <.01 | 1.54 |
| ethylestrenol | <.01 | .01 | <.01 | <.01 | 2 |

RBA of various anabolic/androgenic steroids as competitors for human SIBG binding of DHT, and for receptor binding of methyltrenolone in cytosol from rabbit, rat skeletal muscle and prostate. Source: Endocrinology 114(6):2100-66 1984 June, "Relative Binding Affinity of Anabolic-Androgenic Steroids...", Saartok T; Dahlberg E; Gustafson JA 2, July Section, J H-edits2, Jul 11/1/10 10557 PM Page 19

⁴⁵ A comparative study of the metabolic fate of testosterone, 17alpha-methyltestosterone, 19-nor-testosterone, 17alpha-methyl-19-nor-testosterone and 17alpha-methyl-estr-5(10)ene-17beta-ol-3-one in normal males. Dinick D, Heron M, et al. Clin Chim Acta 6(1961) 63-71.

⁴⁶. Unique steroid congeners for receptor studies. Ojasoo T, Raynaud J. Cancer Research 38 (1978) 4186-98

 Cytosolic androgen receptor in regenerating rat levator ani muscle. Max S.R. Mufti S, Carlson B.M. J Biochem 200 (1981) 77

^{48.} In vitro binding and metabolism of androgens in various organs: a comparative study. Kreig M., Voigt K.D. J Steroid Biochem 7 (1976) 1005

⁴⁹ Androgen concentrations in sexual and non-sexual skin as well as striated muscle in man. Deslypere J.P., Sayed A., Verdonck L., Vermeulen A. J Steroid Biochem 13 (1980) 1455-8

Synthetic AAS Chemistry



Steran Nucleus (All natural and synthetic AAS hormones share this base structure)







Nandrolone

Al analogicalization across a preparations containing one of the above free natural seriod hormosce, or chemicaly alared deviations thereof. In creating new synthesis to provide the contrained particular distribution of the provided particular that the possission of particular tasks that up be hording if the more proparal. For isstance, of the three natural asteroids above, displottatostorence is the only steerid deviation of synthesis and synthesis that had, estrogenic activity and/or collide a more balanced antiogenic to ambidity and and the strongenic activity and/or collide a more balanced antiogenic to ambidity and the strongenic activity and/or collide a more balanced moltance express. Nankubiene adia anomations may and/or collide a more balanced and/or activity data to be conversion to a more potent streaming and openic activity data to be screaming and/openic activity data to be conversion to a more potent more activity than toxistorience.

Testosterone derivatives

Boldenone

Bolkmon is tentostence with an added double-bond between entron atoms one and two. However, this bond tranges the activity of the stread considerable. First, it durantically shows anomatinity, such that bolkmone converts to estability at both all the rate of testostences. Scendby, this bond causes the stread to a vary low substante for the Calphan outcase enzyme. The more active 3-adpha reduced methodic Salphadingehoolschemes expression and materia in humans. The human issues in the stream of the stream of the stream of the stream of the stream both mains and all the stream of the stream of the stream of the stream of the both mains and all more that the stream of the stream of the stream of the stream of the length is buckdown of the stream, iteraway is resistance to 17-ketostenid duscitation and is functional life and con lineouslibility.

Methyltestosterone

This is the most basic derivative of testosterone, differing only by the added 17- alpha methylation that makes the steroid onally active. Conversion to 17-alpha methylastradiol makes this steroid extremely estrogenic, despite the fact that this alteration actually reduces interaction with the aromatase enzyme.

Methandrostenolone

In mure grapits, methanitostolone is very similar to bolkence, as it to cubits reduced subsparies and analysing activity due to the c1-2 double-bond. However, this staroid does have a repatition of being semewhat estrogenic, oxing to the fact that it concerts to a high active form of estrogen (Tapharendepistandi Sae: Adducta Comparate, and Chai active form of estrogen (Tapharendepistandi Sae: Adducta Comparate, and Chai active form of estrogen (Tapharendepistandi Sae: Adducta Comparate, and Chai active form of estrogen (Tapharendepistandi Sae: Adducta Comparate, and Chai adducta Comparate active for a star of the star of the star and the star of the star of the star of the star of the star and the star of estimation of the star of the star of the star and the star of the star and the star of the star and the star of the star and the star of the s

Fluoxymesterone

Fluxymesterone is a \sim 17abpha alcylated ond derivative of testosterone. The 11-beta group functions to inhibit aromatization, so there is no estrogen conversion at all with this steroid. It also works to lower the affitity of this steroid toward restrictive semum binding proteins, increasing its relative activity. Introduction of fluorine at the 9-position also potentiates the action of this steroid.

Nandrolone derivatives

Norethandrolone

Noethandolore is simply matheticate with an added 17-alpha endy group. This aleration is mady used with analobicandrogenic struction, and is much more commonly found with spathack estruggens and progesties. Although 17-endytakin sinkhas 17-Actostoval reduction as a wells a 37-methylation, and Renderal above fisse struct to eithis a similarly high level and a structure and the structure and the structure and the structure and the Needmatochee is clarify a Twodebiscent' hermices in terms of vator relations, fits granted Needmatochee is clarify a Twodebiscent' hermices in terms of vator relations, fit granted and granceomstik, which may near the date to its heighteen buildings to this receptor.

Ethylestrenol

Ethjesterool is an oral derivative of markotoe, very simler is structure to keto group, which is with a lowagen neceptor brinking, as such, ethjesterool is possibly the weakest storoid miligam for miligam ever sold commercially. Any activity the steroid close oldbit is dely form is conversion in northankolous, which does sense to occar with some affitty touble). This is probably the most interesting that of ethylesteroel, which is an undistingished converted other.

Trenbolone

Althcigh a derivative of matrichoux, the two additional double-bends present on meto-bone make any similaries to its parent fromework externels of the sec. First, the 9-10 bend iribbs aromitation. Nandrobene is very dowly aromited, however, some entrogen is still produced from this section. Near sol when benchows. The 11-2 bend additionally neurons, produced from this section. Near sol when the source of the section of the section mathematic and an such does not share the same dissociation between analytic and andoragie effects (networksen is much more androgenic in conversion).

Dihydrotestosterone derivatives

Mesterolone

Mesterobne is a potent orally active derivative of dihydrotestosterone. Similar to methenolone, it possesses a non-toxic 1-methyl group, which increases its resistance to hepatic breakdown. This alteration does not increase the stability of the 3-keto group however, and as such, this steriol is a poor anabole like its parent.

Drostanolone

Drostanolone is simply dihydrotestosterone with an added 2-methyl group. This addition greatly increases the stability of the 3-keto group, vital to androgen binding. As such, the activity of this steroid in muscle tissue is greatly enhanced (see: <u>Anabolic/Androgenic</u> <u>Dissociation</u>).

Oxymetholone

Osymetholone is an orally active derivative of dipdrotestosterone. The 17-methyl group is well understood at this point as we have discussed it with mmy steroids, however, the 2hydrosymethylene group is not seen on any other commercial steroid. We do know that this group group dynathylenese anabole potency by increasing the stability of the 3-shedo group, and that the configuration of this substituent also appears to allow this steroid to bird and activate the estrongen receptor.

Stanozolol

Stanozolol is a potent anabolic steroid, owing to the fact that the 3-2 pyrazol group creates a stable configuration off the A-ring that allows for androgen receptor binding (this steroid is one of the few that does not possess an actual 3-keto group). As such, it is highly active in macle tissue, unlike dhydrotestosterone.

Methenolone

Methenolme also is a potent anabolic steorid, due to the fact that the c1-2 double bond increases the stability of the 3-keto group. The 1-methyl group works to increase its oral bioavailability, making methenoloue (as methenoloue acetate) one of the few orally active non-17-aklylated orals. The c1-2 bond may also help increase hepatic resistance (slightly to 17-ketosteroid duectivation as well.

Oxandrolone

Orandrohene is an onally active derivative of follydrotestosterone, due to is 17-methylation. It also differs from DHT by the substitution of its 2-carbon molecule with oxygen. This is the only commercial steroid to carry this group, and further, the only to have a modification to the base carbon structure of the Steran nucleus. The 2-room group increases resistance of the 3-keto group to metholosim considerable, muching constructions a potent anabole.

Steroid Nomenclature

Perhaps not obvious at first glance, there is a naming convention in place that was used to create identities for the various anabolic/androgenic steroid hormones. This typically involves forming a not word to convey the structural loss of the structi, and signifying order unique structural characteristics by including appropriate prefets or sufficies. Relow, we will book the the common roots, profiles, and sufficies used in stearid momenchatra, and identify them as the common roots, profiles, and sufficies used in stearid momenchatra, and identify them, as they are used in the waves commencial composited mass. As a good waves are sufficient to stear the common roots, profiles and the momenchatra and the stear discussion for the normalized momenta of the stear of the sufficient of the discussion for the normalized mass of the stear of the stear of the stear interval in the cochasion use of the term roots are barrowed and interval the cochasion use of this terminology (and, as is the representation of metanulocatorobate and To-balgoots). (Promotelylandosts 1, 4-dam 5-one).

Common prefixes and suffixes used in steroid naming:

| Structural Property | Prefix | Suffix |
|---------------------|----------------|-----------|
| Carbonyl (C=O) | oxo-; keto- | -one |
| Hydroxyl | hydroxy- | -ol |
| Double Bnd (C=C) | | -ene; -en |
| Methyl | meth-; methyl- | |
| Ethyl | eth-; ethyl- | |

Common roots used in steroid naming:

| Androstane | Base carbon structure of dihydrotestosterone (no double-bond) | | |
|---------------------|--|--|--|
| Androstene | Base carbon structure of or similar to testosterone (one double-bond) | | |
| Androstadiene | Base carbon structure similar to methandrostenolone (two double-bonds; di-ene) | | |
| Estren; Estra | Base structure of nandrolone (19-norandrostene) and estrogen | | |
| also: Norandrostene | | | |

Common Commercial Compound Names:

| Name | Taken From | Incorporated Into Name As |
|--------------------|---|---------------------------|
| Boldenone | [17b-ol, androstadiene, 3-one] | BOL DEN ONE |
| Ethylestrenol | [17a ethyl, estren, 17b-ol] | ETHYL ESTREN OL |
| Fluoxymesterone | [9-fluoro, 11b-hydroxyl, 17a-methyl, testosterone, 3-one] | FLU OXY ME STER ONE |
| Mesterolone | [1-methyl, dihydrotestosterone, 17b-ol, 3-one] | ME STER OL ONE |
| Methandienone | [17a-methyl, androstadiene, 3-one] | METH ANDIEN ONE |
| Methandrostenolone | [17a-methyl, androstadiene, 17b-ol, 3-one] | METH ANDROSTEN OL ONE |
| Methenolone | [1-methyl, c1-2 double bond (en), 17b-ol, 3-one] | METH EN OL ONE |
| Nandrolone | [norandrostene, 17b-ol, 3-one] | NANDR OL ONE |
| Norethandrolone | [19-nor, 17a-ethyl, (nor)androstene, 17b-ol, 3-one] | NOR ETH ANDR OL ONE |
| Oxandrolone | [2-oxy, androstane, 17b-ol, 2-one] | OX ANDR OL ONE |
| Oxymetholone | [2-hydroxymethylene, 17a-Methyl, 17b-ol, 3-one] | OXY METH OL ONE |
| Stanozolol | [Stanolone (androstanolone, DHT), 2-pyrazol, 17b-ol] | STANO ZOL OL |
| Trenbolone | [tri-en, 17b-ol, 3-one] | TREN BOL ONE |

Clinical Applications

Aubolicantoganis attoriks are approved for sak by prescription in virtually every phermenetrical market around the work! Having been applied for many decades to reat a variety of discussed status, today these drugs have a narbre of web-eabbled medical uses. They have been used to near more patiere patientism, ischaffer men and voncen of almost all ages, maging from children to the clefely. In many instances anabolicantloganis starvisk have proven to be like swing medications, which is a fast casily overbooked with all of the discussion about storeid abuse. This section details some of the most common and accepted medical applications for anabolicantloganis storeids.

Androgen Replacement Therapy/Hypogonadism

The most widely used medical application for anabolic/androgenic steroids in the world is that of androgen replacement therapy. Also referred to as Hormone Replacement Therapy (HRT) or Testosterone Replacement Therapy (TRT), this therapy involves the consumeration of the reference make hormone to calonize converting of law supportantiation or use paramy inter institutes enconcenter to activitie symptome to two hormone levels (chicality efferent to as hopportantism). Printers my be adolescent males suffering from childhood hopportandism or a specific disorder that causes androgenic hormone disruption, alhough most of the treated population consists of adult nero over the age of 30, In most cases hormone levels have declined in these men as a result of the normal asing process.

The most common compliairs associated with two testosterouse in adda mon include reduced fibids, orecale displations, loss of energy, calcuend strength and prove outmance, reduced abidly to physic prosts, model flatatations, reduced height (bose loss), reduced work and the strength of the strength of the strength of the strength of the second strength of the strength of the strength of the strength of the second strength of the strength of 20 fight in the first of the strength of the strength of the strength of the 20 fight of the strength of the strength of the strength of the strength of the 20 fight of the strength of the stren

Androgen replacement therapy effectively allovistics must symptoms of low teststateme levels. To begat with using teststateme levels alsow 350 mg/dt (the way low end of the normal maps) will often restore normal sexual function and likels in men with dysfanctions related to hermones institutions. With regard to been rised and endy hormone replacement therapy is also documented to have a significant possible effect. For example, studies administrarg 250 mg of teststateme carefulne every 21 days showed a 5% rearcase in bones metaed clonely address its models.⁶⁰ Over time time imprevent some loss of height and house metaed clonely address its models.⁶¹ Over time time imprevent some loss of height and more some of the sing and may also reaches the risk of stratemer.⁶² There energy and sense of well-being Theory also apports the reterion of team body muss, and improves muck streament and endernace.

Likke storij abue, hormoze replacement fuenzje muj hvo benefis vihi regard to cardiovacidar disease risk. For example, statis tend o solo whome replacement as hvorig a positive effect on sexum ljuki. This includes a reduction in LL and teal chotserol levels, contribed with no singificant funging in HDL (good) chotserol levels.²⁵ To Testosterone amplementation also reduces milection obesity, and inproves inside sensitivity and glorent isolo model. The area energy for the sensitivity and glorent base has a horizontation of attenactive structures the reducent fragmentation plaques and area times. The model contensitivity and glorent has been about in proper the problem through multi-target model in the progression of attenactive base plaques and has the adaptament in plaques and sear times. It models consenses to ship supposes to b fund cardiorascut disease risk, and may actually decrease certain risk factors for the disease in some prators.

These are some concerns with initiating teststateons replacement theory when the individual is in poor hash. To subty control the startly of HTG in run angle G and older with limited melbity and various hash conditions such as obscity. hypertension, diabetes, or hyperpletions¹⁰. Sile control of the startly of the startly (10) (0) major picket gradies are startly as a startly obscit and the startly obscit and the startly obscit points and the startly obscit and the startly obscit and the startly obscit points and startly obscit and the startly obscit and the startly obscit points and the startly obscit and the startly obscit and the startly obscit points and the startly obscit and the startly obscit and the startly obscit points and the startly obscit point factor in a startly assist and the start disease, strong corributing factors to heart disease, or order erdore hashing conditions.

These are other areas of concern with cladely patients. To begin with, testostenere administration may recores prostate values and PAA values.²⁹ 29 White this does set appear to be of disical significance with normal healty patients, being prostate beportuply and prostate carear or as similarial by stacharone. May with prostate carear, high FAA values, or breast carear are generally not prescribed testostenere. Androgon appearation has also been hisked to shop super, which can interface with the most ender (REM) pipes of skeps⁶⁰. The statics have produced coefficing data, however, and de potential existencing memins the short of much dochand.¹² Ladak buschenere endpacement hency has demonstrated regardine, passive, and restand effects on cognitive functioning in delowhyrums.⁶² effects also data gasted multi the doce can catterine the local of approares, with the most possitive effects mored when the androgen hose fractions the neid-to upper-cargo of normal, not surgerspherically.⁶² Ehler partners with presenting deficit in cognitive function should have their cognitive performance and blood hommone levels monitored closely during hommone represent therapy.

Common Treatment Protocols:

<u>Transform</u>? Transform application is the root commonly prescribed method for supplementing tostschore in the LindStatus and Canada, and gas generally the fist coarse of therapy minical with androgan replacement therapy patterns. This method of drug clokery of the single structure is the structure of the structure of the significant structure of the single structure of the structure of the significant structure of the comparison. Transformal application also provides studied drug to-the harmone tests, and does not produce the brand flattation scale provides structure of the structure structure of a structure of the structure of the structure structure of the gal or athiesise transformed path that is replaced athie. Note that due to methods in the data's transformal pathetism of the structure structure structure structure state effects drug the structure of the structure structure structure structure state drugs thermation.

Injection: Testosterone enauthate and testosterone cynionate are the most widely prescribed injectable testosterone drugs in the United States and Canada. In many other markets the blended ester products Sustanon 100 and Sustanon 250 are also commonly prescribed Injection of one of these testosterone ester products will provide the patient supplemental androzen levels for annroximately 2 to 3 weeks after each application. The most common protocol among hormone replacement doctors is to administer 200 mg of testosterone enanthate or cynionate once every 2 to 3 weeks. It is important to remember that testosterone esters will deliver varying levels of testosterone to the body on a day-to-day basis throughout each application window. Levels will be highest the first several days after injection, and will slowly decline to baseline over the following weeks. Physicians are usually encouraged to monitor their patients closely to ensure androgen supplementation is sustaining hormone levels within the normal range (and alleviating symptoms of hypogonadism) throughout the entire therapeutic period. The longer acting injectable testosterone prenaration Nebido (testosterone undecanoate) is undergoing review in the U.S. and has already been approved in other markets. This drug requires only 4 to 5 injections per year for most patients.

One Teststorene undecanote (Audrio) is the only preservition medication that delivers teststorence via an out capacit. This medication is not approved for suls in the Urbit States, bit is a prescription drug in Canada and many other metheds around the world. Platicat compliance and confinit are high via this form of fitnergy, as focus are no special toststorene understored in the state of the state of the state of the state of the toststorene understored is and any state of the state of the state of the which capacits to there to far 40 are gravels. This chosen my be reacted in statespacers weeks to 12 m gper day. The capacits are given in two divided dones per day, with are usually taken with breakst and direct. While this fix mon fitnerging in highly convenient, serum hormone levels, can fluctuate gravity on a days-day basis. The amount of the consumption has a primers of far are econvented when histing the capacits for measurem of the structuation o

Angioedema, Hereditary

Anabolic steroids are commonly preserbed for the treatment of heneflary anginedema, a new and potentially life-frastening disorder of the immuse system. Heredaray angicedema is caused by genetic matations of blood cloting factors, characterized by a decrease in the level of functioning of the protein C1 estemase inhibitor. This protein controls C1, which is a "complement system" protein that plays an inportant role in the control of inflammation. Symptons of heredinary angiodema instellar an intermittene the rapid solution of the hands, ams, legs, lps, eyes, tongue, or throat. Swelling may also be noticed in the digistive tract, resulting in abdominal cramping nansea, or vomiting. In the most serious cases, the patient may notice a swelling of the throat and a blockage of the airway passages, resulting in asphysizion and sudden death. Many attacks occur without a specific trigger, athough stress, trauma, surgery, and dertail work are commonly associated with majoredom attacks.

Orde 1-2 Paphe adylated ambolic/antroperie steroick hnow been shown to be a useff ferm of preservine therapy, subhiring compresent system protein break and reducing the frequency and severy of amgiodeam anticks⁴⁰. They are usually administered in a low does, which is is to stead for large-game anguest of this dowlere? It multiple and the standard standard ducceries, allowagh bistocically umuny other agonts have also been prescribed in aking controller, methylanestical effective data and the standard standard ducceries, allowagh bistocically umuny other agonts have also been prescribed in inking controllers, methylanestical geffective data and the standard methylanestical the lowest threspectial geffective datages in an effort to offer uncertained is standard TDA approved prescribing galdeness for stanoaled incommended in ritid dotage of 2 ang lowed rate participation and the standard standard standard and a subst end therapited respective standard, and and the standard standard and standard s

Anemia

As a closs of dug, anaboki:induges: structus simular the synthesis of cylinopositin in the kinetys, a horner that resports the mundleture of new real bods cells. By doing thigher abinismitation of neurols structure in the synthesis of cell coart and humanscript level, characterized by ministerized real body cells. The synthesis is the synthesis is a structure of the synthesis reaction, specification of the synthesis is a synthesis of the structure of the synthesis reaction, specifications, sinceker-tony, sangevarge migelication metaphysis, and memiss caused by real structure, sinck cell anexis, reference metaphysis, and memiss caused by miniparacy or mytekoise drags. The level of response the munaments of a real metaphysis of the synthesis of the synthesis is a synthesis of the the munaments of a real metaphysis of the synthesis of the synthesy

In the Urida Status, food oxymetholore (Atundis S1) and rundvolore decanates (Daci-Danholin) are approved by the FDA for the number of severe anamis. The galdelines for using oxymetholues with boft multi and formal antenic patients (childron and adult) renormard a dosago of 12 anglegared 60.1 houses as high as 5 mg/kg/by are confinites receasing to achieve the desired therapeutic response. The galdelines for multiboursed for the desired therapeutic response. The galdenines for multibourwork for man. Children (2 to 13 years of age) are recommended a dosage of 25-50 mg every 31 of works.

In recet years, the advect of recordinate arythropoietin as a proceytion dug lues changed the face of around trustmet recordinghol. While anabole infraring strends still offer theraportic value hore, and are still markeded and solid to text americ pattern, they are presently regardle as adjust of thildness indications for use only when therapy with an orthopoint alone has field to achieve a dosined response. The hermaterit morease from antibilization and here scales a garenally be produced by any boots: than the means of the points, and here scales along a strength to produce the point frame the reserve optimization, and here scales and total to produce very metandic task of ficks that garent instances the risks to hereagy strength comparing the boords's of anabolication agars. Each strength to boots's of anabolication agars are strength to boots's of anabolication agars are strength as the first strength to points of the strength comparison. The strength to the strength comparison and the strength to the strength to the strength comparison and the strength comparison and the strength to the strength comparison and the strength to the strength comparison and the strength to the strength to the strength comparison and the strength to t

Breast Cancer

Anabolicantloganic stensis are constinues prescribed to treat beast cancer in postmenopasal sources or permenopasal sources how have have been ourse removed. These drugs are of value when the cancer is hormone responsive, which means that is growth can be affected (positely or arganythese) by hormonal majoritation. Analogans and estogans have opposing acfass on hormone-responses tames, with estogans supporting the growth of the section source and analogues riskings e⁻⁷. The applementation of an factors are have been applemented on the section of the factors are been applemented on the section of t in women, however, so therapy is usually initiated with great caution. An oral androgen such as flaxoprestronce is usually preferred to a solver acting injectable steroid such as randroken deconnet as well as it can be abruphy hilded funderable side identics become too approver. Both primarily anabolic agents, however, have been widely prescribed for this purpose.

In nexet years the dovelopment of newer and more targeted anti-estrogenic danges selective entropm correspondent to the selection of the selec

Decreased Fibrinolytic Activity

Anabolic storoids may be prescribed to truet confittone susceitated with decreased fibrinoistic active, Frenziopies in the process in which a blood chies in borden down and methoducated by far book. It represents a counter to blood congulation, with the two systems methoducated by far book. It represents a counter to blood congulation, with the two systems methoducated by the book in the system of the system. The system of the deprecongulatively, transmitting the system of the system of the system system of the system of the system of the system of the encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system of the encounter, allowing a similar the specific barries can be seen when the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system encounter of the system of the system of the system of the system encounter of the system of the system of the system of the system encounter of the system encounter of the system encounter of the system of the system of the system of the system. System encounter of the system o

Infertility (Male)

In a small percentage of cases, anabolic/andoprist storeds may be prescribed for the treatment of mike hirding. When the cases of inferibly is to specific momentum of the Logic effect section of disferences, an anabogan might be able to allocitate the condition. In such cases the starod may increase the gene moort, speem quality and the fractase concentration,¹⁷ ¹⁷ which can increase the clarace of conception. The oral anabogan measuring (Provisor) is maint commonly presented for this papers, although has no theory measuring (Provisor) is maint commonly presented for this papers, although has no theory measuring the provisor is maint commonly presented for this papers, although and the fractase is intend.

Growth Failure

In the United States, oxandrolone is the anabolic steroid most widely prescribed for the treatment of growth faiture. It is usually given as a supportive medication, used to augment the anabolic effects of human growth hormone therapy. The drug is typically taken for rewrich of 6.127 morths at a time in an offset to accelerate the anather useful and the state useful and anabolic effects. protection or yet, interest in an events the accessing the growth mass means assistantially affecting the ratio of opphysic strains. A dosign of 25 angle of days is during different process, although this may be adjusted upwards or downwards depending on the patient's sex, gas, belowing at a strain strain to advance different bibles model and conditions, the result may be an enhancement of the growth rate and an increase in total conditions, the result may be an enhancement of the growth rate and an increase in total consistently includes and analysis. There are a strain the strain of the growth rate and an increase in total consistently includes and the strain the strain the other of thirds with considering the includes the mice flat to produce a statistically significant effect on stud height questioning in altimute tables.⁷⁶ The down-termborefield routing is strained on the growth rate, however, the running water produces the strain of the growth rate, however, the major that the strain tables are benefits and the growth rate however the running water produces the strain of the growth rate however the running water produces the strain of the growth rate however the running water produces the strain of the growth rate however the running water produces the strain of the strain of the growth rate however the running water produces the strain of the strain of the growth rate however the running water produces the strain of the str

Libido (Female)

The steroid methylactosterone is approved for prescription sule in the United States and many other markets to improve likelis in framel mercopaus patients. Strail doess of the drug are typically included in products that also supplement estrogens, the combination aimed at trusting the fill spectrum of mercopause symptoms, including reduced frameli likels. The dosage used is low compared to those of other clinical applications for methylastosterone, and will susail warmout to no more than Zang per day.

In the United Status, however, the use of an anabole steroid such as markoken decarates for the direct treatment of conceptories in presently wised as concretual. In pipe of substatuit clickal data and history supporting the use of avoids, for this parpose, in the storid term of the status of the storid presently observed as concepts and the storid term of the storid present of the benefits with conceptorized here in stores of the present present clickal store and the store of the dataset, are for supporting how mixed alongly. Anabolic attends remain use for paparities present present and the store of the sto

Osteoporosis

Anabolis strends increase bote mineral density, and may be prescribed for the treatment of ostoporosis. Benefit of therapy riskels for similation of new bote formation, ribbitom to the recording the standard stream and the stream in postmeropausal wortes, and is usually field of the changes in hormonal channels and the stream and the

Nandokee decanote is the ambodi stroid most commody prescribed for the treatment of ostopronsis. The digit tanks to differ measures the bending with regulated to been densing, and may reduce the Bedibodi of bose finates in patients^{10, 01}. The dosage used to treat postmerpostal bounds in usually 50 mg one every 3.0 4 weeks. Adverse reactions are common with thrapp, however, tacking withinton symptoms (hourseess and bodyfield it initiation) and the distribution of the strong of the strong of the strong before its distribution of the strong of 55, who is a group seem to notice lower adverse effects. Make patients are given a methodowa decanota dosigo of 50 mg ocuoring has for one years or longs if necessary. The leng therpositie withow is standy segurid in order to give the disc cound firm to measurable velicity of the strong the strong the strong the strong the strong strong the strong therpositie withow is standy segurid.

Turner's and Klinefelter's Syndrome

Ambolicatingganic steroids may be used to treat certain agencie confiners, most commoly Turn's yandrom in finants and Kinferler's synthemis in males. Boh an erhornsonenti disorders characterized by deviations from the neural XXXV pring; They result in furning other health issue) abrommilies in growth, secand development, and ongoing secand functioning. Males with Kinefler's syntheme are sterile, and typicably have a moder (less mensue) physica. They also develops malt esticistic, introverbidism, and may safer with geneconstain. In these patients the supplementation of losstoatrons (in a scaling factoris to the store for submotion sections). strum astructure to una treez as attruggin representent transprip to contrasts, and can texp abovitos some of the sisses with sexual fractioning and body composition. Fermaliss with Timer's spathence will be of short stature, and develop other physical abnormalities including a broad check, buy misring, how-rest ens, and webbed meet. Low does of a primurby anabole steroid muy be used in adolescert patterns as an adjurct to growth homome herapy to support the inare growth nucl. Courdoose of the stored more commonly used in the United Status for this purpose, and has been clinically accessful at the university full status of the store of the stored more status of the stored more store of the university full heraft when used in disease of Col. Tantgener drawt.¹⁴

Weight Loss/Muscle Wasting

Anabole stretchism have be administered for the treatment of chinally significant weight loss. Common causes include produped controls of theory, extensions surgery, chronic infections or severe traum. In a garenal sense, these agents can be highly useful when a patient is subject to a top hospital at you expired of bed nets, who menuit day muscle stratifications in project and a significant loss of match times is noticed. Severe them injuries may also call for the supportion-application of baseline strations, and its a type of injury and sense that which applies the strategies of the strategies and the strategies and the strategies and sense that which applies the strategies and the strategies and the strategies and sense the strategies and the strategies and the strategies and the strategies and strategies and the strategies and the strategies and the strategies and strategies and the strategies and the strategies and the strategies and strategies and the strategies and the strategies and the strategies and strategies and the strategies and the strategies and the strategies and strategies and the strategies and the strategies and the strategies and strategies and strategies and strategies and strategies and the strategies and str

The significant toss of lean body muss can present to own set of heath issues. Individual that are chronically undersight may differ from two energy and a readed sense of welkness, and are at greater risk of morting¹⁰. Sensere weight loss during recovery from surgary or fluess may also measured heady or comparison that more assessed and the sense problem to the sense that the sense problem sense that head sense that the sense problem sense problem senses problem senses problem senses problem senses problem senses problems sense problem senses that from sense problem senses problem senses that the sense problem sense problem senses that the sense problem senses problem sense sense many three these senses that the sense sense many three senses the sense sense many three senses that the sense sense many three senses that the sense sense many three senses three senses that the sense sense many three senses the sense senses that the sense sense many three senses the sense senses the sense sense many three senses the sense senses the sense many three senses the sense many three se

In the Linde States, coundoknes in the agart most frequently preserved for most links of checking significant weight low. The dosign used for the papers or specifically ong given weights. The dosign used to the present of the state of the

Anabolis strends may also be prescribed to truet mere sevene cause of macel wassing. This is a confinied metanetical by strong ongring proteinci anabolism, which menss that marked proteins its is early medanismently broken advorm (as opposed to to synthesized) in the hody, and a progressive loss of weight, strength, and energy is noticed. In a model a string, severe marked wassing is referred to as cachenia. Cachenia is not modeled attributed (decary malmention), but the as a function cachenia cannot be advorted with instificant field intradic (decary malmention), but the as a function can advorted and the baseling of the strength of the cachenia like (decary malmention), but the as a function can advorted and the advorted and the cachenia like strength in the constraint size of this constant is and write an advorted attraction that which markets in true of these dings as approvide heavy for carrier cachenia has not been well established, however, and carrently the subject of organiz instrainty.

Nandolove decanotic, onandolove, and osynetholove have been the amboles steroids most commonly used in the U.S. to term tanck wasting specifically associated with HV infection. Although no specific FDA recommendations have been adopted, stadies with nandolove decanous three shown a doage of 150 mg every 14 days to have a similar ambolic beendt, and a significantly lower incidence of side effects, as 6 mg (18 JU) of Tumma growth homese per day. ¹⁶ ID 2003, osymptotice was the side of accessible Passe III cisical trinks for HIV susting.²⁶ The dosage of this study (100-150 mg per day) mitrors those that any one convender survey-hold to useducine. In november tame homesen the usise that are most commonly prescribed by physicans. In recent years, nowever, the discontinuance of nandrolone decanoate on the pharmaceutical market and a perceived higher patient comfort profile in oxandrolone has made oxandrolone the preferred agent for HIV cachexia. The dosage of oxandrolone used may range from 20 mg to 80 mg ner day.

The most consistent clinical benefits have been seen with a 40 mg and 80 mg daily dose 89

50. Are related testosterone level changes and male andronause syndrome. Wu CY, Yu TJ, Chen MJ, Chang Gung Med J. 2000 Jun:23(6):348-53.

51. Osteonorosis in male hypogenadism: responses to androgen substitution differ among men with primary and secondary hynogenatism Schubert M. Ballmann C et al. Horm Res. 2003;60(1):21.8

52. Effect of testosterone replacement therapy on lipids and lipoproteins in hypogonadal and elderly men. Zaliczynski S. Ossowski M et al. Atherosclerosis 1996 Mar:121(1):35-43.

53. Testosterone and other anabolic steroids as cardiovascular druas. Shaprio J. Christiana J et al. Am J Ther 1999 May 26(3): 167-74

54. Androzen deficiency as a predictor of metabolic syndrome in aging merc an opportunity for intervention? Kanoor D, Jones TH. Drugs Aging

2008/25/51/357.(9)

 The effect of testosterone replacement on endogenous inflammatory cytokines and lipid profiles in hypogenadal men, Malkin CJ, Push PJ et al, J Clin Endocrinol Metab. 2004 Jul:89(7):3313-8.

56. Adverse events associated with testosterone administration. Basaria S, Coviello A, Travison T, et al. NEJM July 8 2010 Vol. 363 N 2

⁵⁷. Vascular reactivity in hypogonadal men is reduced by androgen substitution. Zitzmann M, Brune M, Nieschlag E J Clin Endocrinol Metab. 2002 Nov37111:5030-7.

58. Testosterone treatment in hypogonadal men: prostate-specific antigen level and risk of prostate cancer. Guay AT Perez IB Fitaihi WA Vereb M Endoor Pract 2000 Mar, Amr6(2):132,8

^{59.} Prostate volume and growth in testosterone-substituted hypogonadal men are dependent on the CAG repeat nolymorphism of the androarn recentor sense a longitudinal pharmacogenetic study. Zitzmann M.Depenbusch M.Gromoll J. Nieschlag E. J Clin Endocrinol Metab. 2003 May:88(5):2049-54.

60. Obstructive sleep annea syndrome induced by testosterone administration. Sandblom RE, Matsumoto AM, Schorpe RB, Lee KA, Giblin FC, Brenner WJ, Pierson DJ, N Engl J Med, 1983 Mar 3:308(9):508-10

61. Testosterone therapy and obstructive sleep apnear is there a real connection? Hanafy HM. J Sex Med. 2007 Sm:4(5):1241.6 Each 2007 Jul 21

62. Intramuscular testosterone treatment in elderly men: evidence of memory decline and altered brain function. Maki PM, Ernst M et al. J Clin Endocrinol Metab. 2007 Nov:92(11):4107-14. Epub 2007 Aug 28.

63. Exogenous testosterone alone or with finasteride does not improve measurements of cognition in healthy older resugnous resolutione acte or with intesterice does not improve measurements of cognition in relating ode men with low serum restouterone. Visiohan C. Golderein FC. Tenower II. 1 Androl. 2007 Nov. Dev 28(6):875-87. Foub 2007 Jul 3

64. Testosterone improves spatial memory in men with Alzheimer disease and mild cognitive impairment. Cherrier MM, Matsumoto AM et al. Neurology,

2005 Jun 28:64(12):2063-8.

65. Characterization of Verbal and Spatial Memory Changes from Moderate to Supraphysiological increases in Symm Testosterone in Healthy Older Men. M Cherrier et al. Psychoneumendorrinology 2007 Jan 32(1): 72-79

66. Danazol and stanozolol in long-term prophylactic treatment of hereditary angioedema. Agostoni A, Cicardi M. J Alleray Clin Immunol, 1980 Jarc65(1):75-9.

67. Endocrine and intracrine sources of androgens in women: inhibition of breast cancer and other roles of androgens and their precursor

68. Effects of androgens on haemostasis. Winkler UH. Maturitas. 1996 Jul;24(3):147-55.

69. Anabolic steroids and fibrinolysis. Lowe GD. Wen Med Wochenschr. 1993;143(14-15):383-5.

70. Plasma fibrinolytic activity following oral anabolic steroid therapy. Walker ID, Davidson JF. Thromb Diath Haemonth. 1975 Sep 30;34(1):236-45.

71. The effect of mesterolone on sperm count, on serum follicle stimulating hormone, luteinizing hormone, plasma testosterone and outcome in idionathic oliaospermic men. Varua TR, Patel RH, Int J Gynaecol Obstet, 1988

72. Mesterolone treatment of patients with pathospermia. Szöllösi J, Falkay GY, Sas M. Int Urol Nephrol.

73. Osandrolone therapy in constitutionally delayed growth and puberty. Bio-Technology General Corporation Cooperative Study Group. Wilson DM, McCauley E, Brown DR, Dadley R, Pediatrics, 1995 Dec;96(6):1085-100.

74. Growth and srowth hormone responses to oundrolone in boys with constitutional delay of growth and puberty (CDGP). Clayton PE, Shalet SM, Price DA, Addison GM. Clin Endoerinol (Oxf.). 1988 Aug29(2):123-30.

75. Osandrolone in constitutional delay of growth: analysis of the growth patterns up to final stature. Bassi F, Neri AS, Cheri RG, Cheli D, Serio M, J Endocrinol Invest, 1993 Feb:16(2):133-7.

76. Oxandrolone treatment of constitutional short stature in boys during adolescence: effect on linear growth, bone age, pubic hair, and testicular development. Marti-Henneberg C, Niirianen AK, Rappaport R. J Pediatr. 1975

77. Anabolic steroids in postmenopausal osteoporosis. Need AG et al. Wien Med Wochenschr. 1993;143(14-15/392.5

78. Nandrolone deamoate: pharmacological properties and therapeutic use in osteoporosis. Geusens P. Clin Rheumatol, 1995 Sep:14 Suppl 3:32-9.

^{79.} Nandrolone decanoate: pharmacological properties and therapeutic use in osteoporosis. Geasens P. Clin Rheumatol. 1995 Sep;14 Suppl 3:32-9.

80. Nandrolone decanoate for men with osteoporosis. Hamdy RC, Moore SW, Whalen KE, Landy C. Am J Ther. 1998 Mar:5(2):89-95.

81. Effects of nandrolone decanoate on bone mass in established osteoporosis. Passeri M, Pedrazzoni M, Pioli G, Butturini L, Ruys AH, Cortennad MG. Maturitas. 1993 Nov;17(3):211-9.

82 Nandrolone decanoate: pharmacological properties and therapeutic use in osteoporosis. Geusens P. Clin Rheamatol. 1995 Sep;14 Suppl 3:32-9.

83. Effects of nandrolone decanoate (Deca-Darabelin) on serum Lp(a), lipids and lipoproteins in women with postmenopausal osteoporosis. Lippi G, Guidi G, Ruzzenente O, Binga V, Adami S. Saand J Clin Lab Invest. 1997 Oct570(6):207-11.

84. Tumer's syndrome. Guarneri MP, Abusrewil SA et al. J Pediatr Endocrinol Metab. 2001 Jul;14 Suppl 2:959-65.

⁸⁵ Undeweight, overweight and obesity as risk factors for mortality and hospitalization. Gunilla Ringhick Weitoft. Sandinavia Journal of Public Health, Vol. 36, No. 2, 169-176 (2008)

86. Catabolic illness. Strategies for enhancing recovery. Wilmore DW. N Engl J Med. 1991 Sep 5;325(10):695-702.

87. A randomized, placebo-controlled trial of nandrolone decanoate in human immanodeficiency virus-infected men with mid to modenate weight loss with recombinant human growth hormore as active reference treatment. Sorer TW,Woordnesse L et ed. J Clin Endocrinol Metals. 2005 Aug/98(9):474-82. Epide 2005 May 24.

⁸⁸. Double-blind, randomized, placebo-controlled phase III trial of oxymetholone for the treatment of HIV wasting. Hengip UR, Stocks K et al. AIDS 2003 Mar 28;17(5):699-710.

 Osandrolone in the treatment of HIV-associated weight loss in men: a randomized, double-blind, placebocontrolled study. Grandeld C, Kotler DP et al. J Acquir Immune Defis Syndr. 2006 Mar;41(3):304-14.

Steroid Side Effects

While antiokicant/ogenic storaids (AAS) are generally equated as therapeutic drugs with light softs, their use can be be sociated with an nutrier of advances constrict, physical and psychological differs. Many of these side diffs are color apparent during frampeuticand psychological differs, Many of these side diffs are color apparent during frampeuticsequences of the storage structure and the storage short differs and the physical set of the storage structure and the storage short discover differs of storage and not noise and checks that its the matches analytical fields to the side differs of storage and not noise and differs and the storage short discover differs are proper storage and performed existing and provide the storage short differs are observed fragmentation and the storage differs and the storage short differs are observed in the storage short differs and the storage storage short differs are observed in the storage storage short differs and the storage storage storage storage storage and not noise and differs that it is a first and the storage storage storage and provide an existence discust differs are not storage and provide and the storage storage storage and provide an existence discust differs and the storage storage storage storage and provide an existence discust differs and the storage storage storage storage and provide an existence discust differs and the storage storag

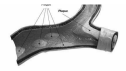
Cardiovascular System

The use of ambokizmloganic storiks in significancymic (and offen theraportic) doess can hav a nuther of advere effects on the cardiovacient system. This may be noticed in scenal areas inkiding unforceable alterations in sourch relations; in this may to noticed in source and an experimental storing in source areas the source areas these drugs are admitted) very safe. The risk of an otherwise healty parson saffraring a hard stark from an isolation storid y exist in scenary mercine. The risk of moties is also etterned by an example of the storid storid storid storid storid storid storid low When these drugs are abased for long parisals, however, their adverse effects on the bard match storid storid storid storid storid storid storid storid by the storid storid storid storid storid storid storid abase of moties is also etterned by the storid storid storid storid storid storid storids and the formed to befort match storid system storids was been available in moties at storid storids that cardiovary storids in scenar by wass.

Cholesterol/Lipids

Anabolicintralinguistic steroids use can adversely affect from HDL (good) and LDL (good) dedisteroid values. The ratio of HDL to LDL (good) dedisteroid values. There and o HDL to LDL (good) dedisteroid values. The ratio of HDL to LDL (good) dedisteroid values. The ratio of HDL to LDL (good) dedisteroid values are dedisteroid values of the origination of plaques in the attrice, generating afterograms or article concentration, which is done contributed with table or a reasonal OL Locksh: Tiggebord values are dedined by the transformed of the origination of the component of the plane is a structure of the structure of the origination of the component of the plane is a structure of the structure of the





Over time, plaque deposits may begin to narrow and clog arteries.

Anabelexitologuise stensits are most consister in the Twening of HIL level. This abserve effect is molical drough the analogues is minimized or Diparties lipses, a here compose responsible for the breakdown of HIL (good) choiseston¹⁰. With more hepatic lipses activity in the hody, the borehub (and-shengers) HIL choiseston with a prime size a characterization more quickly, and their break dops. This is an effect that scene to be very isostances coprime tool at 21% dops in HIL choiseston with a doogs of 700m gree weeks¹⁰. Encreasing this dosage to 600 mg did not here any significant additional effects againstig that the doogs threakly distance HIL supervision in hity how.

Oral stretch, especially ~ 17 apha adylated corporatis, are particularly poter at straining heard incease and appressing BLP. Levels. This is the tot for pass concentration and metabolism in the ber. A drug face stancoid music, therefore, he mikke than a stretch and the tot optical stretch and the stretch and the stretch and the stretch and to top is of migra words? More migra words, for any stretch and the stretch transmission. Exact distribution at the stretch and the

It is also inportant to note that ensogens can have a foureable impact on choisensol profiles. The anomization of testostonene to sensitival of my, harefore, prevert a more dimutis change in secure choisesten A and and caurined this office by comparing the light changes cancel by 20m erg. We also another on the provide set of the anomization of the comparison of the comparison of the comparison of the comparison subscience cauting the distribution of the comparison of the comparison choisense, the group many distribution of the comparison of the comparison of the comparison of the exhestion of an average of 25%. The group taking methylaconstone experimental models and the comparison of the comparison of the comparison of the comparison of the exhestion of an average of 25%. The group taking methylaconstone experimental models and the comparison of the comparison of the comparison of the comparison of the exhestion of an average of 25%. The group taking methylaconstone experimental models and the comparison of the comparison of the comparison of the comparison of the extended numbers of the result. The spectra the comparison of the comparison of the extended numbers of the result of the comparison of the comparison of the comparison of the extended numbers of the result. The spectra the comparison of the comparison of the extended numbers of the result. The spectra the comparison of the compari

The potential positive effect of entrogan on chelstenoi values aito mukes the issue of enorgon miniemeures constribut to consider whet is conso to health risks. To beguit whet, one may want to consider whether or not entrogan miniemeure dungs are actually necessary in any given circumstate. An wisk effects apparent, or is their use a proventine stogan and perhaps surcessary? The miniemeure dung of choice can also huwe a messarable impact on choisened outcomessaries and the storage of the storage storage of the storage net score to eight anti-samparies (effects on choisened values, and in fact tasks to increase size effects instant of momentate inhibitor. The instance, particularly when they are target storoik in for larger periods of time, and are concerned about their candidate candidatescales

Enlarged Heart

The humn heart is a muck. It processes functional androgen receptors, and is growthresponse to musk second hormores. This deterphy accounts for methying larger heart muss can arcoge fram worms.¹⁰ Physical activity can also have a storage effect on the growth of the heart. Resistence exercise (narrowsb) (not its to iterase heart is by a thicking) of the ventricity and usally without an equal expansion of the iteranal carvity. This is known as concernive remodified, Endance (androb) (and these, on the other heart, lead to iteranal least size we expansion of the iteranal carvity, without significant thicking of the ventricles locality preminimizing in the differ listen. The other size heart does not complete the structure of the iteranal carvity is the size of the structure is also complete the structure of the iteranal carvity is the structure is also does not complete the structure of the iteranal carvity is the structure is also complete the structure of the iteranal carvity is the structure is also complete the structure of the iteranal carvity is also also also also also also also complete the structure of the iteranal carvity is the structure is also also also complete the structure of the iteranal carvity is also also also also also complete the structure of the iteranal carvity is also also also also also complete the structure of the iteranal carvity is also also also also also also complete the structure of the iteranal carvity is also also also also also complete the structure of the iteranal carvity is also also also also also complete the structure of the structure of the iteranal carvity expansion tend to reduce.

Anabolis strend abuses are at risk for thickneing of the left and right vertricitient thrown as vertricitical posterophy. Hypertrophy of the left vertricitie (the main parange charaber) particular is cotansidely documented in anabolicant/organic stored abuses.⁷⁷ While lett vertricitient posterophysis, again, also fourt in manal power athless, abstratuabusing athletes tand to have a meet more produced and thickneing. They also letted to develop publicaging answers thirds to find theoring, including paranged inside fractions, and athinably valued heart efficiency.⁷⁷ The level of magnetize the closely associated with men matrixed, and may be histokine of stored accordiance of the curves.⁷⁸ The top developed suggest the publicagial left vertricitar hypertrophy has developed. Additional testing of such patients in the commended.

Let vertricht hypertophy (LVI) is an independer precisive of metally in overweight mitischaus with high-boff personse.¹⁰⁰ II has also been indext of artifications, vertricht affendins, vertricht affendins, vertricht die high-boff, and hubbegal intenses in (17) dispersion are notation in this is be without chait align-factore, publicagie intenses in (27) dispersion are notation in the second dispersion notation in hypertensive patients with 10×10^{11} M Aroung other fings, this could have a second abseing principation with 10×10^{11} M Aroung other fings, this could have as second abseing principation with 10×10^{11} M Aroung other fings, this could have as second abseing principation with 10×10^{11} M Aroung other fings, this could have association between UH and related probability in could have association theorem of probability in out-find and the fingers of the second second second second second second second second second districts (10) and the second second

Heat muse can increase or decrease in relation to the current state of anabolic/strond use, the average dossag, and dominion of finke L. Decressi, the heat sussay hybegies to relace a size once anabolicandrogenic stronds are no longer heigu used. This effect is similar to the way the heat of affects in size once an and there to toger flowlow a rapproxe imilarity schedula.¹⁰ Decrewith this effect, however, some changes in heart musel, size and function caused by the singer present. Static scenaring the effects of search and a size of the state strend in the state of the state of the state and without on left verticale hyperturphy north and latel size along the state strend in the state strend in the state of the state strend in the shifts and state strend in the patholigiest left verticaler hyperturphs following large-strend strends and the matching left strends the subject of present strend strends and the state strends left of the strends where the shifts of strends and the strends and the strends matching left verticaler hyperturphs following large-strend strends and the shifts corresting the solitor of present strend and beats.

Heart Muscle Damage

Ambléindrugnis storid abuse is supperted of producing direct damage to the heart mack in some cases. Studies exposing hard red coluturs to ASA have resported reduced contractile activity, increased cell fugility, and reduced cellular (introchondrina) activity, providing some support for a possible direct twice effect to the heart muckel.¹⁰⁷⁰ Furthermore, a number of case reports have found such puthologies as myocantial fluoriss iccer tasses bulking in the heart, myocardini information of heart insect.¹⁰¹⁰ (action fluora thosis) using terms strong disarders.¹⁰¹⁰ Intil 126 deert link between disard interests (dation floart insiss) in long-terms strong disarders.¹⁰¹⁰ Intil 126 deert link between dige abuse and cardies (antibologies is assamed in these cases, the carned be proven given the skow mange in side these cardies (antibologies devity, and the intelance rung other factors). (such as unet, exercise, messive, and genetics) can have on ment, monotonia remain caunoned about the possibility of cardiac muscle damage with long-term steroid abuse.

Blood Pressure

Anabolicanloganis steroids may elevate biod pressure. Studies of bodybulkent nårgi these drugs in spratformegneti. does how demenstrated neurosais i hosh spatia dara disatois biodo gressare readings¹¹.¹¹ Anaher study manared the average blood pressare randing in a good of studies to be biodyfold, subha so compared to 15200 in vegel filting, studies and studies and the study manared the average blood pressare and analysis of the studies are studied with the studies of the studies allocation of the studies are studied with the studies of the studies allocation of the studies are studied with the studies of the studies of the cased by a narber of factors, rachading trevesoid vater retorion, neurosci vascidar diffuses, and areneas beneficies. A more models, functional blood pressare mp be aused by a narber of factors, rachading trevesoid vater retorion, neurosci vascidar diffuses, and areneas one blood pressare, allocaji devatoris carnot be excluded with antibicarily devater studies have beneficiered from the studies of the s

Hematological (Blood Clotting)

Aubcländingsnis stendis can case a nurber of changes in the benutokegial system that affect blocd changes. The therapeoint is coll that affect blocd changes in the therapeoint is coll these drugs is known to ircrease phornin, antithrombin III, and protein's I beek, simulat fibritopies (else throadours), and appropses a drugs gatelines for anabolicantogarie strendistanti and the strendistanti and the strendistanti and the strendistantogarie strendistanti and the strendistanti and the strendistanti and the strendistanti strendistanti and the strendistanti and the strendistanti and the strendistanti and strendistanti and the strendistanti and the strendistanti and the strendistanti strendistanti and the strendistanti and the strendistanti and the strendistanti significance to healthy individuals using these drugs in the anteropartic drugs the strendist strendistanti and the strengest individual to the strengest different their antronaganter, followed interest, however, could be adversely different brief and strengest and the strengest individual to their the strengest and the strengest different brief and strengest and the strengest individual to their the strengest and the strengest different brief and strengest and the strengest individual to their the strengest and the strengest difference brief and the strengest difference brief and strengest and the strengest and the strengest difference brief and the strengest difference br

Concressly, anaboli:integrates strond abuse has been bisked to increases in bioled cheffing ability. These drugs are needuce these of fromton¹¹ and C execution protection.²² are set also framehouse A2 receptor density,¹²¹ which can support plackit aggregation and the formation of blood CAS Statis of stretch while were have elementate adstatically significant recreases in plackit aggregation tusks in some subject.¹²² There are also a growing number of cases neprot where pointerings fully blood cless, embodies, mat addies, have obscured in standi abuses,¹²³ Life Life Life Albusgh 1 has been difficult to conclusively fulls the second fuel data. (In advectes effects of anabolis extension conceptors ref or the obscure divised abuse, the advectes enviros and avece diffects are now regarded as recognition rises of advect data.

In therapeutic levels, the anti-thrombic effects of anabolic/androgenic steroids seem to dominate physiology and decreases in blood clotting ability may be noted. At a certain supratherapeutic dosage point, however, the pro-thrombic changes appear to overtake the anti-thrombic changes, and physiology begins to favor fast and abnormally thick clot formation (hypercoagulability). The exact dosage threshold or conditions required to increase blood clotting has not been determined and some studies with steroid users taking supraphysiological doses fail to demonstrate increased coasulability 128 Individuals remain warned of the potential increases in thrombic risk with anabolic/androgenic steroid abuse. Blood clotting tendency should return to the protreated state after the discontinuance of anabolic/androgenic steroids, point until the hematocrit issues have been corrected. Minor elevations in hematocrit may be addressed with phlebotomy. For this, 1 pint of blood may be removed periodically during steroid intake, often every two months. Proper hydration is also important, as dehydration can temporarily cause the hematocrit level to elevate, giving a false positive for polycythemia. The daily intake of aspirin is also commonly advised if the hematocrit is above normal, as this will reduce platelet aggregation, or the tendency for platelets to stick together and form clots. Individuals remain cautioned of the potential cardiovascular danger of high hematocrit levels associated with anabolic/androgenic steroid 1162

Hematological (Polycythemia)

constrained access effects of this is polyclycation, as the overpreduction of nel block-off. Polyclycanic can be reflected in the beamscere level, or the percentage of block beams that Polyclycanic can be reflected in the beamscere level, or the percentage of block beams that block because to be the beamscere level. The beams the beams that the beams the beams that the beams the beams the beams that the beams the beams the beams that the beams the beam term of the beam terms that the beam is a beam term of the beam terms that the beams the beam terms that the beam term of the beam terms that the beam term of the beam terms that the beam terms that the beam term of the beam terms the beam term of the beam terms that the beam terms that the beam terms term of the beam terms terms that the beam terms term of the beam terms terms terms that the beam terms term of the beam terms ter

Anabolisticatul admissination toncis to mise the lementer't level by several proceedings points, sometries more As a reads, many selecit-sings holdphaless well have lementer levels that are above the normal range. For example, one study measured the average herearchicatul argued constrained and compared to hold because that the herearchicatul argued compared hold because more herearchicatul argued and services candidouscellar event. Although net likely to be an include classes, high hermitter is is believed to have been and on a contribuing factors in the doubt of a matter of servical hauses, manify particul with high block pressure, hereacyceine, and/or athrenetic ratios and early and within the remain means for health and attractions and early and within the remain means for health and attractions and early and and the service and the health and attractions and early and attractions and early and attractions and a service and a service and a service and and a set of the service and particle of health and attractions and early and attractions are early and attractions and attractions and a service and attractions and attractions and early and attractions and attractions and attractions are early attractions and attractions and early and attractions are attractions and attractions and attractions are early attractions and attractions and early and attractions are attractions after a service attractions and attractions and attractions and attractions and attractions are attractions and attractions and attractions are attractions and attractions are attractions are attractions and attractions and attractions are attractions and attractions are attractions are attractions are attractions and attractions are attracting and attractions are attractin

Many physicains that specialies in hornexe replecement theory consider a hornexter lived of 55% to be an absoluce cardf point. At or above this point, and anabolicantlogonis storial througy cannot be continued addy. Dang sindle would be caused at this point and the humitori sites such been corrected. Whit more elastions in humitoric may be addressed with plathotomy. For this, 1 per of blood may be removed periodically damg staroli strained, othen every to morths. Proper plathotins is also inprotein, as deductions can temporarily cause the humitact head to destate, gaing all data possible for plotyphisms. The moves plathot aggregation, or the indexity of plathotis to addite and the mathematic tasks Indexidant commits competing humitant of the plathotis to addite and mort and indexing the strainest of the posterial environment datages of high hematorit levels movined with ambodicandrearies stream of use.

Homocysteine

Anabolicantingganis staroids may elevate home-spaties levels. Home-spaties is an intermediary aniso and produced as a hypotent of rendening remeables mHg helveds to Trans the results of the end/or spatial star starting and the end/or spatial starting and the spatial sta

Androgens straintic elevations in home-costen,¹²⁷ and men have an approximately 25% higher beel on research and bandbeinandrogenis straintic and have have been associated with hyperhome-systements, or consister distably high home-system beek.¹²⁹ Do estudy faurd that here average home-systemic eccentration in a good O I on that hash been additionating antiokalandogenis storolds (in a local straintic distability) and the same and home-system bandbein and been additionation of the same and do rate transt and dramp and O I on that hash body balances of the same match data in the straintic distability of the same and do rate transt and dramp data in the home-system is a straintic of the same straintic distability of the same straintic distabili

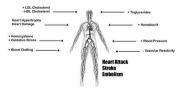
Vascular Reactivity

The endothemins is layer of each that lise the errise circulatory system. These cells are found on the misted of allbod vessels, and help increases or decremes bolds on and pressure by relaying or constricting the vessels (referred to as vascellation and vascesconstriction, respectively). These each has been pregate the pressure of materials in and out of bood vessels, and are involved in a number of important vascellar processes inkeling blood graenally consistent of a start of the presses of materials in and out of bood graenally consistent of denihel for health and. Blooksie, the endothetial solared comported in individuals with endothetial docated for the conference in other and to a notice generative blood of was, helper body presses, bealt influmming, and reduced circulatory capacits, ¹⁴². This may place them at greater risk for hear tanks, stroke, or formboois (blood 40.).

Endoblial cells are autoigns response, which may partly account for men ethibing less uscalar reactivity many usons.¹¹S Strahlar, anabolicantingenis etsendi sen histon shown to impair endohelial activity and suscalar reactivity. Statias at the University of Imbernek in Adamis compared the sele of endohelial altimist. Altivity and the selection of attaces.¹¹⁴ Those influential assignations and the selection of the adamistration of the sense of the selection of the selection

Proving an Association

Direct first between strend above and individual cases of strake and heart anack have been difficulto 1 prove. There are a narrier of lings final have more this difficult. For our, cardiovascult disease is very common in must fiado standy takes decades to develop. This mises individual contributing takes (which have have may strate and a soci, tilledy), having the strate of the strategiest of the strategiest of the strategiest of the strategiest methanism of the strategiest of the strategiest of the strategiest of the strategiest strategiest of the strategiest of the strategiest of the strategiest of the strategiest strategiest of the strategiest of the strategiest of the strategiest of the strategiest strategiest of the strategiest of the strategiest of the strategiest of the strategiest strategiest of the strategiest of the strategiest of the strategiest of the strategiest body of the strategiest body of the strategiest of the s



Anabolic/androgenic steroid abuse can produce changes in a number of areas of cardiovascular health that can work together to increase the risk of heart attack, stroke, or embolism.

Immune System

The humm immure system is responsive to see hormers. This results in fractional differences in immurity between the sector. Worms tend to have a more active immure system compared to mere, and are slightly more resistant to bacterial indication and other types of relative 10^{-10} the minimum system and as a horme prome to developing anothrmure diseases, which may be liked to be lightly beel of activity¹⁻¹⁰. The days-to-day activity of the immure system can also harten for truggooth the messatial cycle, thirde demonstrating the demonstrating the demonstrating the demonstrating the leader to be caused by storatores, which is an immurospression hormers. ¹⁰Andongers may modulate the immure system afreedy, through their convexion to entogens, ¹⁵⁰ or by medifying abcoordinal carbin.¹¹

Anabolicardingsmic stervide how displayed hosh immunosimatory and immunosprepose actions in animi moleck¹¹. Gine that the data can indexe we immune systematic tension in animi moleck stervide area fairly disease class of onlyna, their effects on the immune systema mixed molecular and molecular When used thanpointeally, classing in immune system infractionary and molecular and animated in 3 starting immunosimations or immunospression. Anabolic and mogenation and the systematic with mask working susceitated with HIV infection, without any significant change in immune systematic ²¹³⁴

The use of analysis inducing an inducing sensitive provides the sensitive system functioning relationing in infinitality assistance to certain types of infection. In cost study, steroid abuses were shown to have lower sensus levels of [gfs, [gf], and [gfs immunophysical analysis and gradient provides and the sensitive sensit

Kidneys (Renal System)

Anabolicantioganies structiska angenerally well tokrented by the renal system. These drugs are largely occured from the body frough the kickess, adhrugh them is to a interact strong tuckity in this process. In fact, these are many instances in which these drugs may be used as supportive toximum is painties with compression kickess fraction. For example, anabolic strokits have been prescribed to increase the production of red blood cells in patients with agarnel anabole. (Ian body muss) support, and to treat hypograndism, in patients undrapping dialysis.¹¹¹⁰ Wilk accur must be taken with schult patient, theory may density to conducted usy safely. In otherwise healthy tafixidada, clinical renal toxicity caused by the stort-term administration of anabolic characteris storeks is unlikely.

There have been isolated reports of severe kideny damage in stendi abuses. For example, a landfaf ci divisitation has developed Winn Taruc (respherbiskarum)¹⁶¹⁶¹ witch is a rare form of kidney cancer usually fourd in chiken. Its appearance in adult stendi situations evidence that drags were the statal cancer. There have a base usered, hard a conclusive evidence that drags were the statal cancer. There have a base common from of kidney concer, however, conclusive evidence that drags were the statal cancer. The state is the most common frame of kidney concer, however, conclusives, concache is issue angular, base. The have additionally been case reports of coronised law to area in the fast of state the state of the drags with fast of base. There is the state of the drags with the drags of the drags with the drags of the drags with the drags with the drags of the drags with the drags of the drags with the drags of the drags with the drags

The use of anabolic and openic steroids in superalterapencic dosser may slightly impair immune system functioning reducing an individual's resistance to certain types of inflection. In one study, steroid abusers were shown to have lower serum levels of IgG, IgM, and IgA immunoglobulins (antibodies) compared to bodybuilding controls, consistent with immuscipreprission.¹¹² Although this may bigically increase the charace of cortrarring certain types of theses, Klinely health should be a concern for long-time material-sing biodyblacks and power afficies. To begin with eccession resistance raming cam produce some strain on the real system. A conficion calidar fluidading biologis is a smolely the extense damage of market biods.¹²⁶ In high beaches mogelish and a number of neghetoxic composate is to the biods.¹²⁶ In high beaches mogelish and a number of neghetoxic composate with the hards market and the contrast strain biologism is a strain of a strain strain the strain strain strain the strain st

Liver (Hepatic System)

May cell ambediationgenic storiks (or injectable form of cell storiks) are twice to the low (Papottoxics). These composites can user aviors and sometimes likelytoteting during when abused, and occusionally even under therapontic confitons. These agarts commonly associated with disaid leptotetication of the event of the total commonly associated with disaid leptotetication of the event reset of the event of the model and the event of the storiks is an event of the event of the event of the event of the storiks is a subscription of the event of the event of the storiks is a storik present of the event of the event of the storiks is a subscription of the event of the event of the storiks is a storik present of the event of the event of the storiks is a storik present of the event of the event of the storiks is a storik present of the event of the event of the storiks is a storik present of the event of the event of the storiks is a storik present of the event of the event of the storik present of the event of the event of the event of the storik present of the event of the event of the event of the event of the storik present of the event of the event of the event of the event of the storik present of the event of the event of the event of the event of the storik present of the event of the event of the event of the event of the storik present of the event of the event of the event of the event of the storik present of the event of the storik present of the event o

Algoliton of c-17alpia specifically protects the stread melocie from metholosim by the ensyme 17best–hydroxystenid deploydnesses (Tyber-18EO). This ensyme normally oxidizes a stread's 17best–bydroxyl (17best–ol) group, which must remain itself of the drip to input any analysis of the discussion. Whose protection from this ensyme, way tilt action drug will avoid to activation. Whose protection from this ensyme, way will be active the protection the discussion of the discussion of the discussion doing Algoliton of c-17alphu efficiently protects the samed from 17best–1850 by or compared methods and the discussion of the discussion match, the fact hose scores matrix on the two the protects does in the biodistant match, the fact hose scores matrix not the rule not ensore the rule transmission.

To east mechanism of hepatosocky induced by adjusted anabolicandoguis streads remains unknows, but it is specialized to be due in large part to the nuthal activity of androgen in the ber. This bere processes a high concentration of androgen receptors, and is response to these bermans.¹¹ With hypothesia androgens with the instreme and the instrements of the second strength and the second the second strength and the second strength an

Early her toxicity is smally viable in blocd test reads. For hystic function before physical symptoms or dydarfields of doelby. This is not Bady to include eloutions in animaimaticness ensympts. ACT and ALT, also called serum glasmic-notekent: transmission ensymptation of dydarfields and the symptometry of the symptometry of the eloution. A share physical society of the symptometry of the symptometry and the symptometry of the symptometry of the symptometry of the eloution, the symptometry of the symptometry of the symptometry of the dampet from shore in dynamics. Since a symptometry of the symptometry of the dampet from shore in dynamics. Since a symptometry of the s assessment of liver and full-body health is advised should any signs of unacceptable liver toxicity become apparent.

The most common form of actual here dysfurction caused by the administration of cont anadoxiant/cognize attacks is chocknoics; 2007 The describes a contribution where the flow of his becomes decreased, usually because of obstruction of the small bla dates in the here (implayed). This causes is bus also althicitin to accuratia in the here and blood instant of being properly excited through the digastice tract. Influences in the most result of the states of the state of the abdoming plan, or parsing (ching). This stool may also change to a day cold (chickle state) due to the stade cause of the state. The state most results in the state of the morthermore, due to high beeds of bindrain in the blood (hyperbindments), Intraheguia chocknosis may also concluse with hyperchartment for the state of the state morthermore, due to high beeds of bindrain in the blood (hyperbindments), Intraheguia chocknosis may also concluse with hyperchartment because (hard for the state).

Intrabuptic cholestics will usually resolve locif without scritons jigary or medical intervention with sevent weeks of discontinginal glavelatoria strends. More services cases my take sevenal morths bothen normal hopatic enzyme locik and functioning are restored. Hepatic lisions are likely to hal over time as well, a likela partially. It most cases physicians have initiated apportent treatment with unochooxytchic acid (norodat), which is a secondary bile as all known to possess hapatprotective and march-locktient effects, in an effort to have market ensure that and the second second second second second second second paradice remnit subtravo, however. The two is high residert, and intrabuptic cholestistis is utilely to continue degrading after drug discontinuance unless additional pathologies are present.

Mere errors heptic complications are zne, bet huve irkulical peliosi heptin¹⁰⁰ (bloch filed cysts on the low-portal phyrerbension with variaceal blocking" (blocheng caused) bloch pressure in perturb vin related to obstructed bloch flow), heptinechafter adsomm¹⁰⁰ (non-minigur less transch, blochendkar arcsinom¹⁰) "molganit Fei transch, and heptige angioascentor¹⁰⁰ (legensole milgant causer of the imag of bloch vessels and whote claim complexity of the strange of the physical bloch vessels differs have only support. Although many of these portarity bl-feriodering all effects have often been attributed to very a patients receiving stration medication, a graving matter of case represent a new moloning directive leading your all polyhaldsis haltings bodyblatic developing here cancer after taking high doess of oral anaboleindogene strexit, and one confront data,¹⁰¹ ¹⁰²

Physical

Acne

Androgens stimulate the selencous glarks in the skin to screte an objeststature called shown which is mode of fast and the remnosts of datal flavoratice code. Excess stimulation, as with storoid abuse, may also cause a significant increase in the size of the selencous glarks.⁽¹⁾ Selencous glarks are found at the base of the infiles in all latcordining areas of the skin. If the androgen level becomes too high and the selencous glarks become romeworks, the hard fullesa my begin is clag with selenar and data slikes and the selenar to match the selence of the fact of the selence of the selence of the selence of the selence of the fact of the selence of the selence of the fact of the selence of the selence of the fact of the selence of the fact of the selence of the fact of the selence of

A mill reductee of new valgeris is usually addessed with topical over-the-courter area medications and a regress sinc learning routing that removes eases of and diff. More serious acress may develop in semistric infoldanti, itselfang care, couplebang (source acre acress). Such relations may peak predicatil intervision, which sub-product that with intervision. Topical anti-androgen drugs are also under investigation for the transmet of severe acres, and how shown a grant deal of promise in our by mills." Acress in typical resolution that the coststant of several use, although the corepodaction of sebanning peaks and the sedences angles. There that may also all the self-several grants are been appresented and and and the sedences angles. There that miss the mark place the single second series of the self-several grants. There is the place of the self-second second se



Acne on the chest caused by steroid use. Hair Loss (Androgenetic Alopecia)

Aubclichtragenis stendis my contribute to a form of har hose on the scalp known as androgenicia abpect (AGA). This disords its characterized by pospession-miniarization of hair folds), and a shortening of the angang phase of hair growth, under androgen infraters. The hir proceeding which and balles will approximately a single loss and loss effectively in men, the haldness produced is smally distributed more through the start of the start of the start of the start of the start distribution of the start distribution of the start distribution of the start distribution of the start of t



With male AGA, hair loss is most pronounced on the temples and crown.

Antiopartic algorica is the russ common from of that loss in runs and women affact. It is opecially common immess, and never the 30% of the population with molece by the age of 50, ³⁰⁹ As is runs signifies, antioparcit algorica involves the interplay of both antiogaric hormones and guarcit factors. Individual with this condition appear to be more leading sensitive to antiogarse, and howe higher levels of antiogan receptor protein and displorationstorem in the scaph, in comparison to those unitacellities. Although displorationstorem is does not prosent an ange ability to indirect the confloradingentic algorizer, alongent and market in a sensiti are capable of providing horecomparison guarding and antionalities. Unlike the scape is a sensiti are capable of providing horecomparison guarding and antionalities. Unlike constraint lace, comin the absence of steroids that convert to, or are derived from, dirivdrotestosterone.

The genetics of androgenetic abspectia are not fully understand. At one time I was believed this contains could be inherited addy from the maternal grandfluct, More recent evidence contradicts this notion, however, showing strong support for fullew-to-son transmission in murgue cases.¹⁷³ A matter of garses have been uterfailed as having a potential like to be disorder, relating certain variants (polynom/sims) of the androgen receptor gars.¹⁸⁴ How No single genetic variat abate has yet bound be to ophinal actions of androgenetic algocia. However, AGA is now believed to insolve a scenario garses, full-garses,¹⁷⁴ How control the outer and a wearyly or admogratuation allowers. In Singles 2000 and 2000 are strated to another the outer and actively or admogratuation allowers. This regular to allo known to implant the robundrogenetic alteromy activity and the polynetic scenario and the tophican allowers. The strate alter both androgenetic alteromy activity.

Teatmer for androgenetic abspecti in men usubj involves topical mixedial and call finanticle, a 5-shiph inclusive arrayme mibiok. Women are popularly preserved androgene and companyingosisti dugs. The focus in todo causes is on reaking mikine in midi, may started uses a constraint of the started started and the unconstraint of the started started started started started started unconstraint, may started uses concerned with this is so will are first during infect on mission and the started started started started started started started started started unconstraint, and started with varying degress of starcess.

There has been no study on the role of genetics in balances fixed to stored abuse. Accordult, individual web cusing visible and integrated appear paper to be how more susceptible to the effects of ambiokicandogenic storeds on the scalp. For many of these people, the loss of this appears significantly accelerated when taking these drugs. On the other hand, this side effect is generally a much less significant issue with individual further that not needed training before the size of the size without any visible effect at a function of the size of the size of the size without any visible of the size of the distribution. It is unknown, however, if ambiokinnelogueic stored abuse that and the does not carry wardent socreds with the size of the si

Stunted Growth

Anabolic matrix, These horness standy and huse a distributed before physical matrix. These horness standy and huse as dischoration affances on learn high. These hands, their anabolis effects may increase the potentiar of calcium in the boars, follaring effects and the standard standard standard standard standard standard effects and the standard standard standard standard standard effects and the standard standard standard standard standard growth is index standard in plantard standard standar

While andopens, estrogens, and gluccorricoids all therearly participate in home matrix, estrogens in equalito 6 the primary inhibot of home growth in both men and workma²⁷. Werene are aborter on average fram men, and also text to stop growing at a slightly earder age, due to the effects of this hormore. Antohocimalogenia stored that effect covered to estrogen are inherenfly estrogenia and, knowle, more kindly to inhibit famer growth that of the grant. Further models, which are interesting that the growth that of the grant. Further models, which are interesting that the grant of the prime matcholoxes, and oxynetcholex. These drags must be used with additional earliers in young patients due to the strenger potential for addition growth areas.

Estrogen acts directly on the epiphyseal growth plates to inhibit linear growth. These plates are located at the end of growing bones, and contain a collection of stem-like cells called chondrocytes. These cells probleme and differentiate to form now bone cells, showly expanding the length of the bones and the height of the individual. These cells have a finite like set of the state of the locate and state of the individual. span, with programmed senseconce (cell denti). This will cause the rate of chordhocyte profileration to show over time, and eventually stop. The chordrocytes are replaced with blood and hore cells at the point of physical mattarity, "fissing" the bones and iribiting farthere linear growth. The stimulation of estrogen appears to accelerate hore age advancement by exhausting the profilerative potential for chordrocytes at an earlier time.²⁰⁴

Age will also influence a patient's sensitivity to ophylogeal faion. As young abilityme and the final point of boots multity, the influence effects of hommers hearing values (ability the effects. Statistic trending tennylogeal to gate ($S_{\rm effect}$) statistics trends to these effects. Statistic trending tennyloge to gate ($S_{\rm effect}$) statistics trends of testsotarene curulture (S00 mg every two weeks) was afficient to induce final health values the first statistic effect on the statistic effect on

Tendon Injury

Aubclichtregards strend use is sometimes associated with an increased incidence of tunkton injugod¹²⁷. There are a net your trained causes for this, One of the most basis is the mpid growth of the mackets. Under the influence of anabolichtradrogards attorish, for mackets are capable of becoming gindizanity larger and subsequent and the properties of the trained and the annotar of volget Hilded by the started user tracenses, so to does the load placed on the tantion. If the traineds are origing on sought the to (or string) constraints for the stores are ready on the source of the store of the trained store of the store of the store of the stores in my thirtsely lead to the type of righty in some users, essentially by making them to store for their own concredence issues.

There may also be more direct mechanisms combining to this type of pipes Sinula to sketchi mack biose, the behaviors are responsive to analogons. Statiski show that urched is stratefation of analodi, statiski, the kurchen become more right 2^{10} . As tassat behaviors are strategistical to the strategistic of analodi, as the strategistic of analodi, as the strategistic of the strat

The data concerning the potential role of anabole structurk in concretive issue gives a equivale. Seen studies find an association behaves AAS use and tension gives ²¹¹ Overall, these types of spinois are all fairly uncommon. This could much program association with fidew in this type of spinois and all actual denoid and the spinois of the spinois and these types of the spinois and the more advanced body/balakes and atexual absents. On the other land, tendon sign: (voltated a specific acident error) seems to be externely uncommon in frome that use ASS, meader and spinois that one work that the spinois and the spi

Water and Salt Retention

Anabolicindegenic sterolds my increases the amount of vater and sodian stored in the body. This my viscal the increases is not the strancellate and contendant waves comparisons, intrascellate flast refers to state that has been drawn single the call. While its dots and the strange strange strange strange strange strange strange is done calculated and viscales as part of the life fibre body mass. Ensemblier waters is stored in the circulatory system, as well as is suricus holy issues, in the spaces between cold (restrainfi). Increases in iterativitial flast as the probability of the body (peripheral or width) or distributions from the the physics. Excess that iterative mode the body issues, the space width) of marks functions from the theory. associated with elevated blood pressure,212 which can increase cardiovascular and renal strain.

Estopsis a ragidate of field testorio is hoft men and unrenn²¹¹ This effect appears to be medical in a put by charges in hypothatian ragines usequesses (AVP), the primary hermoze insolved in controlling undar real-neighbor in the kickogs.²¹⁴ Increased levels of estopsin turk in zerose AVP levels, which can promote the increased storage of source. Estopsin also appears to a cto the real tables in the kickogs,²¹⁴ Increased storage of source. Estopsin also appears to a cto the real-tables in the kickogs,²¹⁴ Increased storage of source estopsisting and the real-neighbor regulate the source balance of cells. Higher levels con significantly increases under in the estimational resonance in analysis, and allow the storak that either conserts to estopsion or solarise estocylate the atomic real-neighbor and the social of the increased encoders water to estopsion of a social with increased encoders water torakes.

Energenic anabokizuthogenic storols are generally locored for mess gaining (bulking) proposes. A storols user more ignore varies retoricito during this plane of mining coassistantly even fulfing the short necroses in size to be a sucknown breath. Entropying the storage bulking against storage storagest storagest and storage an individual resistance to ignory. With the use of must storage storagest and storagest an individual resistance to ignore storagest for a large storage storagest and storagest an individual resistance to ignore storagest the storage storagest storage storagest and storagest an individual resistance to ignore storagest near the storage storage storagest and storagest storagest storagest storagest storagest and storagest between storagest and storagest storagest storagest storagest storagest storagest storagest and storagest storagest storagest storagest and storage storagest between storagest storagest storagest storagest storagest storagest storagest storagest between storagest stora

No-aromitizing stories such as controlstone and stanooble hrue also been shown to prometic increased waiter elevations, to the field is not entroly excitation to controllable of estingaria substances.^{217,238} Anthonés astroids with low or no entrogenie action tend to produce models increases in which body start and intrachilitar field attention, however, and not in the which estimates and the system of the start of the start of the start contentials approach and the gamma start and the start in location contentials, approach and the gamma start and the start in location contentials, approach and the gamma start and the start in location contentials, approach and the start interaction in location. Each start and the start models controlled start and the start interaction in location. Each start and the start models are controlled and the start interaction in location. In the start is the start models are controlled and the start interaction in the start in the start interaction in the start models are controlled and the start interaction. The start is the start is the start interaction in the start interaction in the start is the start in the s

Excess water reterior may be addressed with the use of arciliary medications and as the anis-strongen turnoficine circles or an interview inhibit or aches smallers that the the activity of astrogense, these drags can effectively reduce the level of faced value. It must more effective at a shorting this gain the control of the strongense of the competition is to also use a district, which can shell excess water by directly measing rund water excession. This is signaled as the must actually pikely practices as well. Water treatments definition for a can also be one of the must actually pikely practices as well. Water treatments in share heaters and the structure of the must actually pikely practices as well. Water treatments in what heater termines on can also be one of the must actually pikely practices as well. Water termines in share the structure of the struc

Virilization

Physical (Male)

Dysphonia (Vocal Changes)

Although for loss common than dysplonin in sortene, antolocizandogenic staroids may alser vocal physiology me mer. This may technic alseopering of the voluce, Dysplonin is more common when anabolicandogenic staroids are administered daring adolescence, as the doper addit voluce has not yet been calsibiled under the inflament of analogens. The administration of antiolociandogenic staroids before mutriy cart, likewise, casse a progenosice lowering of the woral phik, and miny tigger phenocet out darges in younger Albhough a digit lowering of the wise may be noticed with androgen uses more cases, proport of criscard phigment empty, and the used (analogi example). The same case of anyout of the starbing starter and the starbing and word lowering notice when breating and word howerenses in relation to antholicandogenic start ad abace. "The instartee data is the final to the starter and word howerenses in relation to antholicandogenic starter ad abace." This instartee data involved smoking, however, making the direct influence of steroids more difficult to discern. In general, vocal physiology is well established by adulthood. Aside from very minor reductions in pitch, anabolicandrogenic steroids are not expected to have strong audible effects on the voice.

Gynecomastia

Anabolic microgenic servoits with significant estrogenic or progenitical activity my cases gencomming (inclus) besuts diverporter, in mules). This distortier is specifially characterized by the growth of excess glandkatr issue in muc, due to an inclusate growth, and levels are promoved in the primary specific of marming splat growth, and acts upon receptors in the breast sprome ducal split-failing hyperbasis, ducat i decagnition-families, and flexible at profiles that has a specific flat glandkat these growth, ²²³ High security and the base sprogen usable present files inside advolupment is much for it is prostruct able on storagen subgrowth and the split split split is the split security of the primary split abuse by next users. In externer, cases the breast my take on a vary female hoking appearance, which is official to hisk cover with hoose cohengen.

Gynecomials lends to develop in a series of progensive stags. The severity of this process will used pending on the type and does of stags) used, and attichuids ansensity to hormouse. The first sign is typically pain in the ripple area (genecolynea). This may quickly coincide with mirror soulders manual regulations are the source of the source to as post-genecomentia, as if primmity modes af and not glandhar tissue. At this stags, it may be possible to address mit ripple source (genecolynear) the singer strendition of the source of the source of the source of the source interface of the source of the source of the source of the source strendition of the source of the source of the source of the source of the enably life in the carry stages when princing deeply around the apple. Noticeble proceedings in the source once the cource cource (source bardeet indexing) and source mathematical source of the source of the

It is importent to note that progestorence can also suggest the stimulatory effect of contengen on memory issues growshift and strange strained mass may be also triggest fractional of generozensatis in scenistic antificiality, or write a desaring levels of entropy. Many analosis seriodis, particularly show derived from matchorea, are shown to exhibit straine grospassinization with this site diffect in a matchorea, are shown to exhibit strained are consumely likely to this site diffect in a matchorea, are shown to exhibit strained are consumely likely to find the site of the strained are not strained are shown to exhibit strained are consumely likely and the site diffect in a matching and the site of the site of the strained matching and the site and the site of the strained are strained are shown to exhibit strained are consumely likely and site and strained are strained are strained are strained are income and the site of the site of the strained are s





Early gynecomastia. Physical (Female)

Birth Defects

Ambolicantloganic steroid eposure to a wormu during programy can cause developmental abmoralities in an utroffic metal. Neithation of a frame filters is a particular concern, and may include clickoal hopertophy or even the growth of ambiguous ganital ingreadchematphrolitantiis. Neurona who are pregured, or an admentiate to hoccean beneficiant abnormalities. Worms who are pregured, or an admentiate to hoccean produce, gibe, orientee, patients and an admentiate to hoccean produce, gibe, orientee, patients and antibioinclungstrast streameds can relative speen coant and ferdity in men, they are net fiscal to beth defets when taken by sonceore fichturica edult.

Dysphonia (Vocal Changes)

Anabolicantogan steroids are commonly fished to a despering of the voice in familes. This is is cancel by diret analogane influence on the layers and matche biases involved it voce in physiology, which (in females) are not normally exposed to high androgan lovels. Early changes may include a gliff tomoring of the voce, with anable shafts in plicit at he higher layer and of the vocal spectrum (against quadrag and voca projection).²⁰⁷³ There is to prizally a simulation of the vocal spectrum (against quadrag and voca projection).²⁰⁷³ There is to prizally and to achieve the simulation of the programs, these changes must play on to develop into a mergy and neorginable mode-durated trials vocioe.

Depending of the voice is defined as an anthogenic or muccinizing effect. Anabolicatingpuis actions, with high realise antiogenity work as testostrones, flavoymsterone, and methanoloscolore. Boevies, have high tendency to produce voice changes when used in Branks 2²⁷ 220 a 229 all anabolicationgpuis returing a sensitivity of a sharing the female voice given the right level of similation, however. To this point, would change have been repeated and the Branches and Branches abstraces such as conclusions and mucholoscol 2¹³²³ Case must be taken to meintor the voice during of AdS instate, as changes are taken using generated. Immediately durotized and taken taken and mucholoscol 2¹³²⁴ Case must be taken to meintor the voice during of AdS instate, as changes are endore using generated. Immediately durotized are levely to presist. Anabolic/induogenis stratuli are may, Beevise, permanently alter vocal physiologie information.

Enlarged Clitoris (Clitoromegaly)

The mik and fimits reproductive systems differentiate and develop under the influence of estiogna and testistocome. Even as an addition, the fimits reproductive system remains developmentally neprotexits to mike as hornerss. An developin of the analogen lived in nor advanced angle the maximum system of the strategient remains on advanced angle the may lack to writemation of the external generation. And external horner and the choice (classroomgab), With (chorenegab), the choicemy degin to reserve the same prime, and may never widely ending a daring secand arroads (restriction) in more service cases is association to a mile priori, card by terropic systems (restriction) and prime in advance of the observang advance on the system and intervertion when is own is noted.

Chromengaly is most commonly scen as a congenial disorder, although it may be caused by anabelicatiogosis as scool administration or other pathology in additiond (acquired chromengh). As a vitating side effect, chromengal trank to scear in a dose-dependent disorder and the state of the state of the state of the state of the state indication of the state moder. Through any state of the state of the state of the state of the state state of the s Chromegaly caused by steroid use is both avoidable and progressive. Mitgaing eccess androgenic action early when it is noticed is the most findamental part of treatmert. Reveal of significantly developed tissue, however, will require reconstructive surgery (efforoplasty).³²³ Special caree should be taken to preserve the dorsal and ventral neurovsecular burdles and normal tisse scression.²²⁴



A photograph of distinct clitoromegaly.Here, the clitoris begins to resemble a penislike structure under androgen influence. If left unabated, this may progress to a more defined phalic abnormality. Source: Copcu et al. Reproductive Health 2004 1:4 doi:10.1186/1742-4755-4.4

Aubclichtorganis strockis my cause male-pattern hite growth in familes. Modeildy defined a listication is conclions it characterized by the growth of hite in antogens resistive areas of the body. With hitestation, dark course hair (terminal line) my develop on the face, check, alchorem, and beak, cause of the body roumaly associated that in growth in reartion. The structure of the hitestation typelagie involves immediate abstraces from antibulicantlogeness terminal terminal structures and the structure of motion of a structure of matine antogens (sprice) and the hite facility. The fact material structures antise antogens (sprice) body and the line fact dark and in fatting spring that and antises the structure of the mathematic structure in the structure of matteries of matteries of matteries of mathematic analysis. The structure of the analysis in the line of the structure of

Menstrual Irregularities

Ambléinantograis éstenisk may aler the mestraul cycle in femilies, resulting in infinguent or absort merses (amonthena). Fertily may also be intermpted. Normal mestratanion is expected to resume after anaboléindrograis steroiks are discontinued, and the natural hormean balance is setstend. Complete recovery of the female hormean also area and fertility can take mmy morthe in some cases, however, and long-term interruptions of fertility are possible, thoradh mildely.

Reduced Breast Size

Anabolicanloganic strenisk can iribit the growth- supporting effects of entropen on memory issues, and my cance a visible relation in phenesis the (breast antoph). Androgan use in formals has specifically been shown to cance a reduction in glandhar tissue size, and to promete an increase in throse concredent issues.²⁴ There physiological changes are similar to finder noted after manopanae, when formits sets strends are very low, of other strength strength

Psychological

The effects of nucleoicandogenic stroxis on human psychology are complex, controvensia, and not fair underschool. What is known for certain is future scattering psychology. They phys arek it an individual general invok, alternses, agaessias, stense of differences belowers on run and wrants beaccies of differences is as stensed. Took, and, Revoixe, altergin kernen and wrants beaccies of differences in sets stendie levels, and, Revoixe, altergin kernen and wrants beaccies of differences in sets stendie levels, and, Revoixe, altergin kernen and wrants beaccies of differences in sets stendie levels, and Revoixe, altergin kernen and wrants beaccies of differences in sets stendie levels, and mann psychology. The certain strange of this sociation, however, runnis for subject main psychology for psychological humbits and the base in the sets of several lay areas or psychological humbits.

Aggression

Men tati to be more aggessio flux worms, a durateristic that has been party ambied to higher androgen beek-27³⁹ Physiological, undrogen are known to ext on the angulation and hypotheliums, areas of the first involved in human aggression. They also affect the orbitotical cortex, areas incoled with implementation of the state and the sector of the state and the state of the state and the state of the sta

The psychological effects of examining domages of beststorme esters have been construct an answer of phetoschormold statisks. At thereparkic levels, no adverse psychological effects are apparent. If anyting stessistence replectment therapy tanks to prove model and sense of whether by Wenn such at a constructive domage (200 mg er wesk), agin, to appending effect and a specific statisk and a specific domage (200 mg er wesk), agin, to appending the statisk at constraints whether the statisk and a specific statisk and a speci

Criminality and Violence

Lisks between anabolic indicagenis storoid abuse and vidence have been much mere difficult to establish. Most papers suggesting such an association effert used correlative data, or discussed infisidual cases studies. These help broaden the scope of research, but are not visible for proving cansily. For example, one study questioned a group of 23 storoid abusing mere, and reported that these mere were incoled in a significant data of the study proving the tracks they were in and every study of the study and the study of animatering AAS ongs.⁻⁻⁻ with the known entexts of antisocaranoppene serons on agression, this fulling is competing. It muy very well be that some men are mere susceptible to this type of behavior when absuing AAS thurn others. A paper like this is not sufficient however, to substantiate a violent "violar "get" Futher research is needed to determine if trategies and the state of the trategiestic state of the trategiestic state of the state o

Serior criminally has also been diffical to associate with staroid abuse. When discussed, we again red to see welco cortikine data and cess statis. For example, one paper is storedare property an association between stored abuse and weapons and final crimes.²⁴⁷ It is incertain, however, if stored abuse was actually supportible for his crimenilar, or paper associated with I. It is simply possible those men were more exposed to, or more likely to use, *ligal AMS* for example, *in a lineary fastis*. For advecting the abused analytic dampestic abused and the abused analytic dampestic abused and the store and the store is an extent of the abused analytic dampestic abused abused abused abused abused abused and the abused analytic dampestic abused abused abused abused abused and abused competing for dam, there is no conclusion module abused analytic abused analytic abused analytic or stores crime into hybrids in a provide primely table intribution dates can enance where or series a crimed hybrids in a provide mentily subh infiniciation.

Dependency/Addiction

Anabolicitatioganic starokis are considered to be drugs of abuse. Albough there is no university accepted facilities for this, abuse is controntly described is a bistance of a substance in spike of adverse consequences. Given the magnities hash consequences that are subscritted with synthemperiod close of AdVA sings, the closelitation is a difficult on to dispute Drugs of abuses are very often also drugs of dispersion-y, which in this context dubts correctly one or an anabolic starokis also that do offen years of the drugs of dispersionfluence on the start of the dispersion of the dispersion of drugs of dispersion-y. Furthermore, among those that speptor the rokis of an anabolic steroid dispersion-y, there is a spik with argued to the nature of this dependency (speciohigical or physich).

Physical dependency is usably regarded as the most serious form of drug dependency, advogable hot process on the voy extreme and two tobing depending on the stantors. Physical dependency is defined as the need to administer a substance in order for the body to function normally. A physical dependency is usually demensioned by the dependency and with the substantiant of the drug as discontinued advergth. The most well known examples of drugs of physical dependency or queuing such as morphics, hydrocochone, successful and Optates cannot very difficul drugs for dependent individual to qui using size stopping the's use tanks to produce cannot well-known by morphics and physical physical services physical optendency cannot be drugs to weeks after for drugs in discontinued, while the procludesign average more mysical for morthe locarc.

Anabolizationgenis storoid abase could be associated with many of the DSMV1 vertein messary for a diagonis of both probledgical and physical diag designedres, For instance, it is not measurement for someone to take the drugs in lighter does or for longer periods of time then the hydrid high planed (clients) [11, Many Jakens abase to a close to coperiormers may present this decision (criterii 82). Individual of the common to abase storoids in spite of regulative heat concentrative comparison (criterii 87). Should abase is also associated with a drainsking level of effect and escalating dosages (criterii 86). Individual robiced see clients, fingue, depression, menuna, saicial floright, realescence, lack of storoids.²

According to the American Psychiatric Association and its Diagnostic and Statistical Manual of Menial Disorders (DSMIF), three or more of the following criteria must be met for a diagnosis of psychoactic drug dependency.

1. Substance is taken in higher doses or for longer periods than intended.

2. Desire or unsuccessful efforts to cut down or control substance use.

3. Excessive time spent obtaining, using, or recovering from the substance.

4. Important activities are given up because of substance abuse.

5.Continued substance use despite negative psychological or physical consequences.

6. Tolerance, or the need for higher amounts of the substance to achieve desired effect

7. Withdrawal symptoms.

A drug dependency that is isolated to criteria #1 to #5 would be described as associated The meeting of criteria #6 or #7 indicates the dependency is also a physical one.

The physical benefits of anabolic/androgenic steroids complicate the matter of drug dependency a great deal. Unlike parcotics, the main motivator behind the use of steroids is their positive effect on muscle and performance. With this in mind, steroid addiction could actually be a misdiagnosis for muscle dysmorphia in many cases. This is a psychological disorder characterized by nersistent feelings of physical indequacy in spite of extreme muscular development. Steroid abuse (often extreme) is highly common in muscle dysmorphics along with compulsive resistance training ²⁵⁰ But steroid abuse is regarded as a symptom of this disorder, not a cause. In a similar sense, the physique-, strength-, and nerformance-improving qualities of anabolic/androgenic steroids could be driving much or all of the abuse. An analogy would be the so-called addiction to chocolate. Some individuals develop tangible psychological issues surrounding the consumption of chocolate, with uncontrolled binging and negative social and health consequences.²⁵¹ But we do not regard chocolate itself as a substance that causes dependency.

There is some evidence that the reinforcing qualities of steroid use so beyond an attraction to their physical benefits. Lab animals such as mice and hamsters will repeatedly self-administer testosterone and other anabolic/androgenic steroids for example, an effect that cannot be caused by a perception of physical change.252 Testosterone is also known to interact with the mesolimbic donamine system, which is common with other drugs of abuse.²⁵³ 254 Studies additionally suggest that anabolic/androgenic steroids influence donamine transporter density. and increase sensitivity of the brain reward system²²⁵ Steroids are known to influence is interested with the second system of the second system of the second se confidence when taking AAS drugs. Some speculate this is due in part to an inherent psychoactive effect. Further research is needed to determine if anabolic/androgenic steroids are actually mild respective drugs

Anabolic/androzenic steroids are not drugs of marked intoxication 256 which makes them very different from other drugs and abuse or dependency. This makes diagnosing a drug dependency difficult. By definition, drug dependency is related to the abuse of a psychoactive substance, and it is unknown if AAS drugs can accurately be classified as psychoactive substances. At the present time, most experts do not regard anabolic/androgenic steroids as drugs of true physical dependency. It is difficult to correlate the post-cycle hormone imbalance with traditional withdrawal symptoms and tolerance is really a function of metabolic limits on muscle growth not necessarily a diminishing biological effect. Individuals remain warned, however, that steroid abuse is commonly associated with the sions of nsychological dependency. Further research is needed to evaluate the biological and psychological nature of steroid abuse.

Depression/Suicide

Anabolic/androgenic steroids abuse may be associated with bouts of depression. This is most common after the administration of AAS drugs has been discontinued, especially following high doses or long cycles. During the time that steroids are being administered. natural hormone production is diminished because the body recognizes the excess hormone levels. When the steroid drugs are abruptly discontinued, however, the body can enter a state of temporary hypogonadism (low androgen levels). This may be associated with a number of psychological symptoms including depression, insomnia, and loss of interest. This condition is usually referred to as anabolic steroid withdrawal depression, and can persist for

weeks or even months as the body slowly resumes normal normone production.~...

The not common method of addressig anabols stored withdrowed depression in men is prompticely, with the implementation of an agressive poor-yeak hermone recovery program. These programs are typically based on the combined use of HCG (thurma chronice grandcorport) and aris-storagarc inform gost and tearolings. They are used together in a way that can stimathe and sensitize the hypothalmic pittatry testicular axis, allowing mattrix hormese production to return more agaidy. Advantably of depression fallowing action with the stored protection in the hypothalmic pittatry postcular depression fallowing action with the stored protection with the depression is prolonged on servere.²³⁰ These drugs must be used with cardine, however, as they also have been linked with increased houtget of solate its norm praters.²³⁰

Albuqh isso commo, depression is sometimes reported during the active administration of anaboliant/organic sension. This may be caused by an intributer of oss served i levels, particularly with regard to relative anthoganizity or estrogenizity. In more cases that more, it will move a studient where addicert andoparity is not present, study when primulandboke during probability howers, it is difficult to early outline the promotion strengest with human probedings howers, it is difficult to early outline the promotion strengest with human probability howers, it is difficult to early outline the promotion strengest can insolve either elevated or suppressed levels of central set strengest strengest end and the strengest of the strengest difficult to early difficult to provide the probability strengest end and strengest of the strengest difficult to the strengest action.

Sacide hus been linked to anabolicitential abuse in new instance.²⁰⁰ Such reports are usually case studies, convoluping individual huw ore belowed to be psychological yabibe before abusing AAS, and who committed assisted during our after use of the drugs. It is known that depresents in a controm comparint during anabole stared withfarmed. It is able known that a small percentage of users are especially sensitive to the psychological abuse insubiointrologous extends, and noted committee model strong, miss behavior, and/or secree depression with their use. It is unknown by these infinduals have such enterme reactions, while the varies of the strong of the strong the strong strong the strong the strong stron

Insonnia

Anabolic miningenis thereid uses in induced to Themos levels, and has been noted during both caccoss and imaffactor humanol states. For example, mean its is a common compliant of the state of the states of the states of the states of the states of the reported by started bases a damage the projective mixed may project when an advance of the states of the line of the states of

Reproductive (Male)

Infertility

Anaboliciandrogenic steroid use may impair farility. The lamma hody strives to matinai balance in its sec hormore levels (hormostissi). This balance is regated largely by the hypothanine'-patiany-testicalar axis (HPTA), which is responsible for controlling the production of testistorone and sperm. The administration of anabolic-inhologenic storoids provides additional sex storoid(s) to the body, which the hypothalamas can recogate as excess. It reasonable to this excess by recularia starot the modulation of obtainary gradotropris laterizing hormone (LH) and folicle-stinulating hormone (FSH). LH and FSH normally stirulate the release of testosterone by the testes (gorads), and also increase the quarity and quality of sperm. When LH and FSH levels drop, testosterone levels, sperm concentrations, and sperm quality may alb perduced.

When given in superphysiological levels, anabolic/indipagic structile controlly ruleace objaconceptern. This is a fram of related efficitly characterical by hiving lass fram 20 milition spermitona per ni of igicalute. The quility of the sperm may also be impaired under the inflaxec of AAS, as noted by an increase in the marker of absential of hypothesis (motivity append) and the high hypothesis in the marker of absential of hypothesis frame structure of the structure of the structure of the structure of AAS. (We append and miles by the hody.) Frank dots of conception are any singilizarily house that when spent concentrations are around. In many cases acrospormit is readed during Comparison in the probability with the structure dots of conception are specification of the structure of AAS. (Strilly mean set and the birth hody.) Frank dots conceptions are produced in the structure of AAS. (Strilly mean set and the birth hody.) Frank dots conceptions in the structure of the structure of AAS. (Strilly mean set and the birth hody.) Frank dots conceptions in the structure of the structure is conduction of the CO 3th.

Drainshold fertility is considered a revensible side effect of ambolicandopsein stread abase. Sperm concentrations usualty retrain to normal within several methods of discontingdrug trainale. A substatiati post-cycle recovery program based on the use of HCG, traination, and choinghne muy significantly solverts the reflactory posti, and is highly recommended among those in the storoid-sating community. In a smill percenting of cases, professional and the storoid-sating community with a percenting of cases, professional and use to a specific of the storoid storage. Concervo, The Star Star Storage concentration of the story of the store to a store of the store

The ability of ambolicandogenic strenks to appress LIF SNL and fertily he initiated a gene dual of research arrowing their to see a mic contraceptics, high-tike lettstrenks has been extensively studied by the World Heah Organization, for example, and was determined to be as als and moderable deficient endel of mith bit hortcorn it. In states which admissioned 200 mg of its structures are under the work to heahly mere, asconpermic was calsedue a fieldy of patients which its morthel²⁰⁰. Most of the comming patients was disponsement. This distributed bettily was fully researched, and baseline speem of agrossopermic. This distributed heatly was fully researched, and baseline speem of high ascopermics in the desired edspect of mice correspective, however, and this insectional has desired edspect of mice correspective, however, and the lines or how no fieldly achieved with AAS drags along, even in high doses, ²⁰ Autobicandogenic strenks have a rest approved price as mice correspective.

Libido/Sexual Dysfunction

Anabolicintegenis steroids may alter sexual dosise and fractioning. The nature of their effects, however, can use of performing on the dorg(s) and dossay(s) used, as well as the individual assistion of the user to hormanial manipulation. One of the most commuscent and the second steroid st

AX50 are my also be associated with diminished Bioli and exual functioning. This could be due to several factors. One is innelficient analogues activity. These sourcements is noted with the use of primarily analosis attended as an enderstockee, methodenee, or consultatione, and the several factors is and as an enderstockee. The source of the source o are also common after steroid discontinuance, when endogenous steroid levels are low.

Priapism

In very rare instances, antibolizmentopagies attorish have been lisked to pripigaries²² 278-271 first is a conflict characterized by the development of an excertion late of late trans to be flucied status within face house. Pripagin is a potertially very serious condition, which can popula damage creation of the status of the status of the status of the priori. More pripigarin is field to started use, testosterore is usually responsible. Furthermore, this condition appears to be more flucation transport priority and proposable transmer test pripagarism. This may be due in part to a rapid is access in and/agenerity, in pripagina very million dovelopm in addition of anisotation.

Prostate Cancer

Prostate cancer is dependent on androgens. This disease will not develop f androgens are derivened from the body as a young age one with constrained.³⁷² and laberents of androgens activity in patterns with active disease is regarded as a standard path of treatment. A complete pixture of the non-bornet of androgens, however, termins uncles. Stadies show there is no sociation between the testosterome level and likelihood of developing grounds cancers.³⁷² On the same che, the administration of composite testosterone their grantegaregarder and the statistical statistical statistical statistical statistical statistical statistical pixels of the model between the testosterone level and benchmark the statistical statistical pixels of the model between the testosterone level and statistical statistical statistical pixels of the model between the testosterone level and statistical st

New diagons of prostate caree an continues reported daring teststerone replacement theory and startin discess $^{-734}$ Scie theorytic may be here stud of pervisely indigonous confinito en urrelated development of this discess, with androgen stimulation assisting the turne growth rate. May form of prostate career prossess functional androgen receptors, and are highly androgen responsive. As such, they can be simulated to grow under the rithtence of teststerence or other antiobiciantogenesis steensis. Games administration approach with the steeroit administration approach with which a bisety of prostate career, arXiV able steeroit administration approach with the steero of the real-bioliteat career, arXiV able steeroit administration approach with the use of teststerence or other antibic strate career, arXiV able provides and the use of steststerone or other antibic able career print/able antibic real strate strate and miggant prostate acree or advelues of the transport match with previously undigrowed (latery miggant prostate career of advelues).

Prostate Enlargement

Anabolicardoganie starodis may inflance the size of the prostate. Androgani may impact to the development of the prostate in onely like, and are essential to maintaing prostate structure and liketion throughout additional. Increases in the androgani bed often result in volume has been compared as the prostate of the provide structure and the of hypogenetic list owner paties: receiving testotence relations for the data outdrafts, prostate work one does tend to reacting testotence and the major data androgan the structure and the structure of the prostate of the structure of the androganic structure of the struct

AmbiGrandroganis strend abase may result in significant increases in prostate volume. In more severe cases, this my lead to being protect hypertophy (FeBT) BPT is a commo condition in older men, characterized by reduced urine low, diffically or disconfit training, and changes in intrusy frequery. Accordant propts of FBT among seriod-isangle bodybalates are not common, but do occar with enough frequency to warnet concern. Such reports a men colom hield to more and/nogic drug such as testosterone and terbolone, or the accessive dosing of AAS in gazend. Due of the most eatherm reports of prostate hypertophy came from Dr. John Zealer, the U.S. Ohmer, physicain accerding with introducing Dianabol to sports.²⁰³ Dr. Zegler noted that during the mid-1950s, many Russian weightfilters were abusing so much testosterone that they needed catheterization to urinate. Dianabol was released soon after, which is structurally a close derivative of testosterone with reduced ant/orgenicity.

Statis of antholeinthogenic acroid abuses show a preferential simulation of the irner prostate under the situation of these damps, and an area where beings prostate logarcheights is known to originate.²⁴⁴ In contrast, prostate curcer usually develops in perplexal areas of the given. Stress association between BHP and prostate, curcer is known to osist, however, albrough the exact rature and strength of this association remains userstain.²⁶⁶ PAA values are done (albrough rature) solution biodicolitorities are number of protential curvinitions and biodic testing for PAA beek, Autobecknindogetic strend is the sprently invincitatly discontrastic figure of BHP or curced TFAA walls sets and the strength of the strength of the solution of the strength of the sprently invincitatly discontrastic figure of BHP or curced TFAA walls sets and the strength of the strength of the strength of the strength of the sprently invincitatly discontrastic figure of BHP or curced TFAA walls become approxet.

In addition to anticogene, estrogene are also loxone to be involved in prostate growth and functioning.²⁰⁰ When handprogene are generally initiatively ownsh prostate growth, haveentingen receiptor (Eds-bas shoppe) may help protects the prostest from influentizon, estrogene receiptor (Eds-bas shoppe) may help protects the prostest from influentizon, esla properties, and carcinogenesis. Conversely, stimulation of the alpha subtype of the estrogene receiptor is liked to showmice of proferints, influentizon, estimations of the shopse of the automation of tastistence and AAS (blokh will result as stimulation of the receiptor shopped shopse) will effect protects the prostestic genomical and cascing estimation of tastistence and AAS (blokh will result as stimulation of the receiptor shopped statistical and the statistical strategies and the shop of the company strategies and the protective genomical methods and enough strategies and the genomical strategies and the strategies and the shop of the prostate 2²⁰ Presently, the most accessed strategy to minimizing prostate hepotrophy some to be located enclosing relative analysis, genomica, texture, sciento.

Testicular Atrophy

Anabolicantingamic strends may produce anophy (stringago) of the testicks. Fostosterone is sopatistical and secretably but Legited in the tests. Its relates in regulated by the Legited in the tests is the relates in the strends. The relation is a system that is vary sensible to sex strends. When anabolis strends are and diministend, the HFT WA and receipts the deviated homore levels, and respond by reducing the synthesis of lesistaterones that are more both as so of resistant when an adverse that the relation of the strengs of may not be obvious to the individual. In strength many drings are by these to the individual in strength energy drings are supported. When strends are requery to the obvious of the individual in strength second the more comes, of adversal show, appearing in more than 50% of all anabolicandogonic steroid abuser, ³⁰⁰ 30

Although testicular attrophy is very common in frequency, it is also regarded as a temporary neurobic kide effects¹⁰. The granch, by their ratter, will vary as its under hormout inflatore. Anotyby should not produce permanent during, Node, however, that it can be a single structure of the str

Some of the mere potert analysic/indugris storolds, including testosterons, multiolose, therbohar, and oxynetholise, appear to be more suppression of estosterore release than many other AAS drugs. This may be explained in part by the additional estogastic organisational activity interest in these activity, and estogastic model activity in provide regarise feedback inhibition of testosteneore release.^{40,40} It is important to most, severeton. This involves more model, conversal, and its proposition and activity in the estimation of testosteneor estates and its proposition activity in the estimation of the estimation of the estimation of the severeton. This is involves more model, conversal, and its more than the enversa. coundrolone, which are normally regarded as milder in this regard. While these compounds may be less inhibitive of testosterone synthesis under some therapeutic conditions, when taken in the sapratherapeutic doses necessary for physique- or performance-enhancement, significant attrophysical supersistion are common, and distinctions less pronounced.

Other

Anaphylactoid Reactions

An amphibicatil exection is a series and poteriably file freaturing allergic response to the administration of a foreign substance. Symptome of this indexer funck letting of the suband topics, seeling of the meson membranes, hore, howeved blood pressure, abdominal contrast, thereby executing barreling. Aquation of the subtional method by unconsciousness, corns, or don'th. The physical symptoms of this disorder are medication by the reisson of limitsrine. Isokorize of 20, possignation 12, and typicase. An amphibication of the subtion of the sub-transmission of the sub-transmission of the disorder are medication by the reisson of limitsrine. Isokorize of possignation 12, and typicase. An amphibication control medication is more characteristics as amphibical standard on rungs (offen) significable medications, however, Mid skin reactions may be effectively treated with an ambinations. How earlies many heating the sub-transmission of the sub-transmission of the sub-transmission may reage the effectively treated variable medications, however, Mid skin reactions of the sub-transmission of the sub-transmission may reage the effectively treated variable medications, however, Mid skin reactions of the sub-transmission of the sub-transmission may reage the effectively treated variable medications, however, Mid skin reactions of the sub-transmission of the sub-transmission may reage the effectively treated variable medications however. Mid skin reactions of the sub-transmission of the sub-transmission standard to sub-transmission the scape of an amphibication reaction develops.

Cancer, Brain

Anabolizatioganis strends are not susceited with hera encarec. Complications relating to a new and usually fails from of carece called primury certain renews asystem (rimalymphorm caused the dash of finous American football player Ljek Atado. This type of their ancers rene sourced appears in intrame-compounding builties, such as those stifting from Hamm Immandelecticny. Vite, (HTV), or organ transplart recipients taking immanageneous data to presert organ pricedon.²⁰⁰ 300 Bellet ambolizationganic streates and an anti-ball immandepression taking ambolizationganic streates and an anti-ball immandepression.²⁰⁰ Mellet ambolizationganic streates and an anti-ball immandepression theorem the machine tensors called a more methanol to call that to go and schemenmethanic and the streates and the streates and the streates and streates the streates and the streates and the streates and the streates the streates and the streates and the streates and the streates and streates and the streates and the streates and the streates and streates and the streates and the streates and Ataba's cameer.²¹¹ this streates and streates and the media to correct the streates and Ataba's cameer.²¹¹ this streates and streates and the streates and t

Cancer, Breast

Albade atorney me, mic breast cancer has been associated with the admission of tostostocous²¹ H is atoron, however, fit hormore harpeny so relied to be once of this disease, or f it was just incidental to its progression and discoursy. Androgen generally oblit inlikely effects on hormore-response breast cancer, and have statuly been used in their tensment.²¹¹ Estingent, on the other brand, can support the growth of many breast tarms. It is not accounting for each of early more than the statule breast cancer than the statule of the statule of the statule of the statule breast cancer. The breast cancers and basistence admission resumes meaker. Testostence is present/containdicated in patients with breast cancer.

The data concerning the effects of analysis/induced is a strain or starking is a difficult to interpret. It does append that when these data gas as used a ringle, reactions in hody that are common particularly viscent adapted uses that the strain of the

Insulin Resistance

Anabelicandeganies starolsk may aleer inadin sensitivity, an important measure of metabolic heahth. The effect of these drags may be variable, however, for example, testostores administration may improve inadia sensitivity in max with hpogenedism²¹¹⁴ O (20) map et adv) has also been shown to impore inadia sensitivity and detra me(d) to 53 yeass).²¹³ These hereficial metabolic outcomes were correlated with reductions in viscent adapose tisses (APA). This is a doep layer of that surround the addormal outgons, and a space tisses (APA). This is a doep layer of that surround the addormal outgons and a space tisses (APA). This is a doep layer of that surround the addormal of the AIA is subscitted by hortensed in existence. The Audeing May and and sufficience and AAS may improve indin sensitiva, and potentially metabolic health.

Conversely, anabolic/androgenic steroid abuse has been associated with impaired plucose metabolism³¹⁷ In one study, nowerlifters that had abused AAS in high doses for up to seven years were shown to have diminished plucose tolerance and increased insulin resistance 318 In spite of a long history of resistance exercise, these subjects secreted more insulin in response to measured placase ingestion than even obese sedentary control subjects Additional studies with methandrostenolone demonstrated significantly increased insulin secretion and notential resistance 319 A similar outcome is not found in all AAS studies. however. For example, testosterone enanthate in doses as high as 600 mg ner week for 20 weeks failed to produce any changes in insulin sensitivity in healthy young men.320 Nandrolone decanoate (300 me ner week) also did not immair placose tolerance, and actually improved insulin independent plucose disposal 321 The data concerning the effects of anabolic/androgenic steroids on insulin sensitivity is difficult to interpret. It does appear that when these drugs are used initially reductions in body fit are common particularly visceral adipose tissue. This may actually improve insulin sensitivity and the overall metabolic state. and reduce certain specific risk factors for diabetes and cardiovascular disease. Beyond this the effects of AAS on plucose metabolism are not fully understood, and difficult to predict. Studies using supratheraneutic doses of testosterone and randrolone have failed to produce any negative changes, suggesting that moderate AAS abuse is probably not associated with impairments in insulin sensitivity. At the same time, studies do suggest that there could be concerns with heavy steroid abuse. Further research is needed is assess the impact of steroid abuse on metabolic health

Nosebleeds

Anabolicantingamic stored use may be associated with predict mosehoods. According to one study, approximately 20% of disk stored users reported this sidely, maintig disk community. "Nosebheads are not a direct result of antroparite action, but are secondary to storio induced increases in blood pressure and/or moderism in blood chriting factors. Although they can be scarry, most mosebheads are laminess, and will not require emergrary medical antiruli, Whore include to stored action. Showever, the ymmy medical theorem (who theorem the indical antiruli, Whore include to stored action, stored actions and will not require emergrary inflavor tunally upon properting shortly during discontinuere, as blood pressure and/or column factors. The store of the result stored store.

Sleep Apnea

Obstructio skep aprox (OSA) is a disorder characterized by brief passes in horaling drarg skep, which core when the soft sizes as in the free of lose at the passages, productively of kkep. It may also choose the hermateer, it fixed nut histod, and passages in other than the state of other hermateers in the state of the obstruction drarg loss including hypermetation and conflocated drares. Since parameters are not confirmed as the state of the state of the state of the state of the obstruction drarg loss (hyperbase) (CSA) (hold defines (hyperbase), soriering, notating analoging, and meeting headbacks. Obstructive skep apress access to occur most and the state of the state

Anabolic/androgenic steroids may be associated with the development of obstructive skeep apnea in a small percentage of individuals. The exact relationship between AAS and OSA, between the sterior modern and the obstructive sterior to anabolic income activate modeling nwever, remute tacker. In a soverse reaction scents to appear in some patients recoving testosterione drugs to treat hypogonalism³²⁶ More detailed studies have shown that high does of testosterione can disturg shep and breating as well as increase skep-related hypoxenia, effects that may precipitate obstructive skep aprea.³²⁷ While OSA has not been clearly documented in steroid abases: andhores have been shown to have the structure and

function of the oropharynx in ways that can predispose an individual to this disorder.²²⁸ More research is needed to determine if steroid abuse can trigger OSA in an otherwise healthy person. Individuals with a history of obstructive skep apnear should not use anabolicantrogenic steroids. Physicians are advised to monitor their patients closely for sime OGSA durine AAS theraro.

90. Use of anabolic steroids and associated health risks among gay men attending London gyms Bolding G.Sherr L,Elford J.Addiction.2002 Feb;97(2):195-203. ⁹¹. Hepatic lipase activity influences high density lipoprotein sabelass distribution in normotifylyeridenic merc genetic and pharmacological evidence. Grandy S et al. J Lipid Res 1999 40: 259-34.

92. Changes in lipoprotein-lipid levels in normal men following administration of increasing doses of testosterone cypionate. Kouri EM et al. Clin J Sport Med 1996 Julp(3):152-7.

93. Contrasting effects of testosterone and stanozolol on serum lipoprotein levels. JAMA 261:1165-8,1989

 High-Density Lipoprotein Cholesterol Is Not Decreased if an Aromatizable Androgen Is Administered. Metabolism. 39:69-74, 1990

 Relationship of cardiac size to maximal oxygen uptake and body size in men and women. Hutchinson PL, Careton KJ, Outz H, Wilson G. Int J Sports Med. 1991 Aug12(4):369–73.

96. Androgenic anabolic steroids also impair right ventricular function. Kasikcioglu E et al. Int J Cardiol 2008 Feb 11 E Pub.

Gardiac effects of anabolic steroids. Payne J. et al. Heart 2004; 90:473-75.

98. Adverse effects of anabolic androgenic steroids on the cardiovascular, metabolic and reproductive systems of anabolic subscription. Tuomo Karini, Publications of the National Public Health Institute ISBN 951-740-388-2.

99. Absense of left ventricular wall thickening in athletes engaged in intense power training. Pelliccia A et al. Am J Cardiol 1993; 72: 1048-54.

 Left ventricular hypertrophy by Sokolow-Lyon voltage criterion predicts mortality in overweight hypertensive subjects. Artikainen RL, Grodzicki T. J Hum Hypertens. 2008 Aug 28. [Epub]

 Hypertensive heart disease. A complex syndrome or a hypertensive cardiomyopathy? Lip GYH et al. Ear Heart J 2000; 21: 1653-65.

102. Adverse effects of anabolic androgenic steroids on the cardiovascular, metabolic and reproductive systems of anabolic substance abusers. Tacono Karila, Publications of the National Public Health Institute ISBN 951-140-388-2. 1031. Jeft vertricular Internetworks and OT discoversion in Invertencision. Maxvet 1et al. Historetencis 1996: 28: 291.

%.

104. Cardiac arrest following anaesthetic induction in a world-class bodybuilder. Angelilli A, Katz ES, Goldenberg RM. Acta Cardiol. 2005 Aug60(4):443-4.

105. Effects of training on left ventricular structure and function. An echocardiographic study. Shapiro CM et al. Br. Heart J 1983; 50: 534-39.

106. Are the cardiac effects of anabolic steroid abuse in strength athletes reversible? Urhausen A, Albers T, Kindermann W. Heart. 2004 May;90(5):496-501.

107. The effects of anabolic-androgenic steroids on primary myocardial cell cultures. Melchert RB et al. Med Sci Sports Exerc 1992; 24:266-12

 Candiovascular effects of anabolic-androgenic steroids. Melchert RB et al. Med Sci Sports Exerc 1995;27: 1252-62

109. Cause and manner of death among users of anabolic androgenic steroids. Thiblin I et al. J Forensic Sci 2000;45:16-23

110. Anabolic steroid abuse and cardiac death. Kennedy MC et al. Med J Aust 1993; 158:346-48.

111. Serious cardiovascular side effects of large doses of anabolic steroids in weight lifters. Nieminen MS et al. Eur Heart J 1996; 17:1576-83.

112. Sadden cardiac death during anabolic steroid abuse: morphologic and toxicologic findings in two fittal cases of bodybuilders. Fineschi V et al. htt J Legal Med 2007 Jar(121(1):48-53.Epub 2005 Nov 15. Review.

113 Blood pressure and rate pressure product response in males using high-dose anabolic androgenic steroids (AAS), Grace F, Sculthorpe N, Baker J, Davies B J Sci Med Sport, 2005 Sep(5(3):07-12.

114. Are the cardiac effects of anabolic steroid abuse in strength athletes reversible? A Uthausen et al. Heart 2004;90:496-501.

115. Cardiovascular side effects of anabolic-androgenic steroids. Herz. 2006 Sep;31(6):566-73.

116. Anabolic steroids and fibrinolysis. Lowe GD. Wen Med Wochenschr. 1993;143(14-15):383-5.

117. Effect of anabolic steroids on plasma antithrombin III. alpha2 macroglobulin and alpha1 antitrypsin levels. Walker ID, Davidson JF, Young P, Conkie JA, Thromb Diath Haemorth. 1975 Sep 30;34(1):106-14.

118. Depo-Testosterone. Pharmacia. U.S. Prescribing Information. Revised August 2002.

119. Anabolic-androgenic steroid abuse in weight lifters: evidence for activation of the hemostatic system Am J Hemrool. Ferenchick GS, Hirokawa S, Mammen EF, Schwartz KA, 1995 Aug49(4):282-8.

120. Raised concentrations of C reactive protein in anabolic steroid using bodybuilders. F M Grace, B Davies et al. Br J Sports Med 2004;38:97-98.

121. Testosterone increases human platelet thromboune A2 receptor density and aggregation responses. Ajayi AA, Mathar R, Halashia PV. Circulation. 1995 Jun 19(1)(1):2742-7.

122 Androgenic-autholic steroid abuse and platelet aggregation: a pilot study in weight lifters. Ferenchick G, Schwartz D, Ball M, Schwartz K, Am J Med Sci. 1992 Feb;203(2):78-82. 123. Pulmonary embolism associated with the use of anabolic steroids. Liljeqvist S, Helidén A, Bergman U, Sideberg M. Fur I Intern Med. 2008 May 19(3):214-5. Enub 2007 Sep. 19.

124. Converse thrombosis and ectasia of coronary arteries after long-term use of anabolic steroids. Tischer KH, Heyny-yon Haussen R. Mall G. Doenecke P. Z. Kardiol. 2003 Apr:92(4):326-31.

125. Massive pulmonary embolus and anabolic steroid abuse. Gaede JT, Montine TJ. JAMA. 1992 May 6/267/17/2328-9

126. Stepid appholic dross and attenial complications in an athlete-a case history. Laurche GP Anaiology 1990 Nex:41/11/064.0

127. Death caused by pulmonary embolism in a body builder taking anabolic steroids (metanabol). Siekierzy/ska-Czanecka A. Polowiec Z. Kulawijska M. Rowinska-Zakrzewska F. Wad Lek. 1990 Oct 1-15:43(19-20):972-5.

128. Completion abnormalities associated with the use of anabolic stemads. Ansell JE: Tiarks C: Enirchild VK: Am Heart J 1993 Edv 125(2 Pt 1):367,71

129. Hematocrit and the risk of cardiovascular disease-the Framingham study: a 34-year follow-up. Gagnon DR, Zhang TJ, Brand FN, Kannel WB, Am Heart J, 1994 Mar:127(3):674-82.

130. Homoevsteine induced cardiovascular events: a consequence of long term anabolic-androgenic steroid (AAS) abuse M R Graham F M Grace et al Br 1 Sports Med 2006d0544.48

131. Homocysteine and cardiovascular disease: time to routinely screen and treat? P. O'Callaghan et al. Br J Cario 2002:10/2)115.7

132. Promotion of vascular smooth muscle cell growth by homocysteine: a link to atherosclerosis. Proc Natl Acad Sti 1994-91-6369.73

133. Homocystinaria: metabolic studies on three patients. Brenton D et al. J Pediatr 1966;67-58-68.

134. Homocysteine, and atherogenic stimulus, reduces protein C activation by arterial and venious endothelial cell. Rogers G et al. Blood 1990:75:895-901.

Homocysteinethiolactone disposal by human arterial endothelial cells and serum in vitro. Dudman N et al. Arberioscher Thwenh 1991:11:463,20

136. Plasma homoevsteine levels and mortality in patients with coronary artery disease. Nygard O et al. N Engl J Med 1997:337:230-6.

137. Effects of sex steroids on plasma total homocysteine levels: a study in transsexual males and females. Giltay FI Hoorson FK et al. 1 Clin Endocrinol Metab. 1998 Eeb/83(7):550.3

138. Homocysteine levels in men and women of different ethnic and cultural background living in England. Carouccio FP. Bell R. et al. Atherosclerosis. 2002 Sep:164(1):95-102.

139. Hyperhomocysteinemia in bodybuilders taking anabolic stemids. Ebenhichler CE: Kaser S et al. Fur 1 Intern Med 2001 Feb:12(1):43:47

140. Homovosteine induced cardiovascular events: a consequence of long term anabolic-androgenic steroid (AAS) abuse. M R Graham, F M Grace et al. Br. J Sports Med. 2006;40:544-48.

141. The effect of summhysiologic doses of testosterone on fasting total homocysteine levels in normal men. Interence of supriphysiologic uses of essistence of fusing total nonceystene evers in a Zmada JM, Bausseman LL, Maceroni D, Thompson PD. Atherosclerosis. 1997 Apr;130(1-2):199-202.

142. Hypertension. stroke, and endothelium. F. Consentino, M. Volpe. Cur Hypertension Rep. January 2005: 7(1);68-71

143. Differences in Vascular Reactivity Between Men and Women. Bob J. Schank, MS et al. Anniology, Vol. 57. No. 6, 702-708 (2007)

144. Flow-mediated, endothelium-dependent vasodilatation is impaired in male body builders taking anabolicandrogenic steroids. Ebenbichler CF, Sturm W et al. Atherosclerosis. 2001 Oct;158(2):483-90.

145. Impaired vasoreactivity in bodybuilders using androgenic anabolic steroids. Lane HA et al. Eur J Clin Invest 2006 Jul: 36(7): 483-8.

146. The more effective immune system of women against infectious agents. Müller HE, Wen Med Wochenschr. 1992:142(17):389-95

147. Sex hormones, immune responses, and autoimmune diseases. Mechanisms of sex hormone action. Ansar AS et al. Human Repr Upd. 11(4) 411-23. 2005.

148. Sex hormones and the immune response in humans. A Bournan et al. Human Reprod Update 11(4) pp.411-23, 2005

149. Effect of a single administration of testosterone on the immune response and lymphoid tissue in mice. Fuji H et al. Immunology 20:315-26, 1975.

150. Estrogens and health in males. Lombardi G. Zarrilli S et al. Mol Cell Endocrinol. 2001 Jun 10:178/1-2):51-5.

151. Dichotomy of glucocorticoid action in the immune system Asadullah K, Schicke H, Cato AC. Trends Immunol. 2002 Mar;23(3):120-2

152 Anabolic steroid effects on immune function: differences between analogues. Mendenhall CL, Grossman CJ, et al. J Steroid Biochem Mol Biol. 1990 Sep:37(1):71-6.

153. A trial of testosterone therapy for HIV-associated weight loss. Coodley GO, Coodley MK. AIDS. 1997 Sep;11(11):1347-52

154. Effects of nandrolone decanoate compared with placebo or testosterone on HIV-associated wasting. J Gold, 1 MJ Batterham et al. HIV Medicine (2006).

155. The effect of anabolic steroids and strength training on the immune response. L Calabrese et al. Med and Sci in Sports and Exer. 21(4) pp. 386-92, 1983

156. Androgens potentiate the effects of erythropoietin in the treatment of anemia of end-stage renal disease. Ballal SH. Dometo DT, Polack DC, Marciulonis P, Martin KJ, Am J Kidney Dis. 1991 Jan:17(1):29-33.

157. Use of androzens in patients with renal failure. Johnson CA. Semin Dial. 2000 Jan-Feb;13(1):36-9.

158. Anabolic effects of nandrolone decanoate in patients receiving dialysis: a randomized controlled trial. Johansen KL, Mullian K, Schambelan M, JAMA

1999 Apr 14:281(14):1275-81.

159. Testosterone metabolism and replacement therapy in patients with end-stage renal disease. Johansen KL, Semin Dial. 2004 May-Jun:17(3):202-8.

160. The effect of anabolic steroids on the gastrointestinal system, kidneys, and adrenal glands. Modlinski R, Fields KB Curr Sports Med Rep. 2006 Apr 5(7):104-9

161. Wilms tumor in an adult associated with androgen abuse. Prat J. Grav GE. Stolley PD. Coleman JW JAMA. 1977 May 23-227(21) 2222 2

162. Anabolic steroid abuse and renal cell carcinoma. Martorana G, Concetti S, Manferrari F, Creti S J Urol. 1999 Dev:162(6):2089

163. Anabolio steroid abuse and renal-cell carcinoma. Bryden AA, Rothwell PJ, O'Reilly PH. Lancet. 1995 Nov 11/2/6/2025/1206 2

164. At what price, alory? Severe cholestasis and acute renal failure in an athlete abusing stanozolol Voshida FM, Karim MA, Shakh JF, Scos JG, Eth SR. CMA1 1994 Sen 15:151(6):791.3

165. Severe cholestasis with kidney failure from anabolic steroids in a body builder. Habscheid W. Abele U. Dahm HH. Dtsch Med Wochenschr. 1999 Sen

166. The incidence of post-operative renal failure in obstructive jaundice. Dawson JL, Br J Surg 1965: 52: 663-665.

167. Acute renal failure complicating muscle crush injury. Abassi ZA, Hoffman A, Better OS. Semin Nephrol. 1998 Sm 18(5):558.65

168. Rhabdomyolysis in a bodybuilder using anabolic steroids. Hageloch W, Appell HJ, Weicker H. Sportverletz Sportschaden, 1988 Sep:2(3):122-5.

169. Atraamatic rhabdomvolysis in a 20-year-old bodybuilder. Morocco PA. J EmergNurs. 1991 Dec;17(6):370-2.

170. Continuous veno-venous hemofiltration for the immediate management of massive rhabdomyolysis after falminant malianant hyperthermia in a bodybuilder. Scherk MR, Beck DH, Nolte M, Kox WJ, Anesthesiolosy, 2001 Jun:94(6):1139-41.

171. Rhabdomvolvsis in a bodybuilder using steroids. Duniels JM, van Westerloo DJ, de Hon OM, Frissen PH. Ned Tildschr Geneeskel 2006 May

172. Hypertension and the kidney. Hohenstein K, Watschinger B. Wien Med Wochenschr. 2008;158(13-14):359-64

173. Henutic effects of 17 alpha-alkylated anaboli-androgenic steroids. HIVHotline. 1998 Dec;8(5-6):2-5

174. Methyltestosterone, related steroids, and liver function. A deLorimier et al. Arch Inter Med Vol. 116 Aug 1965 n 789,94

p 207-94 175: Jaundice associated with norbolethone (Nilevar) therapy. Shaw R K et al. Ann Intern Med 52:428-34 1960.

176. Androaenic/Anabolic steroid-induced toxic hepatitis. Stimac D, Milic S, Dintinjana RD, Kovac D, Ristic S J Clin Gastroenterol. 2002 Oct;35(4):350-2

177- Review of oxymetholone: a 17alpha-alkylated anabolic-androgenic steroid. Pawlatos AM, Fultz O, Monberg MJ, Vootkar A, Pharmd. Clin Ther. 2001 Jun;23(6):789-801; discussion 771

178 A pilot study of anabolic steroids in elderly patients with hip fractures. Sloan JP,WingP, Dian L,Meneilly GS. LAm Geriatr Soc. 1992 Nov:40(11):1105, 11

179. Effects of long-term use of testosterone enanthate. II. Effects on lipids, high and low density lipoprotein cholesterol and liver function parameters. Tvati A. Rajalakshmi M. et al. Int J Androl. 1999 Dec;22(6):347-55

180. Cellular distribution of androgen receptors in the liver. Hinchliffe SA, Woods S, Gray S, Bart AD. J Clin Pathol 1996 May: 49(5):418-20

181. Liver toxicity of a new anabolic agent: methylmienolone (17-alpha-methyl-4,9,11-estratriene-17 beta-ol-3-one). Kruskemper, Noell. Steroids. 1966 Jul;8(1):13-24

182. T. Feyd-Cabanes. Compt. Rend. Soc. Biol. 157, 1428 (1963)

183. Anabolic-androaenic steroids and liver injury. M Sanchez-Osorio et al. Liver International ISSN 1478-3223 p.

184. Androgenic/Anabolic steroid-induced toxic hepatitis. Stimac D et al. J Clin Gastroenterol. 2002 Oct;35(4):350-

185. Mechanisms and sites of action of ursodeoxycholic acid in cholestasis. Beaers U. Nat Clin Pract Gastroenterol Hepatol 2006; 3:318,28

186. Peliosis hepatis in a young adult bodybuilder. Cabasso A. Med Sci Sports Exerc. 1994 Jan;26(1):2-4.

187. Bleeding esophageal varices associated with anabolic steroid use in an athlete. Winwood PJ et al. Post-Grad Med 1 1990 (6:864.65

188. Benipp liver-cell adenoma associated with long-term administration of an androgenic-anabolic steroid (methandienone). Hernandez-Nieto L. Bruauera M. Bornbi J. Carnacho L. Rozman C. Cancer. 1977 Oct:40(4):1761-

189. Henatocellular carcinoma in the non-cirrhotic liver. Evert M, Dombrowski F. Pathologe. 2008 Feb;29(1):47-52 190. Intratesticular biomyosancema in a young man after hish dose doning with Oral Turinabal: a case report Cancer. Froehner M, Fischer R, Leike S, Hakenberg OW, Noack B, Wirth MP. 1999 Oct 15:86(8):1571-5

191. Hepatocellular carcinoma associated with recreational anabolic steroid use. Gonyski P, Thompson CH, Sabhash HS, Thomas AC, Br J Sports Med. 2008 Jan:42(1):74-5: discussion 75.

192. Bodybuilder death steroids warning. Express and Star 2008 09/04. Epub. www.expressandstar.com

193 Effect of testosterone and anabolic steroids on the size of sebaceous glands in power athletes.Kiraly CL et al.Am J Dermatoriathol, 1987 Dec. 9.6, 515-9.

194. RU 58841, a new specific topical antiandrogen: a candidate of choice for the treatment of acne, androgenetic alopecia, and hirsutism. Battmann T. et al. J Steroid Biochem Mol Biol. 1994 Jan;48(1):55-60

195. Androgenetic alopecia and current methods of treatment. Bienová M, Kucerová R. et al. Acta Dermatovenerol Alp Panonica Adriat, 2005 Mar;14(1):5-8.

¹⁹⁶. Molecular mechanisms of androgenetic alopecia. Trüch RM. Exp Gerontol. 2002 Aug-Sep;37(8-9):981-90. 197. The inheritance of common haldness: two B or not two B? Kuster W, Happle R. J Am Acad Dermatol 1984; 11:921-26

198. Polymorphism of the Androren Receptor Gene is Associated with Male Pattern Baldness. Justine A Ellis, Margaret Stebbing and Stephen B Harrap. Journal of Investigative Dermatology (2001) 116, 452-455.

199. EDA2R is associated with androgenetic alopecia. Prodi DA, Pirastu N, et al. J Invest Dermatol. 2008 Cas-176/0/7569 70 Each 2009 Aug 2

309,128(9).2288-70. Epub 2008 Apr 3.

 Current understanding of androgenetic alopecia. Part I: Etiopathogenesis. Hoffmann R, Happle R. European Journal of Dermatology. Volume 10, Number 4, 319-27, June 2000

201- Estrogen and progesterone receptors in androgenic alopecia versus alopecia areata. Wallace ML, Smoller BR. Am J Demutopathol. 1998 Apr;20(2):160-3.

2022. Hormonal doping and androgenization of athletes: a secret program of the German Democratic Republic government. Funke WW, Beendork B. Cin Chern. 1997 Jul;45(7):1262-79.

203. Role of estrogen on bone in the human male: insights from the natural models of congenital estrogen deficiency. Fastini-Fustini. M. et al. Md Cell Endocrinol. 2001 Jun 10:178/1-21:215-20.

204. Effects of estrogen on growth plate senescence and epiphyseal function. M Weise, S De-Levi et al. Proc Natl Acad Sci June 5,2001 pp. 6871-6876.

205. The results of short-term (6 months) high-dose testosterone treatment on bone age and adult height in boys of excessively tall stature. Brimswig JH, von Lengerke HJ et al Eur J Pediatr. 1988 Nov;148(2):104-6.

206. Orandrolone in constitutional delay of growth: analysis of the growth patterns up to final stature. Bassi F, Neri AS, Gheri RG, Cheli D, Serio M. J Endocrinol Invest. 1993 Feb;16(2):133-7.

 Pectoralis maior tendon rupture and anabolic steroids in anarmesis—a case review. Milez J, Novotn⁺ P. Rozhl Chir. 2008 Jul;87(7):380-3.

 Saccessive ruptures of patellar and Achilles tendens. Anabolic steroids in competitive sports. Isenberg J, Prokop A, Skoums E. Unfallchiurg. 2008 Jar;111(1):46-9

 Biomechanical responses of different rat tendoms to nandrolone decanoate and load exercise. Marqueti RC, Prestes J, WangCC, et al. Scand J Med Sci Sports. 2010 Jul 29.

210. The effect of local use of nandrolone decanoate on rotator cuff repair in rabbits. Papaspiliopoulos A, Papaparaskeva K, Papadopoulou E, et al. J Invest Surg 2010 Aug23(4):204-7.

211. Self-epoch and bio-androgenic stroub use and masculosketal injuries: findings from the center for the study of retired athletis health survey of retired NFL players. Hom S, Gregory P, Guskiewicz KM. Am J Phys Med Rehabil. 2009 Mar;98(3):192-200.

212. Salt, hypertension, and edema. Rössler R. Internist (Berl). 1976 Oct;17(10):489-93. Review.

213. Sex hormone effects on body fluid regulation. Stachenfeld NS: Exerc Sport Sci Rev. 2008 Jul;36(3):152-9.

214. Effect of ovarian steroids on vasopressin secretion. Forsling, M. L., P. Stromberg and M. Akerhand, J. Endocrinol. 95: 147-151, 1982

215. Estrogen influences osmotic secretion of AVP and body water balance in postmenopausal women. Nina S. Stacherdeki, Loretra Dipietro, Steven F. Patter, and Ethan R. Nadel Am J Physiol Regal Integr Comp Physiol 274: RI37:R195, 1998.

216. Independent and combined effects of testosterone and growth hormone on estracellular water in hypoplituitary men. Johannsson G, Gibney J, et al. J Clin Endocrinol Metab. 2005 Jul/90(7):3989-94. Epub 2005 Apr 12.

217. Casner, S. W., Early, R. G., and Carlson, B.R. Journal of Sports Med and Phys Fitness, 1971 11,98.

218. The effects of anabolic steroids on growth, body composition, and metabolism in boys with chronic renal failure on regular herrodialysis. Jones RW, El Bishri MM et al. J Pediarr. 1980 Oct;97(4):559-66.

219. A madenized, placebo-controlled trial of nandrolone decanoate in human immunodeliciency virus-infected men with mid to medenate weight loss with recombinant human growth hormone as active reference treatment. Storer TW, Woodhouse LJ, J Clin Endocrinol Metah. 2005 Aug/08(9):4174-82. Epub 2005 May 23.

220. Bodyhaiders' Body Composition: Effect of Nandrokne Deamone. VAN MARKEN LICHTENBELT, W. D., F. HARTGENS, N. B. J. VOLLAARD, S. EBBING, and H. KUIPERS. Med. Sci. Sports Esere, Vol. 36, No. 3, pp. 484-489, 2004.
231. Soore larvatiris following chronic ambolic steroid abuse. Ray S. Masood A. Pickles J. Moarmolidis L. J.

221. Severe laryngitis following chronic anabolic steroid abuse. Ray S, Masood A, Pickles J, Mournoulidis L J Laryngol Otol. 2008 Mar;122(3):230-2. Epub

2007 May 14

222 Estrogen regulation of mammary gland development and broast cancer: amphiregalin takes center stage Heather L LaMarcal and Jeffrey M Rosen. Broast Cancer Res. 2007; 9(4): 304.

223. Androgens and mammary growth and neoplasia. Dimitrakakis C, Zhou J, Bondy CA. Fertil Steril. 2002 Apr;77 Suppl 4:252-33.

224. Surgical treatment of gynecomastia in the body builder. Alache AE. Plast Reconstr Surg. 1989 Jan;83(1):61-6.

225. Roles of estrogen and progesterone in normal mammary gland development insights from progesterone receptor nall matant mice and in situ localization of receptor. Shyamala G. Trends Endocrinol Metab. 1997 Jan-Feb/8[1]:34-9.

226. A report on alterations to the speaking and singing voices of four women following hormonal therapy with virilizing agents. Baker J. J Voice 1999 Dec;13(4):496-507.

227. Fundamental voice frequence during normal and abnormal growth, and after androgen treatment. Massenkoski V, Lenko HL, Tjernhard P, Vuorenkoski L, Perheentupa J. Arch Dis Child. 1978 Mar;53(3):201-9.

228. Fluosymesterone therapy in anemia of patients on maintenance hemodialysis: comparison between patients with kidneys and anephric patients. Acchiardo SR, Black WD. J Dal. 1977;1(4):357-66.

229. Testoterone therapy in women: its role in the management of hypoactive sexual desire disorder. Abdallah RT, Simon JA. Int J Impot Res. 2007 SepOct;19(5):458-63. Epub 2007 Jun 21.

230 Mrilization caused by methandrostenolone-containing cream in 2 prepubertal girls. Sorgo W, Zachmann M. Helv Paediatr Acta. 1982 Sep;37(4):401-6.

231. Change in speaking fundamental frequency in homene-treated patients with Turner's syndrome-a longitudinal study of four cases. AnderssonWallgren G, Albertsson-Wildard K. Acta Paediar. 1994 Apr;83(4):452-5.

232. Valization of the voke in post-menopassal women due to the anabolic steroid nandrolone decanoate (Deca-Dambolin). The effects of medication for one year. Genitsmu EJ, Buocaar MP, Hakkesteegt MM, Birkenhüger JC. Cin Ocelaryagal Alliad Sci. 1994 Petryl 81,758-84.

233. Idiopathic isolated clitoromegaly: A report of two cases. Enay Copcu1, Alper Aktas et al. Reproductive Health 2004, 1:4.

²³⁴. Two Cases of Citoromegily. Citoral Reduction Preserving Sensation of Citoris. NODA KOJIRO (Chiha-ken Kodomo Byoin, UDAGNWAAKIKAZU (Chiha-ken Kodomo Byoin) et al. Journal of Japan Society of Aesthetic Plastic Sargery VOL22NO.37Ac6E9.995(2000).

235. Evolution and Tourmont of Bluman with Elevation. METICCA ELEBATED M.D. and DETED 1. CADER

CARDENE HILLING HILLING IN MUCH MANY PROVIDENT AND LONG TO THE VERY AND AN AVAILABLE. M.D. Am Fam Physician 2003;67:2565,72

236. Effects of lona-term androgen administration on breast tissue of female-to-male transsexuals. M Slagter, L Goorn et al. L'Histochem Cytochem 54/87 905,910,2006

237. Bolowioural effects of androaen in men and women. Christiansen K. J Endocrinol. 2001 Jul;170(1):39-48.

238. Experiments testostemme enhances responsiveness to social threat in the neural circuitry of social appreciation in Exogenous residence entrances responsiveness to social mean in the neurin circuitry of social aggressic humans. Hermans FI. Ramsev, NF, van Hank I. Biol Powhiatry. 2008 Feb 1:63(3):763-70. Fruib 2007 Aug 28.

239. Undewround Steroid Handbook II. Daniel Duchaine. 1989. HLR technical books. Venice, CA.

240. Metabolic and behavioral effects of high-dose, exogenous testosterone in healthy men. Bagatell CJ, Heiman JR, Metabole and belaviour effects of high-case, outgetous residence in reality field Matsumoto AM Rivier IF Reemer WI 1 (Tin Endocrino) Metab 1994 Aug 29(2):561,7

241. The effects of exogenous testosterone on sexuality and mood of normal men. Anderson RA, Bancroft J, Wa EC 1 Clin Endorrinol Match 1992 Day 75(6):1502 7

242. Psychological and serum homovanillic acid changes in men administered androgenic steroids. Harnan CJ Jr, Friedl KE. Zold A. Kettler TM, Plymate SR. Psychoneuroendocrinology. 1991;16(4):335-43.

243. Powhoostaal effects of three doses of testosterone cycling in normal men. Yines WR et al. Biol Psychiatry. 1000-15-251 (0)

244. Effects of supraphysiological doses of testosterone on mood and aggression in normal men. H Pope, E Kouri et al. Arch Gen Psychiatry, 2000;57:133-140.

245. Psychiatric side effects induced by supraphysiological doses of combinations of anabolic steroids correlate to the severity of abuse. T Pagonis et al. Eur Psych 21 (2006) 551-62.

246. Unlower toward women and illicit androgenic-anabolic steroid use. Choi PY, Pope HG Jr. Ann Clin Psychiatry, 1994 Mar.6(1):21-5.

247. Criminality among individuals testing positive for the presence of anabolic androgenic steroids. Klötz F, Garle M, Granath F, Thiblin I. Arch Gen Psychiatry. 2006 Nov;63(11):1274-9.

248. Homicide and near-homicide by anabolic steroid users. Pope HG Jr, Katz DL. J Clin Psychiatry. 1990 Janc51(1):28-31.

249. Evidence for physical and psychological dependence on anabolic androgenic steroids in eight weight lifters. K Bower G Flionelos et al. Am J Psychiatry 1990; 147:510,12

²⁵⁰ Muscle dysmorphia. An underrecognized form of body dysmorphic disorder. Pope HG Jr, Gruber AJ, Choi P, Olivardia R. Phillips KA. Psychosomatics.

1997 Nov-Dec:38(6):548-57.

^{251.} "Chocolate addiction": a preliminary study of its description and its relationship to problem eating. Hetherinaton MM, MacDiarmid JI. Appetite. 1993 Dec;21(3):233-46

252. Reinforcing aspects of androgens. Wood RI. Physiol Behav. 2004 Nov 15;83(2):279-89.

253. Expression of testosterone conditioned place preference is blocked by periphenal or intra-accumbens injection of alpha-flupenthiol. Packard MG. Schroeder JP, Alexander GM, Horn Behav, 1998 Aug;34(1):39-47.

254. Role of donamine receptor subtypes in the acquisition of a testosterone conditioned place preference in rats. Schroeder JP, Packard MG. Neurosci Lett. 2000 Mar 17:282/1-2):17-20

255. Increased dopamine transporter density in the male rat brain following chronic nandrolone decanoate administration, Kindhadh AM, Rahman S, Lindblom J, Nyberg F, Neurosci Lett, 2004 Feb 12:356(2):131-4.

256. Abuse liability of testosterone. Fingehood MI, Sullivan JT et al. J Psychopharmacol 1997; 11(1):59-63.

257. Anabolie steroid withdrawal depression: a case report. Allnutt S, Chaimowitz G. Can J Psychiatry. 1994 Jun:39(5):317.8

258. The use of fluoretine in depression associated with anabolic steroid withdrawal: a case series. Malone DA Jr, Dimeff RJ. J Clin Psychiatry. 1992 Apr;53(4):130-2.

259. New-generation antidepressants, suicide and depressed adolescents: how should clinicians respond to changing evidence? Dudley M, Hadzi-Pavlovic D, Andrews D, Perich T, Aust N Z J Psychiatry, 2008 Jun 42(6):456-66.

260. Anabolic androgenic steroids and suicide. Thiblin I, Raneson B, Rais J, Ann Clin Psychiatry, 1999 Dec:11(4):223-31

261. Testosterone deficiency and mood in aging men: pathogenic and therapeutic interactions. Seidman SN. World J Biol Psychiatry. 2003 Jan;4(1):14-20

262. Treatment strategies of withdrawal from long-term use of anabolic-androgenic steroids. MedraÉ M, Tworowska U. Pol Merkar Lekarski. 2001 Dec:11(66):535-8. Review

263. Psychological moods and subjectively perceived behavioral and somatic changes accompanying anabolicandrogenic steroid use. Bahrke MS,Wright JE, Strauss RH, Catlin DH. Am J Sports Med. 1992 Nov-Dx:20(6)717.24

264. Insormia: pathophysiology and implications for treatment. Roth T, Rochrs T, Pies R. Sleep Med Rev. 2007 Feb:11(1):71-9. Epub 2006 Dec 18.

265. When does estrogen replacement therapy improve sleep quality? Polo-Kantola P, Erkkola R, Helenius H, Irjala K, Polo O. AmJ Obstet Gynecol. 1998 May;178(5):1002-9.

266. Adverse effects of anabolic androgenic steroids on the cardiovascular, metabolic and reproductive systems of anabolic substance abusers. Tuorno Karila, Publications of the National Public Health Institute ISBN 951-740-388-2

267. Reversible hypoxonadism and azoospermia as a result of anabolic-androgenic steroid use in a bodybuilder with personality disorder. A case report. Boyadjiev NP, Georgieva KN, Massaldjieva RI, Gueorgaiev SI. J Sports Med Phys Fitness. 2000 Sep;40(3):271-4.

268. Conservative management of azoospermia following steroid abuse. M.R. Gazvani et al. Human Reprod 12(8) (1997) no. 1706-08

269. Contraceptive efficacy of testosterone-induced azoospermia in normal men. Lancet, 1990 20;336(8721):955-9.

270. Effects of chronic testosterone administraton in normal men: saferty and efficacy of high dose testosterone and parallel dose-dependent suppression of lateinizing hormone, follicle-stimulating hormone, and sperm production. Matsumoto AM. J Clin Endocrinol Metab 1990; 70:282-87.

271. Restorative increases in serum testosterone levels are significantly correlate to improvements in sexual functioning A Seftel, R Mack et al. J Androl 25(6) 2004 pp.963-72.

200 2009 pp. 200 22

*** Sexual functioning of male anabolic steroid abusers. Moss HB, Panzak GL, Tarter RE. Arch Sex Behav. 1993 Feb;22(1):1-12.

273. Low see hormone-binding globalin and testosterone levels in association with erectile dysfunction among human immanodeficiency virus-infected men receiving testosterone and oxandrolone. Wasserman P, Segal-Maarer S, Rabin D J, Sex Med. 2008 Jure(1):2041-7. Epub 2007 Oct 24.

274. Of mice and men: the many guises of estrogens. Simpson ER, Jones ME. Ernst Schering Found Symp Proc. 2006;(1):45-67.

²⁷⁵ Testosterone-induced priapism in Klinefelter syndrome. Ichioka K, Utsanomiya N, Kohei N, Ueda N, Inoue K, Teni A.Uroloov. 2006 Marx57(3):622 e17–8. Evub 2006 Feb 28.

276. Severe priapism as a complication of testosterone substitution therapy. Zelissen PM, Stricker BH. Am J Med. 1988 Aug85(2):273-4

277. Testosterone induced priapism in two adolescents with sidde cell disease. Slayton W, Kedar A, Schatz D. J Pediatr Endocrinol Metab. 1995 JulSep;8(3):199-203.

278. Benim hypertrophy and carcinoma of the prostate. Moore RA. Surgery 1944;16:152-67.

279. Guilt by association: a historical perspective on Huggins, testosterone therapy, and prostate cancer. Morganialer A. J See:Med. 2008 Aug;5(8):1834-40. Epub 2008 Jun 10.

280. Testosterone therapy in hypogenadal men and potential prostate cancer risk: a systematic review. Shabsigh R, Crawford ED, Nehra A, Slawin KM. Int J Impor Res. 2008 Jul 17. [Epub shead of print]

281. Long-term psychiatric and medical consequences of anabolic-androgenic steroid abuse: A looming public health concern? Kanayama G, Hudson JI, Pope HG Jr. Drug Alcohol Depend. 2008 Nov 1;98(1-2):1-12. Epub 2008 Jul 2.

282. The many faces of testosterone. Bain J. Clin Interv Aging. 2007;2(4):567-76.

283. Testosterone treatment in hypogonadal men: prostate-specific antigen level and risk of prostate cancer. Guay AT, Perez JB, Fitalhi WA, Vereb M. Endocr Pract. 2000 Mar-Apr;6(2):132-8.

284. Adenocarcinoma of prostate in 40 year old body-builder. Roberts TJ, Essenhigh DM. Lancet 1986;2:742.

 Testosterone therapy for men at risk for or with history of prostate cancer. Morgentaler A. Curr Treat Options Oxeol. 2006 Sep;7(5):363-9.

266. Testosterone replacement therapy and the risk of prostate cancer. Is there a link? Barqawi A, Crawford ED. Int J Impor Res. 2006 Jul-Aug 18(4):323-8. Epub 2005 Nov 10.

 Warne change of the prostate and seminal vesicles in male hypogenadism after androgen replacement therapy. Sissagova I, Nakada T, Kazaran T, Satomi S, Tenah T, Katayaran T. Int Urol Nephrol. 1990;22(3):279-84.

288. A four-year efficacy and safety study of the long-acting parenteral testosterone undcarnone. Minnernam T, Schubert M, Hühler D, Goani-Berthold I, Freade S, Schumram C, Oettel M, Ernst M, Mellinger U, Sommer F, Krone W, Jockschole F. Angin Mule. 2007 Sept. 1003;1155-8.

289. Effect of Testostorone Replacement Therapy on Prostate Tissue in Men With Late-Onset Hypogenadism A Randomized Controlled Trial Leonard S. Marks, MD; Norman A. Mazer, MD, et al. JAMA. 2006;296:2351-2661.

290. Prognostic value of serum markers for prostate cancer. Stemman UH, Abrahamsson PA, Aus G, Lilja H, Banarna C, Handw FC, Boccon-Gibod L, Ekama P, Scand J Urol Newhool Sarod. 2005 Mar/1216/654-81.

 Relationship between prostate specific antiger, prostate volume and age in the benign prostate. Collins GN, Lee RJ, McKelvie GB, Rogers AC, Hehir M. Br J Urol. 1993 Apr;71(4):445-50.

292. Long-term treatment with finateride results in a clinically significant reduction in study prostnet volume compared to placebo over the full range of baseline prostate sizes in next enrolled in the MTOPS trial. Kaplan SA, Reachtern CG/McContell JD et al. Ubol. 2008 Sept. [90(6))(3103-2). September 30, Papir 2008 Jul 17.

293. Winning is the Only Thing, Randy Roberts, James S. Olson, JHU Press, 1991. ISBN 0801842409

294. Androgen or Estrogen Effects on Human Prostate. B. Jin, L. Turner et al. J Clin Endocrinol Metab. 1996;81(12):4290-95.

295. Fingerprinting the diseased prostate: associations between BPH and prostate cancer. Shah US, Getzenberg RH. J Cell Blochem. 2004 Jan 1;91(1):161-9.

296 Estrogen-regulated development and differentiation of the prostate. McPherson SJ, Ellem SJ, Risbridger GP. Differentiation. 2008 Jul;76(6):660-70. Epub 2008 Jun 28.

297. Estrogen action on the prostate gland: a critical mix of endocrine and paracrine signaling. G Risbridger, S Ellem et al. J Mol Endocrinol (2007) 39, 183–88.

 Effects of androgen therapy on prostatic markers in hemodialyzed patients. Teruel JL, Aguilen A, Avila C, Otrafo J, Sand J Urol Nephrol. 1996 Apr;30(2):129-31.

299. The role of aromatization in testosterone supplementation. Effects on cognition in older men. M.M. Cherrier, A.M. Matsumoto et al. Neurology

2005;64:290-96

 Use of anabolic steroids and associated health risks among gay men attending London gyms. Bolding G, Sherr L, Elford J. Addiction. 2002 Feb;97(2):195-203.

 Indications of prevalence, practice and effects of anabolic steroid use in Great Britain. Kotkia P, Stimson GV. Int J Sports Med. 1997 Oct;18(7):557-62.

302. A combined regimen of cyproterone acetate and testosterone enanthate as a potentially highly effective male contraceptive. C Meriggola et al. J Clin Endocrinol Metab 81(8) 3018-23, 1996.

303. HPGA normalization protocol after androgen treatment. N Vergel, AL Hodge, MC Scally. Program for Wellness Restoration.

304. An update to the Crisler HCG Protocol. John Crisler, DO. 2004.

305. Testicalar responsiveness following chronic administration of hCG (1500 IU every six days) in untreated hypogenadotropic hypogenadism, Baldacci R, Toscano V, Casilli D, Maroder M, Sciarra F, Boscherini B. Horn Metab Res. 1997 Mary15(5):216-21.

306 Estrogen suppression in males: metabolic effects. Mauras N, O'Brien KO, Klein KO, Hayes V J Clin Endocrinol Metab. 2000 Jul/85(7):2370-7.

307. Progesterone and testosterone in combination act in the hypothalarnas of castrated runs to regulate the secretion of LH.Turner AI,Tilbrook AJ, Clarke IJ, Scott CJ. J Endocrinol. 2001 May;169(2):291-8.

308. Football; Alzado Tumor is Rare and Deadly. Elisabeth Rosenthal. NY Times. July 4, 1991.

309. Primary central nervous system lymphoma. O'Nelli BP, Illig JJ. Mayo Clin Proc. 1989 Aug64(8):1005-20.

310. Sports People: Football; Alzado Talks. NY Times. Associated Press Report. June 28,1991.

311. Alzado believed drus killed him-Ex-Raider star dead from brain cancer. Associated Press. May 15, 1992.

312 54-year-old man with breast cancer after prolonged testosterone therapy. Sorscher S, Krause W. Clin Adv Hernatol Oncol. 2005 Jan;2(6):475; discussion

476.

Androgen therapy, Longson D. Practitioner. 1972 Mar;208(245):338-48.

314. Androgen treatment of middle-aged, obese merc effects on metabolism, muscle and adipose tissues. Milrin P, Krotkiewski M, Björntorp P. Eur J Med.

1992 Oct;1(6):329-36.

1315: Effects of androgen therapy on adjoose tissue and metabolism in older men. Schroeder ET, Zheng L, Ong MD, Martinez C, Flores C, Stewart Y, Azen C, Sattler FR, J Clin Endocrinol Metab. 2004 Oct;89(10):4863-72.

316. Insulin sensitivity, insulin secretion, and abdeminal fat: the insulin resistance atherosclerosis study (IRAS) family study. Wagerknecht LE, Langerfeld CD et al. Diabetes 52:2490-2494.

 Recent developments in the toxicology of anabolic steroids. Graham S, Kennedy M. Drug Saf. 1990 Nov-Dec5(6)(458-76.

Jack (1990) - ... 318. Insulin resistance and diminished glucose tolerance in powerlifters ingesting anabolic steroids. Cohen JC, Hickman R. J Clin Endocrinol Metab. 1887 May;64(5):960-3.

319. Insulin action and dynamics modelled in patients taking the anabolic steroid methandienone (Dianabol). Godsland IF, Shenran NM, Wym V. Clin Sci (Lond). 1986 Dec;71(6):665-73

320. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Singh AB, Hsia S, et al. J Clin Endocrinol Metab. 2002 Jan;87(1):136-43.

321. Nandrolone, a 19-nortestosteore, enhances insulin-independent glucose uptake in normal men. Hobbs CJ, Jones RE, Plymate SR, et al. J Clin Endocrinol Metab. 1996 Apr;81(4):1582-5.

222. Indications of prevalence, practice and effects of anabolic steroid use in Great Britain. Korkia P, Stimson GV. Int J Sports Med. 1997 Oct;18(7):557-62.

 Castionascular complications of respiratory diseases. Chowdhuri S, Crook ED, Taylor HA Jr, Badr MS. Am J Med Sci. 2007 Nov;334(5):361-80.

324. Obssity and hormonal factors in sleep and sleep apnea. Wittels EH. Med Clin North Am 1985 Nov;69(6):1265-80.

325. Metabolic aspects of sleep apnea. Grunstein RR. Sleep. 1996 Dec;19(10 Suppl):S218-20.

326. Testosterone replacement therapy for older men. Borst SE, Mulligan T. Clin Interv Aging 2007;2(4):561-6.

327. The Short-Term Effects of High-Dose Testosterone on Steep, Breathing and Function in Older Men Peter Y. Liu, Breadon Yee et al. The Journal of Clinical Endoerinology & Metabolism Vol. 88, No. 8 3605-3613

328. Induction of the obstructive sleep apnea syndrome in a woman by exogenous androgen administration. Johnson MW, Anch AM, Remmers JE, Am Rev Respir Dis. 1984 Jun;129(6):1023-5.

Acute Steroid Safety: Studies with Real-World Dosages

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The commits dateid user will usually point on the fact that a review of the medical licentre over the part 50 years will also with the overall staty prodied of these duggs has been quite forwards. Statesid approximate, not the there hand point out that an filtra user takes the state of the states of the darger than the patients using the multi box in given by the states in the lack of particular medical states. Moreal and the multi plant has above, conformed in the states with an above multimatic states of the medical states with an above multi-state state states are the states. Moreal and the multi-states with an above multi-states are the states of the the multi-states above of the medical states with an above multi-state states are the states of the states and the states of the medical states with an above multi-states and the states of the states and the states of the medical states with an above multi-states and the states of the statest and bas end backs of the statest constructions.

In this section, we carrine three medical studies that appear highly relevant for examing networds actue amphoicamlogene stored safety. They concern of thempeteic does, but a supathempetic level and duration of findse that any first stored user would recogate as a difficult for improving mucks may, attending and performance. In fact, the doegas and aggressive stored-insign body huddens and storeght athletes. A fairly comprehensive set thanh murkers were assessed during theme three investments, relating its stored in sensitive. serum cholesterol and triglyceride, prostate specific artigen (PSA) levels, and liver enzymes. Because of the protocols that were used, these studies give us a fairly good basis to evaluate the negative health impact of anabolic/androgenic steroids, at least as it relates to an isolated cycle.

600 mg/wk of Testosterone

The first is a testosterone dose-response study published in the American Journal of Physiology Endocrinology and Metabolism in July of 2001, which looked at the effects of various doses of testosterone enanthate on body composition, muscle size, strength, power, sexual and cognitive functions and various markers of health 329 61 normal men ages 18-35, participated in this investigation. They were divided into five groups, with each receiving weekly injections of 25-50-125-300 or 600 millionants for a period of 20 weeks. This treatment period was preceded by a control (no drug) period of 4 weeks, and followed by a recovery period of 16 weeks. Markers of strength and lean body mass gains were the oreatest with larger doses of testosterone, with the 600 mg group gaining slightly over 17 pounds of fat-free mass on average over the 20 weeks of steroid therapy. There were no significant changes in prostate-specific antigen (PSA) liver enzymes (liver stress) sexual activity, or cognitive functioning at any dose. The only negative trait noted was a slight HDL (good) cholesterol reduction in all groups except those taking 25 mg. The worst reduction of 9 points was noted in the 600 mg group, which still averaged 34 points after 20 weeks of treatment. All groups, except this one, remained in the normal reference range for males (40-59 noints)

600 mg/wk of Nandrolone

Next we look at a study conducted with HIV+ men which charted the lean-mass-building effects of randrolone decanoate330 30 neonle narticipated in this investigation, with each given the same (high) weekly dose of this drug. Half underwent resistance training so that two groups (trained and untrained) were formed. The dosing schedule was quite formidable, beginning with 200 mg on the first week, 400 mg on the second, and 600 mg for the remaining 10 weeks of neak therany. Doses were slowly reduced from weeks 13 to 16 to withdraw patients slowly from the drug. Potential negative metabolic changes were looked at closely including cholesterol and linid levels (including subfractions of HDL and LDL). trielycerides, insulin sensitivity, and fasting glucose levels. Even with the high dosages used here, no negative changes were noted in total or LDL cholesterol, trighcerides, or insulin sensitivity. In fact, the group also undergoing resistance exercise noticed significant improvements in LDL particle size distribution. Incorrotein(a) levels, and triglyceride values. which all indicate improved cardiovascular disease risk. Carbohydrate metabolism was also significantly improved in this group. The only negative impact noted during this study was a reduction in HDL (good) cholesterol values similar to that noted with the testosterone study. with an 8-10 point reduction noted between both groups.

100 mg/day of Anadrol

Lastly we find a study looking at the notent oral steroid oxymetholone (Anadrol) 331 This steroid is thought to be one of the most dangerous ones around by bodybuilders, who as a group seem to treat it with both a lot of respect and caution. It is not common to find them exceeding the doses and intake durations of this investigation, making it a very good representation of real-world Anadrol usage. This study involves 31 elderly men, between the ages of 65 and 80. The men were divided into three groups, with each taking 50 mg, 100 mg, or placebo daily for a 12-week period. Changes in lean body mass and strength were measured, as well as common markers of safety including total, LDL and HDL cholesterol levels, serum triglycerides, PSA (prostate-specific antizen), and liver enzymes. Muscle mass and strength gains were again relative to the dosage taken, with the end results being similar to those noted with 20 weeks of testosterone enanthate therapy at 125 mg or 300 mg per week (about 6.4 and 12 lb of lean body mass gained for the 50 mg and 100 mg doses respectively). There were no significant changes in PSA, total or LDL cholesterol values, or fasting triglycerides: however, there was a significant reduction in HDL cholesterol values (reduced 19 and 23 points for the 50 mg and 100 mg groups respectively). Liver enzymes (transaminases AST and ALT) increased only in the 100 mg group, but the changes were not dramatic, and were not accompanied by hepatic enlargement or the development of any springs lister condition

Adding It All Up

One handle and tweety-one runn participated in these three statics, which into hold the use of noderate to injudy ones of starshift for profits of there to the remain. Altopoilt in mysshecking in most opposents of analohic starburgness starshift (in the starburgness) and the starburgness of the starburgness of the starburgness of the min negative impacts of starshift is not influence cases was a reduction in good (HTGL), chotsteroit values, which is a highing to cores to assessing one' risk for declering cardinocated fiscase. It is uscernin, however, I a short-foil tensors in the particular risk factor reducts to any tangles durings to core's handlo oue'r the king declering the mathematicate of the starburgness of the starburgness of the starburgness of the mathematicate of the starburgness of

Logie wordl seem to sagget that the isolated use of storoids, nucler parameters similar to those used in these true studies, should match neitherly minimal risks to handh. At the work least, it is externely difficult to argue that an isolated cycle with a moderate dung dose is tattenuout to ploying assimo notient with your body, as mort meficial campians against the use of these drugs would seem to saggest. Bat make no missile. These same study results consistently demonstrating to outstorogers in chosed lips with the doses necessary for ploying or performance enhancement, and uncefre low it is that long-term ambedinationgen stermid abuse cam injure controloscular health.

The Endocrinology of Muscle Growth

The road to anaboke insight must include a biological understanding of what muscle growth actually earths. Other any simplifies by the term protein synthesis," means the edge could be a lightly complex process involving much more than just building proteins from minim casks. Moules hypertraphysic loce courts scientific terms for the source source hiterance build keeled theses. Since this discovery of statistic colls in 1961, a great deal of research arts the mechanism of muscle, loce theory and the statistic coll and the statistic coll and that unlike normal muscle cole, these satellite cole cas be regreared throughout and the mechanism of muscle cole, these satellite cole cas the provide source of the mechanism of muscle cole and wheat lamaged muscle, cole. These satellite cole are three, vasing for enorming to sufficient actionation.

Injury or trauma will provide the stimulus necessary to activate satellite cells. Once activated, they will begin to divide, multiply, and form into myoblasts (myoblasts are essentially donor cells that express myogenic genes). This stage of hypertrophy is often referred to as satellite cell *proliferation*. The muchbasts will then fisse with existing muscle fibers, dorating their nuclei. This stage of the process is usually called differentiation. Skeletal muscle cells are multinucleated, which means they nossess many nuclei. Increasing the number of nuclei allows the cell to regulate more cytoplasm, which allows more actin and muosin, the two dominant contractile proteins in skeletal muscle, to be produced. This increases the overall cell size and protein content of the muscle cell. Incidentally, the number of nuclei in relation to cross-sectional area also helps to determine the fiber type of the cell namely slow twitch (aerobic) or fast twitch (anaerobic)332 333 It is innortant to note that we are not increasing muscle cell number with muscle hypertrophy. We are only increasing cell size and protein content, even though we are using satellite cells to help accomplish this. It is possible for myoblasts to fuse together and actually form new muscle fibers. This is called muscle hyperplasia, and equates to the legitimate growth of new muscle tissue. This is, however, not the primary mechanism of muscle growth in adult life.

^{329.} Testosterone dose-response relationships in healthy young men. Shalender A, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: e1172-81 2001

Physiaa Enddatase structure and characterized and resistance training in men with HIV Sattler FR, Schroeder ET, Dabe MP, Jaque SV, Martinez C, Blanche PJ, Azen S, Krauss RM. Am J Physiol Endocrinol Metab. 283(6) Dec (2002):E121-42: Epide 3020 Aug 27.

^{331.} Effects of an oral androgen on muscle and metabolism in older, community-dwelling men. Schroeder et al. Am J Physiol Endocrinol Metab 284: E120-28

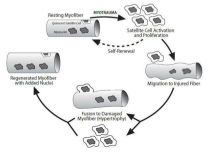
Now that we know what masch hypertrophysis really about, Lris look at anables distinks and ongoing regulation. The following is a number of the chino the formeness and growth factors that mediate mascle growth, from the initiation of dumage, to themeses and growth and growth. For the sale of organization, 11 theme presented them is what I consider to be three logical primes of action. These are not scentifically accepted definitions. Additionally, we coald contine to go detoper and deeper into each of the various composation, messengeness, buding proteins, and reception is moded in the initiate and analogical budged in the state of the loce state of the state of the state of the loce state of t

Trigger

We all uncleasted that weight training is influenceful to growing mack issue. To date, no 'is on your as and gring and inpropril pills been intered. The sensor in that a mather of changes take place in your local macket issues during inframe training that are visit to the growth process. Within these early images, growthis afficial frame prossels to simulate. So for our proposes, we will start have. Training is the "Yingar" in the anabole process. We specifically, it is hockadow chalte manges that weight training powers that on History they proceed with the hockadow chalte manges that weight training powers are will be the process with a typic backet (mange) been excessive to have any specific formation.

Phase I: Initial Response

The hitli Response phase covers those changes in marke chemistry that begin immediately, daring mining, which will be the prosphored for their repair and growth, in more grapher, the hitling Response Phase will corter the potential magnitude of other signals to follow. In the anables process, the phases is a dargared by the relases of anablesine and formanesed code, and the formation of active massengers including mostightmins, cyclukese, substratese, and proceedings in the breakdown of the cost prophorphic type of market, each, which is initiated by the cellular disquiroit of damaging energies.¹¹ Hospitolytic strates the outer hypothese the cellular disquiroit of damaging energies.¹¹ Hospitolytic strates the outer hypothese threaks the bit hospitolity that and of the cortexies movement is of particular profutance here, which is the "negative" part of the its, where the market is strateful and messation.



Fusion to Produce New Myofibers (Hyperplasia)

MUSCLE HYPERTROPHY AND THE 4 STAGES OF THE SATELLITE CELL CYCLE

During the Activation stage, dormant satellite cells are stimulated to enter the cell cycle. Proliferation marks the formation of new myohasts (active donor cells). These myohasts will face with existing damaged macke filters during the Difformation places. This allows for greater protein synthesis and the expansion State (Structure) and the state of the state of the state of the state state (Structure) and the state of the state of the state of the state of the state militator of market growth, is believed to be a key regulation that is stage.^{333,35}

The moure of anchiclonic acid, which is the certanl bioactice lipid in the analosis process. Biornal of Hingys control what coccas thereing its physics. Anchicles card is converted leading and immediately via enzymes is a narrher of archice analosk, and produced the model of which its terms of marked ground the genesizations, the most methy of corrol marked ground the genesizations in the genesizations, markey will coronid marked ground biological strategies and the strategies of the model of the genesization of the strategies of the strategies of the strategies of the strategies. Additionally, the protaginging ICPC 2014 works to resuse the strategies of the which is also an achieve melecular in the analosis process. It is such actions as during blood vessels (to increase the flow of matrixin star downers not in markets) and increasing the protections to information and principality as well, and to choose part indication and protections to information and principality as well, and is choose parts integrations of the protections or information and principality as well, and is choose parts in the strategies of the strategies of the transmitter of the protections or and principality and PCC2.

Training intensity and the relative density of arachizotte acid in the phospholial layer (metabolized acid analysis) with a metabolic machining stops in the formation of analysis, prostagalantisis) will distate how much of this potent ipid can be Bornard darage excesse. The distance of the stop of the distance of the stop of the distance of the stop of

Phase II: Localized Tissue Priming

Pluse II is churacterized by a localized increase in growth factor expression and tissue sensitivity to anabole hormores. Those who have always workedned with sub-marked dings of net or localized the actient of these administration of the sensitivity of the sensitivity of the sensitivity anabole of certain receptors in those specific muscles (these really) where it needs to is intract expression of order, and/nego, IC(4), MGF, and its safe to increase the damay of certain receptors in those specific muscles (these really) where it meeds to instance approximation of the sensitivity of the sensitivity of the sensitivity receptors. Steach-statuced marke damage and the These Teopores are both principle time similarity instanced in action of the body that of an engent e. Receptor density can, therefore, be as strong a regularing face on the planmacological activity of nubelic damages at the semint-body face of the damage muscles.





Note: Imhibition of the cyclooxygenase-2 enzyme with anti-inflammatory drugs like likeprofen, accetaminophen, or a spirin, prevents the formation of active prostaglandins. The anabolic cascade is stalled without sufficient prostaglandin formation (Au) Physiol Enderchning Metab 282:2552-16, interching with the normal increase in protein synthesis rates after exercise. It is often advised to use such drugs only when necessary if mancel growth is a key focus.

To put it is perspective, we need to remember that there are two separate components that interact before any message is set to a macch cell obligg to its enserge growth like two is composing neuropeon in the other lipsering coognona analysis from fastisticating and there is the second second second second second second second second lines (oblicable). The more hormsoss or growth likens year has a second second its discus (oblicable). The more hormsoss or growth likens year has a second second its discus (oblicable). The more hormsoss or and second like the lines of the horizing mark composition of the second second second like the process. The second second second second second like the process the horizing means the semiclule message is seen more quickly, and once completed that the analosh messager with like message is seen more quickly and once completed the message) before it is broken down by organics. It is all abort horizing and the second message like it is broken down by organics. It is all abort horizing means the second is the second message like it is broken down by organics. It is all abort horizing the second second in the second second second second message is a second second message horizing the second second second second second second message horizing the second second message horizing the down the second second message horizing the second second message horizing the down the do

White orose hard we have an icrease in tissue sensity to analoke hormons and growth factors, also vial directly first located biases. Priming primes is in microse in the located expression of certain vial growth factors thereabers. This includes $|GF^-|$, MG_r , FGF, HGF, TNK, EI, -1, and L-G. These components will be released, and will work together on the caking damaged markel: Brear and stellite ceds, in a sort of grand septoper of mackandolosis, with each hybrig is now vial size in the process. In marcine sectors and the stellar sectors are also the stellar sectors and the sector of the stellar sectors and the stellar sectors are also also also the stellar sectors in a data the internet owned in a web sector with the scene of the scene of the scene of the rate of the scene of the

Phase III: Repair

Voir local macke issues are prired dring Plases I and II. Dring Plass III, the hormcess and growth factors go to work to fish the Johe Catagorize the plases as no of congeting anabolic action, action mediated by the confined effects of many anabolic hormcess and growth factors including medosyns, resistif, (Lei, Lei, Z., M., G.F., HE, TN, IL, LI, and IL-6. This is the time when requer and hypertrophy are physically taking place in your meaks, and each corespond on β play an intrast role in the process. Hensit set of key the hormcess, the start of the procession in Places I & III has all been hormcessing that are experimely lading up to this point (the starts of in the process. Hensit set of key the hormcessing that the core previous W. veil of Medow the infibiatal sectors of the anabolic components very closely here. During the find places, tissue repair and growth will be finded with the hole of the following hormcess, and around factors.

Hepatocyte Growth Factor (HGF):

HGF is a heparinbinding growth factor that resides on the outer surface of uninjured cells.

opon ngury, a maganes to samme cens where a ungers user activation and entry into use cencycle.³⁰ HGF expression is regulated via infric oxide release,³⁰ which is stimulated upon rigury to also aid in the flow of nativerts and hormones to the area. PGE2 plays a pixotal role in infrito oxide synthesis and HGF release.³⁴¹

Androgens:

Androgens (the hormness that anabolic/androgenis stansiks minici) are strong supporters or potent synthesis ratios in skeletal marker issues. They are also known to stimulate beal [GF-1] expression, so the effects of these hormones cotend to the satellite cell cycle (perhaps explaining why they are such strong simulators of marker growth). It is also of note that anachidone i acil increases androgen receptor density in skeletal masket issue. This helps to fitther piece together the biochemical lists between the Phane Land Phane III neponee.

Insulin-Like Growth Factor I (IGF-I):

IGT-1 is an insulin-like hormone with marked anabolic effects. Owing to its reame, it also has some insulin-like effects as well. IGT-1 increases protein synthusis, and supports the proliferation and differentiation of satellite cells. The prostaglandin PGF2alphn is known to strongly up-regulate local IGT-1 exceptor expression.^{342,343} PGE2 is also believed to play a role in increasing local IGT-1 strongles.³⁴⁴

Insulin-Like Growth Factor II (IGF-II):

IGF-II is a second insulin-like growth factor that plays a role in the proliferation of satellite cells. Unlike IGF-I, IGF-II expression does not appear to drastically increase in response to training.²⁴⁵

Mechano-Growth Factor (MGF):

Mechano-Growth Factor is a recently discovered variant of Insufa-Like Growth Factor I. This growth factor is produced during an alternate spicing sequence of the IGF protein, and plays a strong role in the support of mpoblast proliferation MGF expression, like many of the growth factors discussed here, is strongly up-regulated in muscle tissue in response to stretch simula;³⁶

Fibroblast Growth Factor (FGF):

FGF is a rankly a family of growth factors, with nine different isoftems (FGF) it through FGF. 90. The fail robs that FGF pays is muscle hypertrophy in additional is not faily understood, however, it is believed to be a strong proliferator of stabilite calls, serving to equand their production.²⁴ FGF from galator physica rules in call differentiation: As with many growth production of the equation of the strong production of the strong physical FGF-2 and FGF-4 seems to be the most profile responsitions of this family in matter mack times.

Insulin:

In addition to having some ability to increase protein synthesis and rinhib protein breakdown, insini is the body's chief rarinerit transport hormore. The actions of risaini advo cels to transport glacose and anino acids through the plasm membrane. Insidin receptor expression is storody up-regulated after transmite, exercise, so as to provide more immediate narinfon to the affected area. This up-regulation has been closely linked to the prostaglandin PGE2.³⁴⁰

Cytokines (IL-1, IL-6, TNF):

Cyckiess are a group of immunomchildatory compounds, though in the cortext of this sections were boosely referring to them as growth factors. The IL cyckiess are called interleaking, and TNF is short for furner Neurosis Factor. Among other things, cyckiess are flown to strainable the impairion of physical tyckies, neurophysik, moncetysis, and tother hading such as singling in the removal of chamiged colds and regulating certain influentuoty responses. Including the physical of some prostagationality. Providentian are alrease as chosen to phy important roles in the expression of all three of the cytokines mentioned here, ^{531 352} however, they may not be the sole stimulus. Other pathways of arachidonic acid metabolism may also be involved.

Prostaglandins:

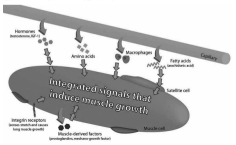
Although these are the key initial reactionary chemicals, prostaglandins continue to play a role throughout the muscle building process (including Pfasse III). This includes their support of hormone receptor proliferation, the enhancement of protein synthesis rates, and an intensitication of the anabole signaling of IGF-1 via a shared pathway (PBK).³⁵³

Estrogens:

Although not specifically highlighted in this outline, estrogens also play a minor role in the anabolic process. This includes helping to increase androgen receptor density in certain tissues (though perhaps not skeletal muscle), stimulating the GH/IGF-1 axis, and enhancing glucose utilization for tissue growth and repair.

Bringing it All Together

So that, in a very loose nutshell, is what is going on inside your body from the time you pick up a weight to the time your muscles are renained, stronger, and ready for more. If the above seems confising to you it should. The fact is, the whole process of muscle growth has been confounding scientists for decades, and undoubtedly will for decades more. We still have a oreat way to so before being able to explain fully how it is that muscle hypertrophy occurs in humans. But as you can see, we have traveled a great distance as well. During the mid-1960s, scientists were only first learning that we grow muscle with the help of satellite cells More than forty years later we have identified, and are experimenting with, dozens of growth factors that were unheard of back then. It is a new world today, and despite not having all the answers, we know enough to enhance human performance in many exciting new ways. But please don't mistake the intention of this section. It is not here to give you a functional roadmap of the entire anabolic process, or to guide you in the ultimate polydrug program. It is here simply to open your mind to the true complexity of anabolism. When we start to see muscle growth from its various angles and intricacies, we begin to see our own potential opportunities for successful exploitation. How many of these opportunities you act upon will depend on your own goals and interests. But no matter how much or how little you actually apply this information. I hope you feel better equipped by having it.



Skeletal muscle growth is a complex process that involves a variety of signaling compounds.

332. Behind the Scenes: Hynertrophy. Gene. Mind and Muscle Magazine 5/2005

333. Regulation of skeletal muscle fiber size, shape and function. J Biomech. 24(suppl1):123-33 (1991)

334. Initial events in eventises induced muscular iniury. Med Sci Sports Fuere 77(4):479.35 (1990).

335. Myostatin neutrively resultes satellite cell activation and self-renewal. J Cell Biol. 162:1135-47 (2003)

336. Similar satellite-cell activation in skeletal muscle: markers, models, stretch, and potential alternative pathways. A Wozniak, J Kong et al. Muscle Nerve 31: 283-300 (2005)

337. Efforts of physical exercise on phospholipid fatty acid composition in skeletal muscle. Agaeta Andersson et al. Am J Physiol. 274 (Endocrinol. Metab. 37): E432-8 (1998)

338. Effects of Exercise on parameters of blood complation, platelet function and the prostaglandin system. H Sinzinger I Virgolini Sports Medicine fr

738.45 (1988)

339. Mechanical stretch induces activation of skeletal muscle satellite cells in vitro. Tatsuni R, Sheehan SM et al. Exp Cell Res 267(1) 107-14 (2001)

340. Release of hepatocyte growth factor from mechanically stretched skeletal muscle satellite cells and role of pH and Nitric Oxide. Ryuichi Tatsumi et al. Mol Biol of the Cell 13 p 2509-18 (2002)

341. Hepatocyte growth factor as a key to modulate anti-ulcer action of prostaglandins in storach. J Clin Inv 98:2604-11

342 The role of prostaduadus in bone formation. Handa SL Balena R et al. Connect Tissue Res. 1995;31(4):279-82

343. Prostaelandin F2a stimulates proliferation of clonal osteoblastic MC3T3-E1 cells by up-regulation of Insulinlike Growth Factor L recentors, Yoshiyuky Hokeda et al J Biol Chem 266(31): 21044-50 (1991)

344. Prostsolvadin E2 stimulates insulin-like growth factor I synthesis in osteoblast-enriched cultures from fetal bone. McCarthy TL, Centrella M et al. Endocrinology 128(6):2895-900 (1991)

345. Secuence of IGF-L IGF-II, and HGF expression in regenerating skeletal muscle. Hayashi S et al. Histochem Cell Biol 122(5):427-34 (2004)

346. Expression of insulin like growth factor-1 splice variants and structural genes in rabbit skeletal muscle induced by stretch and stimulation. G McKov, W Ashley et al. J Physiol 516(2) 583-92 (1999)

347. Expression of fibrolast growth factor family during postnatal skeletal muscle hypertrophy. P Mitchell, T Steenstrup, K Harmon, J Applied Physiol

348. Fibroblast growth factor is stored in fiber estracellular matrix and plays a role in regulating muscle hypertrophy. Medicine and Science in Sports and Exercise 21(5) S173-80 (1989)

349. The insulin-like effect of muscle contraction. Ivy JL. Exerc Sport Sci Rev. 1987;15:29-51.

350. The role of prostadandins as modulators of insulin-stimulated glucose metabolism in skeletal muscle. Leighton B et al Horm Metab Res Sanol

22:89.95 (1990)

351. Differential effects of prostaglandins derived from n-6 and n-3 polyunsaturated fatty acids on COX-2 expression and IL-6 secretion. Dilorit Baam et al. PNAS 100(4) 1751-56 (2003)

352. The role of arachidonic acid metabolism in the activities of Interleukin 1 and 2. W Farrar, J Humes, J of Immunol 135(2) 1153-9 (1985)

353. Regulation of protein synthesis associated with skeletal muscle hypertrophy by insulin-, amino acid and exercise-induced signaling. D Bolster, L Jefferson, S Kimball, Proc of the Nutrition Society 63: 351-56 (2004).

Practical Application

Steroid Cycles

Anabolic/androgenic steroids are not medically approved to promote excessive muscle mass gains (bodybuilding) or improve athletic performance. Aside from early experimentation on athletes by a handful of sports physicians, an extensive effort to study the physique- and performance-enhancing properties of these drugs, specifically with an eve on developing strategies for using them to maximize benefits and minimize adverse effects, has not been undertaken by the medical community. Because of this, illicit users have been left to develop their own protocols for administering these drugs. The result has been a large variety of different approaches to using these agents, some safer or more effective than others. While it would not be possible to comprehensively evaluate all known approaches, this section will discuss some of the most fundamental and time-proven methods for using AAS.

Steroid Selection

When first considering what steroid(s) to use, one will endoe there are many different medications that fill and there the categoor of analysis and the steroid of many sams of development, where specific painters and needs are addressed with dings that have specific characteristics. For example, some dang are considered miker (here androgenic, which miss them better at a payment second factoring in mrs. Some are injectable medications, and others made fails are used to an advect the mission provide matching papersite. In due words, while different ASA shares my have some differing properties. If your objective is to gain nucleon areas.

While all AAS drags may be capable of improving macke mass, strength, and performance, Is would not be correct to say these are no advantages to choosing one gast or or analytic for a particular purpose. Most influentially, the quarky and quality of muck gained may be to have been approximately and the strength of the strength of the produce while water (and sometimes fair) netteriors, however, and are governally stored who may use its more important frammatic definition. Days with bo or on significant consignities that the produce loss durantic star gains in the gain gain haves on the strength of the strength of the strength of the quality in higher, consignities that the produce loss durantic star gains in temperatures to the the quality in higher, consequently then its the text own and angeneries as falsows.

Mass (Bulking):

Methandrostenolone - Oral Oxymetholone - Oral Testosterone (cypionate, enanthate) - Injectable

Lean Mass:

Boldenone undecylenate – Injectable Methenolone enanthate – Injectable Nandrolone decanoate – Injectable Oxandrolone – Oral Stanozolol – Oral

The potential for adverse reaction should also be considered when choosing a stemule to use, especially first. Site us is to lengthy repeated. For coursels, he finded our medications pressere greatest strains on the cardiovascular system, and are also fore truck. For most of all, horized coursest is also first the most observable and the strain strain of all and truth coursest is also first the most observable and the strain methods, and and the strain observable and the strain observable and methods, methods, and and a strain observable and the strain observable land, may isolate their use to prodominarly analysis drags, such as muchoises, notentials de (Brick Cardhour is advised) before coursing to an ΔA is methods.

Dosage

The dosage used is important in determining the level of benefit received. Anabolic/androgenic steroids tend to be most efficient at promoting muscle gains when taken at a moderately separheraportic dosage lved. Enkv mis (chearportic), postratil anubolic benefina are oftice controlutioned, at last to some extext, by the suppression of endogenous lastoatenere. At vary high droses (accessive separheraportic), smiller optimules, and the second second second second second second second generality second second second second second second second generality second second second second second second second (moderate sequencing), solvest, eff drog in high efficient at sporting may be proprinted, for carring losses of the other second second second (moderate sequencing) and the second second second second (moderate sequencing) and the second second second second (moderate sequencing) and second second second second second second (moderate sequencing) and (moderate second second second second second (moderate sequencing) and (moderate second second second second second second (moderate sequencing) and (moderate second secon

- Boldenone undecylenate: 200-400 mg/wk
- Methandrostenolone: 10-30 mg/day
- Methenolone enanthate: 200-400 mg/wk
- Nandrolone decanoate: 200-400 mg/wk
- Oxandrolone: 10-30 mg/day
- Oxymetholone: 50-100 mg/day
- Stanozolol: 10-30 mg/day
- Stanozolol: 10-30 mg/day
- Testosterone (cypionate, enanthate): 200-600 mg/wk

These are additional considerations ofter than the cost effectiveness of a particular dosage. To beyin which high does of mable chardwords strends for target barbace strends regardly connectic, psychological, and physical side effects. In high of drinnishing returns, the tradooff between results and adverse reactions becomes less and less forwalls. Gains made on lower does also tend to be better tenined after strend discontinuance than those resulting fram cascoise indice. It is garently not results to expect the most allowed barbace and a valued. It is also very invertion to remarks the high doebs-dist weight gains induced by musice dosing of memi leng data expects a voice in the strend strends and the strend distribution of the strends of the strengs of the strends of the

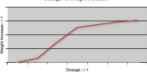




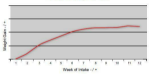
Figure 1.Anabolic/androgenic steroids tend to be most effective in moderately supratherapeutic doses. The anabolic benefits diminish in relation to the amount of drug given at both the high and low ends of the dosage range. Duration (Cycling)

The administration of anabolic androgenic staroits at a given dosage well typical produce molecular increases in macke size and strength for approximable 8-8 weeks. After this point, the net of new marke gian typically slows significant programs to so maler, where all downed memortamines accessed. To contraine uniting significant programs beyond the point can entail escalating dosages, which is likely to cortecide with a greater incidence of advecement excitons and draining anabolic returns. Even whitted dosage corrected diary agakkly. The practice of entended or contraines stored administration is discoursed for they servers. It is averable to correct works advecement advecement and advecement is a store source. It is averable to corrective stored administration is a final store of the store store advecement in the store of than 8 weeks at a time (10-12 weeks at the maximum), followed by an equal or longer period of abstinence before another steroid regimen is initiated. This pattern of rotating between "on" and "off" periods is referred to as cycling.

Off-Cycle (Recovery, Bridging, and Tapering)

The point immediately following actival cassion can intole a state of programmlinm (low manopan level), and as reading point indications in an offert to minime marker, how it dedottimes it simulation, and rearring additionality in programmling and approximation of the state of

When corcluring a cycle, some strend users also follow a practice of field solvy reducing their doagest (neuror). This sporring may proceed for a 3-4 week speering, and will income an even stopping down of the dose each week will the post of dhug discontinues. It is uitationan, however, if a with taptient going and tapable suite. The parts the maxes bear with some other data guards and the proceeding and the processing of the each AAS administration study can also be that the year of standard dose particip. One first in the logic of using a tapering program is that they are standard dose parts of a therman encourse. Focus ye is not processle, however, which does AAS administration study can also be found to eard the measure doses, however, which does AAS administration study can also be found to eard the measure doses, however, which does AAS administration study can also be found to eard the measure doses at the year of the standard doses and the parts of the standard dose tapering is not a proven usy to reduce poor - took muck antibolism.



Duration vs. Weight Increase

Figure 2.Anabolic/androgenic steroids tend to be most effective at a given dosage for approximately 6-8 weeks.After this point, the rate of new muscle gain will slow and soon after will usually hit a full plateau. Stacking

As individuals become more experienced with muboliconflogaries steorid use they may begin experimenting with the use of more than one stood at a time. The practice is referred to as instantical Stacking is most common with advanced to boluntations with a first base of the proviso single-agent approach. In many cases, however, it may simply be the grader carnalistic stood doogs that is necessary for the resamed progress. Stacking usually include the combined of a more androgene statemost with the other than the approximation of a more androgene statemost with one per terming analysis, and and the analysis and an experiment statemost of the transfer the other the agents. On the analysis and, and an experiment is there of most statemost are inducationable with stress and as malorized has one of the statemost and mutantocatorshow the stress as the androgene has one state. In reasons at succear annuogane man annovas sources loganet in un immer are non 646. Of the one hand, high does of estoratorence, conjerdendore, or methandonschoel neue prone to producing strong androgenic and estogenic side effects. Stacking first because very popular durig the Dybos, in mit when effective entropy methan which also strong the behavior behavior and the strong main strengenic and the strong behavior effective and the behavior behavior and the strong strengenic which also would be behavior but in signer androgen, attached annotation and the strengenic effective and the strengenic strenge

Today, the availability of dangs that can reduce settings at richly makes the corritance (u see single gater cycles) based on a strong androps like test-storeme carathat or exploring much more width than it was decades ago. Side effects like genecomstain and water restorinc can row be effectively minitated with anti-settoppen or availability in the when taking lighter doeses. Individuals should be aware that stacking is, Review, not a necessary practice. It is dealy to small commonly applicable in competitive helphalinght circles, however, or when an individual is are they have progressed as far as they possibly bolyhalikes, the periodic use of a single stared will be more than suffacer to maintain optimal levels of mucch muss and performance, and it may never be necessary to deviate from this approach.

Sample Steroid Cycles

The following cycles are presented as examples of common steroid administration protocols. These programs have not been evaluated in a cincil sating for sharp and efficiacy, and are provided for informational parposes only. These are not recommendations for economic years one incidental hands statistic, the histic has an effortune could be decising it is imposed to a statistic to the statistic hardwork of the decising it is imposed to the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the statistic hardwork of the statistic hardwork of the statistic hardwork of the medical hardwork of the statistic hardwork of the stati

Single Agent Cycles

Dianabol Cycle #1 (Mass)

Products: 100 tablets 5 mg Methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4 g/day).

Estrogen Support: tamoxifen (10-20 mg/day).

Comments: This is a very common first cycle for building mascle mass, and uffizes a single standard borth of methandrostenolone. This cycle is likely to produce very noticeable muscle growth an a first-time started user, often in excess of 8-10bs of weigit gain. This is usually not accompanied by significant visible side effects such as gynecomustia and water retention. Adhough this is considered a beginner 's cycle, methantostenolone is a C-11 alpha adylated Athough this 'considered a beginner 's cycle, methantostenolone' is a C-11 alpha fadylated set of the s oral steroid, and presents significant cardiovascular and liver toxicity. The repeated use of such drugs should be limited.

| Week Methandrostenolon | | |
|------------------------|-----------|--|
| 1 | 10 mg/day | |
| 2 | 10 mg/day | |
| 3 | 15 mg/day | |
| 4 | 15 mg/day | |
| 5 | 20 mg/day | |

Dianabol Cycle #2 (Mass)

Products: 200 tablets 5 mg Methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is a common follow up to the first Dianabol cycle, trilizing a slightly higher dose and longer duration of intake. The dosages used here are more common for bodybuilting purposes. A slightly greater intervity of adverse nearcitors is likely.

| Week | Methandrostenolone | |
|------|--------------------|--|
| 1 | 20 mg/day | |
| 2 | 20 mg/day | |
| 3 | 25 mg/day | |
| 4 | 25 mg/day | |
| 5 | 25 mg/day | |
| 6 | 25 mg/day | |

Testosterone Cycle #1 (Mass)

Products: 10 mL 200 mg/mL Testosterone (enanthate or cypionate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4 g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5 mg/day).

Comments: This muss building cycle is likely to yield similar quantitative results as an early Diambol cycle, but is favored over the oral for its lower cardiovascular and hepatic strain. The doors used are expected to cause mild shifts in the HDL/LDL cholsterol ratio, but not the substantial changes normally seen with oral anabolic storoids. This sample cycle is fledy to present the least arourd or fleah halis def flex for flexible in this section.

| Week | Testosterone | rone | |
|------|--------------|------|--|
| 1 | 200 mg | | |
| 2 | 200 mg | | |
| 3 | 300 mg | | |
| 4 | 300 mg | | |
| 5 | 300 mg | | |
| 6 | 350 mg | | |
| 7 | 350 mg | | |

Testosterone Cycle #2 (Mass)

Products: 20 mL 200 mg/mL Testosterone (enanthate or cypionate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1 mg/day).

Comments: This cycle is a common follow up to the first testostence only cycle, with a higher doogs and 3 week loared during to minke. The total testosterme doorse given is double in comparison, and is likely to produce more prorozoared estrogenic and antiogune is determined and the second second second second second second mean assistantialy lower than cycles with our IAAS. Testosterone is arguably the safest, and at the same time one of the most effective, music-bailing second second exclusions the cycles like the sword be achieved and the second exclusions are assisted as a second second second exclusion second second second exclusions are assisted as a second exclusion second second exclusions are assisted as a second second exclusion second exclusions the second exclusion second exclusions that the second exclusion second exclusion second exclusions exclusi

| Week | Testosterone |
|------|--------------|
| 1 | 200 mg |
| 2 | 400 mg |
| 3 | 400 mg |
| 4 | 400 mg |
| 5 | 400 mg |
| 6 | 500 mg |
| 7 | 500 mg |
| 8 | 500 mg |
| 9 | 500 mg |
| 10 | 200 mg |

Sustanon 250 Cycle (Mass)

Products: 15 mL 250 mg/mL Sustanon (testosterone blend)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1 mg/day).

Comments: This mass building program is similar to the other testosterone cycles, but utilizes Statamon 250, a form of blended testosterone more widely used in Europe and other regions outside the U.S. The total starced dosage of this cycle is 3,750 mg, externily close to the amount used in testosterone cycle #2. A similar level of cardiovascular strain and visible side effects are exercted.

| Week | Sustanon | |
|------|----------|--|
| 1 | 250 mg | |
| 2 | 250 mg | |
| 3 | 500 mg | |
| 4 | 500 mg | |
| 5 | 500 mg | |
| 6 | 500 mg | |
| 7 | 500 mg | |
| 8 | 500 mg | |
| 9 | 250 mg | |

Products: 50 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: Operativose is controlos regarded as the most poter muss bailing storeid walshiel. It is also prote to causing both storeg estrogaria and anticognic side effects. A stored rovec may gain 15-20 pounds or more on this cycle, altough a significant annuat of this will be sume restriction, which will assist accound far day documance. Operativolution more conversient to use thin an injectable testosterour, it is not regarded as a significant Repeated use of (-1) apha hadylated on the first document.

| Week | k Oxymetholone | |
|------|----------------|--|
| 1 | 50 mg/day | |
| 2 | 50 mg/day | |
| 3 | 50 mg/day | |
| 4 | 75 mg/day | |
| 5 | 75 mg/day | |
| 6 | 75 mg/day | |

Oxymetholone Cycle #2 (Mass)

Products: 100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is a more popular version of the oxymetholone only cycle. The doses here are more common with experienced steroid users, and more than sufficient to promote strong mass and strength increases. Side effects may be more noticeable than the lower dose cycle, of course, which may necessitate a higher dose of tamoxiden.

| Week | Oxymetholone | |
|------|--------------|--|
| 1 | 50 mg/day | |
| 2 | 50 mg/day | |
| 3 | 100 mg/day | |
| 4 | 100 mg/day | |
| 5 | 100 mg/day | |
| 6 | 100 mg/day | |
| 7 | 100 mg/day | |
| 8 | 100 mg/day | |

Stanozolol Cycle #1 (Lean Mass/Cutting)

Products: 200 tablets 2mg Stanozolol

An weeks: Liver Support: Liver Statu, Liv-52, or Essentiate Forte (anter recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: This is a common first-cycle for an athlets looking for performance improvements on a holphadier looking for a karm mass or carring starout. This cycle was more common when stanozold was widely available in 2 mg tablets. Such preparations are now ancommon occupied in a trayer. The doogs and there is two bolyholphading standards, especially during the 1970s and 80x. Significant visible advence reactions are utiliably at this doogn.

| Week | Stanozolol | |
|------|------------|--|
| 1 | 8 mg/day | |
| 2 | 8 mg/day | |
| 3 | 10 mg/day | |
| 4 | 10 mg/day | |
| 5 | 10 mg/day | |
| 6 | 10 mg/day | |

Stanozolol Cycle #2 (Lean Mass/Cutting)

Products: 200 tablets 5 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: This is a stronger version of a cutting/han muss building cycle tabling stronolab. The doogue such here is shartarilab/higher hum the first staraoobile cycle, a fact that makes this cycle more properly sited for bodyheading parposes than Staraoobil Cycle 41. Cardiovascetar and heptite strain wild be more notabilized, and wish site deficies more pronounced, than the first cycle. Three should be no need to addition an estrogen minitemace drug.

| Week | Stanozolol | |
|------|------------|--|
| 1 | 20 mg/day | |
| 2 | 20 mg/day | |
| 3 | 25 mg/day | |
| 4 | 25 mg/day | |
| 5 | 25 mg/day | |
| 6 | 25 mg/day | |

Stack Cycles

Deca/Dianabol Cycle #1 (Mass)

Products: 10 mL 200 mg/mL nandrolone decanoate

100 tablets 5 mg methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is an externely old and widely repeated storoid contribution, based on the prodominutly analysis is storial randomic decanosis. Mediumdostanchuse arsvers as the antiogene component of this stated, and is added driving work 3, which is a time that side differst or forelasci antioopasieby (with the occusion use on transforme documents) are are more highly most bodybading standards, but are sufficient to impart a noticeable increase in muck size and strength.

| Week | Nandrolone | Methandrostenolone |
|------|------------|--------------------|
| 1 | 200 mg | |
| 2 | 200 mg | |
| 3 | 200 mg | 10 mg/day |
| 4 | 200 mg | 10 mg/day |
| 5 | 300 mg | 10 mg/day |
| 6 | 300 mg | 15 mg/day |
| 7 | 300 mg | 15 mg/day |
| 8 | 300 mg | 15 mg/day |

Deca/Dianabol Cycle #2 (Mass)

Products: 20 mL 200 mg/mL nandrolone decanoate

200 tablets 5 mg methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: A more popular manifestation of the Deca/Danabol Cycle, with more commonly accepted dosages for a moderately experienced steroid user. Incidences of side effects are expected to be higher at these dosages, although overall this stack is likely to be less problematic than a contribution of testosterone and osymetholone.

| Week | Nandrolone | Methandrostenolone |
|------|------------|--------------------|
| 1 | 400 mg | |
| 2 | 400 mg | |
| 3 | 400 mg | 10 mg/day |
| 4 | 400 mg | 10 mg/day |
| 5 | 400 mg | 20 mg/day |
| 6 | 400 mg | 20 mg/day |
| 7 | 400 mg | 20 mg/day |
| 8 | 400 mg | 20 mg/day |
| 9 | 400 mg | 20 mg/day |
| 10 | 400 mg | 20 mg/day |

Testosterone/Anadrol Cycle (Mass)

Products: 20 mL 200 mg/mL testosterone (enanthate or cypionate)

100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: A contribution of testosterore and oxymetholne is generally regarded as the most potert 2-days takes for pairing rank much must. Both days will present significant ostogenerity, and will be likely to induce genecomstati quickly unless an estogen minitenance drag and a tempolities in additional lange linear data data the best for the state of the state of the state of the state of the state. Inoverse, and a rapid loss of water weight (possibly up to 10 pounds or more) is expected soon after the cycle is discontributed.

| Week | Testosterone | Oxymetholone |
|------|--------------|--------------|
| 1 | 200 mg |] |
| 2 | 400 mg | 1 |
| 3 | 400 mg | 50 mg/day |
| 4 | 400 mg | 50 mg/day |
| 5 | 400 mg | 100 mg/day |
| 6 | 500 mg | 100 mg/day |
| 7 | 500 mg | 100 mg/day |
| 8 | 500 mg | 100 mg/day |
| 9 | 500 mg | 100 mg/day |
| 10 | 200 mg | 100 mg/day |

Testosterone/Deca Cycle (Mass)

Products: 10 mL 200 mg/mL nandrolone decanoate

10 mL 200 mg/mL testosterone (enanthate or cypionate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1mg/day).

Comments: Testosterore with matchone is considered to be one of the most findamental 2-deng constraints matchs. Nandrokec compliants the androgenit lake of classotteroe by supplementing additional anabola activity without strong estrogenicity. The resulting stack is admost as production as a cycle utilizing, a ligher does of testosterone alone, but less problemating in terms of estiogaria side effects such as water reterion, gatecomstai, and fat buildag. Estogen conversion is still formibialle compliable compliable compliable match the use of an estogena matietance ding, however, and this stack remains in the realm of mass building instead of law muss or entire.

| Week | Testosterone | Nandrolone |
|------|--------------|------------|
| 1 | 200 mg | 200 mg |
| 2 | 200 mg | 200 mg |
| 3 | 200 mg | 200 mg |
| 4 | 300 mg | 300 mg |
| 5 | 300 mg | 300 mg |
| 6 | 300 ma | 300 ma |

| 7 | 300 mg | 300 mg |
|---|--------|--------|
| 8 | 200 mg | 200 mg |

Andriol/Anavar Cycle (Lean Mass)

Products: 360 capsules Andriol 40 mg

400 tablets oxandrolone 2.5 mg

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: This is an effective but raid cenk-ordy lean muss bailing cyck: Archivia is used as the antrogenic bases, but in abous that do not graudy exceed remain thempartic i-best. Orandrokene is non-normatizable, so significantly elevated estrogenicity is utilisely limited in the product set of the problem limit. This stack is popular armung older men and shous net wishing to use spicetime. The provide domains starts that the same of the should be statisticable and should be statisticable and the set of the start of the set of the s

| Week | Andriol | Oxandrolone |
|------|------------|-------------|
| 1 | 240 mg/day | 15 mg/day |
| 2 | 240 mg/day | 15 mg/day |
| 3 | 240 mg/day | 15 mg/day |
| 4 | 240 mg/day | 20 mg/day |
| 5 | 280 mg/day | 20 mg/day |
| 6 | 280 mg/day | 20 mg/day |
| 7 | 280 mg/day | 20 mg/day |
| 8 | 280 mg/day | 20 mg/day |

Anabolic-Androgenic Bi-Phasic Stack (Lean Mass)

Products: 18 mL methenolone enanthate 100 mg/mL

50 mL boldenone undecylenate 50 mg/mL

20 mL testosterone (enanthate or cypionate) 200 mg/mL

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1mg/day).

Comments: This is a 3-month non-bite-toxic cycle that has 2 distinct phases, muss and lean mesoritating. The first workeds of training and the are focused on mossibility. Significant estogarizity will be present in these weeks, and may necessitate the use of tarnoxidian or an anomation influible studies of the present generometry and the costs of works and the studies of the studies of the studies of the studies of of plane 2, which focuses on increasing the molegoed to estogarize that the start of plane 2, which focuses on increasing the molegoed to estogarize that and subliding the effects to present second objection or hose of linkide, which often occurs with the use of protominity analysis storik along.

| Week | Testosterone | Boldenone | Methenolone |
|------|--------------|-----------|-------------|
| 1 | 500 mg | 200 mg |] |
| | | | |

| 2 | 500 mg | 200 mg | |
|----|--------|--------|--------|
| 3 | 600 mg | 200 mg | |
| 4 | 600 mg | 200 mg | |
| 5 | 600 mg | 200 mg | |
| 6 | 600 mg | 200 mg | |
| 7 | 100 mg | 200 mg | 300 mg |
| 8 | 100 mg | 200 mg | 300 mg |
| 9 | 100 mg | 200 mg | 300 mg |
| 10 | 100 mg | 200 mg | 300 mg |
| 11 | 100 mg | 200 mg | 300 mg |
| 12 | 100 mg | 200 mg | 300 mg |

Testosterone/Anadrol/Trenbolone Cycle (Mass)

Products: 30 mL 200 mg/mL testosterone (enanthate or cypionate)

20 mL 75 mg/mL trenbolone acetate

100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20 mg/day) and anastrozole (.5-1mg/day).

Comments: One of the more extence mass balling cycles in common use among bodyhakkes. This set well appent rup glues in new macks less all entergell. This doucombination is highly proces to causing estragantia and antoquinits side effects, lickling in the cycle. These cost of macmutates inhibiter is lickly by encreasing to a down one is the cycle. These cost of macmutates inhibiter is lickly by encreasing to a down one morning, however, which may necessating the address of the start highly problematic with regard to aide effects, and therefore mady recommended to trubohome for bullet rup dranket mass.

| Week | Testosterone | Oxymetholone | Trenbolone |
|------|--------------|--------------|------------|
| 1 | 300 mg | | 150 mg |
| 2 | 600 mg |] [| 150 mg |
| 3 | 600 mg | 50 mg | 150 mg |
| 4 | 600 mg | 50 mg | 150 mg |
| 5 | 600 mg | 100 mg | 150 mg |
| 6 | 600 mg | 100 mg | 150 mg |
| 7 | 600 mg | 100 mg | 150 mg |
| 8 | 600 mg | 100 mg | 150 mg |
| 9 | 600 mg | 100 mg | 150 mg |
| 10 | 600 mg | 100 mg | 150 mg |
| 11 | 300 mg | | |

Masteron/Primobolan (Lean Mass/Cutting)

20 mL 100 mg/mL methenolone enanthate

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day)

Comments: This is an effective stock for hardening, catting and gaining lean massel mass. Neither agent is capable of converting to estrogen, so this cycle should significantly elevate the androgen to estrogen ratio. This may assist in the breakdown of this tosus, enhancing mascle definition. This stack should not present significant liver toxicity, although cholesterol ratios may be significantly altered in light of reduced estrogenia existy.

| Week | Drostanolone | Methenolone |
|------|--------------|-------------|
| 1 | 200 mg | 200 mg |
| 2 | 200 mg | 200 mg |
| 3 | 200 mg | 200 mg |
| 4 | 300 mg | 300 mg |
| 5 | 300 mg | 300 mg |
| 6 | 300 mg | 300 mg |
| 7 | 300 mg | 300 mg |
| 8 | 200 mg | 200 mg |
| | | |

Winstrol/Proviron/Trenbolone Cycle (Lean Mass/Cutting)

Products: 250 tablets stanozolol 5 mg

100 tablets mesterolone 25 mg

20 mL trenbolone acetate 75 mg/mL

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: Stanoold and terebooke use popular stavids during cutting phases of training, and inputs storg mathics and modern and angosic effects with no significant estoparised. This contribution helps to imput a storug fit basic/difficience-stratering effect. Thus 25 mg tables is of metaricone have been added per day to sapplement additional andacativity which should help matrian normal Bakis and searching. Additional andmory of theorem and the storage storage storage and the storage and the should help matrian normal Bakis and search factoring. Additional and the should help matrian normal Bakis and search factoring. Additional and the should be the storage storage storage and the storage and/societar and help the storage and the storage and/societar and helps the storage and/societar and helps the storage and and the storage and/societar and helps the storage and/societar and helps the storage and and the storage and the s

| Week | Stanozolol | Mesterolone | Trenbolone |
|------|------------|-------------|------------|
| 1 | 20 mg/day | 25 mg/day | 150 mg |
| 2 | 20 mg/day | 50 mg/day | 150 mg |
| 3 | 20 mg/day | 50 mg/day | 150 mg |
| 4 | 20 mg/day | 50 mg/day | 150 mg |
| 5 | 25 mg/day | 50 mg/day | 225 mg |
| 6 | 25 mg/day | 50 mg/day | 225 mg |
| 7 | 25 mg/day | 50 mg/day | 225 mg |
| 8 | 25 mg/day | 50 mg/day | 225 mg |
| | | | |

PCT: Post-Cycle Therapy

Pose-Cyck Therapy, or PCT for short, nefros to the practice of using certain medications to assistant in the discontrance of analysis stressis. What sensols are not addicide dargs in a classical anese, they do suppress your own hornness production, at least emportant). This is an usen that should be addressed in the concellos of the stressistical and and abruptly without addressing internal hornness production, the result could be a prolonged static of hypogenetic (bus adresses) to else concellos and according the stress is an originated static productions and the stressistical in so of much much concellos and the stressistic and the stressistical in so of much and and the stressistic and the stressistic and the stressistic and the concellos much concellos and the stressistic of the stressistic and we will also do doness certain medications that and correct the forward in relative window to help simulated natural control of format of during the postcycle window to help simulate natural to testosteome productions in stressistic relative stressistic and the stressistic much and control of the normal in relative stressistic and the stressistic and the contrast of the stressistic much and control to the normal in the stressistic and the stressistic and to stressistic and contrast the hornness in the stressistic and the stressistic and to stressistic and contrast the hornness in the stressistic and the stressistic and to stressistic and the stress

The HPTA Axis

In the lumm body, the Hpothanine-Phinkary-Tesischar Avia (HPTA) controls teststemere biosynthesis. The PHY is a highly engagined system of obeless and hances that works to assure the correct test of siststearcore is maintained. We can look at this engalating as hiving three levels. At the usy is the hopothanine region of the bars, which relaxes and the start of the start of the start of the sist of the start of the Hormon (LH). LH is this mesh can suggest the levels, which relaxes (GRH seads a signal to the second level of the axis, the phinkary to produce Lueixing Hormon (LH). LH is this meshs can suggest on the levels (level freed) was result of the LH simulation serve as a constrabution: They provide negative feedback to lower the securitor of LH and lesistatores, operating userproduction, Synthest analosis, theorem (L), LH and lesistatores, operating userproduction, Synthest analosis, theorem (L), and relaxed to the security as a methylic strateging the theorem (L) and relaxed the security of the size (level freed) as the security of the securitor of LH and lesistatores and other security strateging the security of the theorem (L), and relaxed to the security of the security of the security of the security of the securitor of the security of the security



The Hypothalamic-Finitary-Testicular Axis: The hypothalamic releases foundatorian Recissing Bormonic Guildin, builds ortimatises the pinitary to release harinary in the release of testistarease from the testics. Androgena, as well as the provide the release of testistarease from the testics. Androgena, as well as hypothalamism and pinitary, lowering the output of goundatrophics and testosterones when to mach hormonic is present.

Unaided HPTA Recovery

The appression of matral testoarce synthesis by strend use is typically a temporary homorrown. Ever Jyou do northing your holy's normal andregon synthesis will usually return a few to sevent morthe after the cycle is concluded. The problem is, this can be a very long time when you are orbhyge on testoarcene for so may these, itselding the matteratures of matche tissue. In fact, much of the matche mass achieved during AAS administration can be to at the works and much too folds. We handle work the administration can be to at the works and much too folds. We handle work the HPTA, so normal hormore production howke may come hack more quickly in order to accound that and which, housever are normal to fits out HPTA more and HPTA more and the HPT normally looks like without assistance. Only then can we identify the levels of the HPTA that are most open to manipulation with support medications.

Statics on the post-costation appect of anabole staroit use, especially in AAS abatesr, are blering. In more cases we mater refor a single-appert taking, stangle of hormour repleneement patients. Due of the most detailed views of what a post-cycle crash probably looks like cosmes from an investigation into isotosterone annulta-¹⁵ ¹⁵ In topolo a group of ment were given weekly rejections (250 mg fr c21 weeks, a dose that admittedly deeg as posport normal HBT use. Hornos hormous were measured each week during the statigy and for a first star of the star star of the star of

The above study suggests that one of the first things to lappen after strend cosmicn is that the brain recognize strendermer. Before are using in This of class GMH and L1 Howes begin correcting first/ quickly. The substantial delay between this and an increase in instantome brech is caused largely by including arrowsponsessors to bidding howes of mose introphedic. This is a weld-concrated side effect of mabelia stared use, even if a size difference may not be introducely which is all cases. When the low-base instantome is the interaction which is all cases. When the layes have a significant on the introduced which is all cases. When the layes have a staring magnetized the stare of the stare is always brains the size of the other mass. While appendix of the post-syste recovery predicatandy braing characterized by effective.

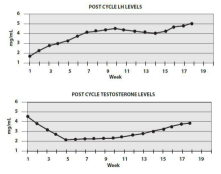


Figure 1. LH and Testosterone measurements starting 1 week after the last injection of 250mg of testosterone enanthate (pretreated measures were 5 mU/ml and 4.5 mg/ml respectively).Note that between weeks 1 and 5, as testosterone levels are declining due to the essation of exogenous androgen administration, LH levels are beginning to correct. From weeks 5 to 10, testosterone levels

near baseline, although LH is increasing by this point.No notable correction in testosterone occurs until after the 10-week mark. bCG in PCT

Henron Cheroixe Gonzáloropen (CCC) is a farilig dug turt minist for actions of cheroixing hermone. It is commenty used during the post-sck-periodi to distributes testicikar mongles, which as we have seen is one of the fardamental conditions to intermediate the schematic schem

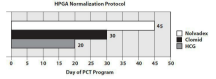
Anti-Estrogens in PCT

It is important to note that the use of arti-estrogens alow is generally not regardled as an effectively strategy of addressing hormera convey at the conclusion of a strated yeel. This is because these drugs only work by lostering the heightened release of Marinized hormerae. We expect that the post-cycles whole as already partly charactered by normalingh. It levels. Thus, while artis-emogens may have an additise effect in this regard, they do not efficiently and directly addieses the mini andicatofic to hormatic additional theory affect should be directly and and end addieses the mini andicatofic to hormania for additional theory affect should be directly and and end addieses the mini andicator and the ord for mechanism directly and the formabile peeted of AAS has; which at likes all these of the methanism directly and the section. There are some exceptions when an abbreviated PCT program may be desirable, which we will discuss ther on.

Traditional PCT Program

The following PCT program was developed by Dr. Mehnel Scalb, one of the most well known and accomplicat inviduals in the differ diardinois distortional multitomorally and generated inviduals. In the distortional hypothesis of the formation of the distortional structure of the distortional hypothesis of the distort invidual of the distortional hypothesis of the distort interime and trade and done block work on hardreds of patients, and while doing so developed the following PCT program. A skiptly motified from of the program works called a structure and the distort block of the distortion of the program works. Scalb's "HTCM-Normalizations Protocod" leaves on the construct are of the distort of the distort of Normalism Protocod" leaves on the construct are of the distort of the distort of motified and and distort approach paris-jetch leaves program scalability.

This PCT program begins with a substantial dose of LGG (2000 U) every other day for 20 doy). Arti-storgans are also used during this proint. This is potertially inportant because LGG may up-regulate testicular aromatese activity.³³⁸ Thus, their use can minimize both compare, side effects and rocken engine feedblock infihtion of storstoremore release. The arti-storgane taken are tamovalien of that (20 mg twice per day) and cloniphene crime (50 mg twice per day). Clondi is used for a shorter period of time, in a storging down of the program's mcKatafors. While in the first couple of wecks the arti-estrogens may not be lighly effective, hey should prove more circula lowards the mddk and end of the program. In the published version of Scally's program (which is slightly mddified from the above), normal hormound larticon retrander on a sloghest with rd's dyst. This is a definite success, far more flowards than the protoacted recovery window reported in the study with 250 mg/weck of restoratore curranture.



Protocols: Human chorionic gonadotropin (hCG) is taken at 2000IU every other day for 20 days. Cloniphene citrate 50 mg is taken twice per day for 30 days.Tamoxifen citrate is taken 20 mg twice per day for 45 days.

The timing for a Post-Cyck: Thempy program can be as important as its compositon. If it is initiated to bia, valuable days of oreand Hermone beside and also down made massing buy be last. Flow usart far program its order, you may miss the optimal variation of differencess. FUCT program monte in mechanics in particular, we want to make a we that ICG a beside applied right anough the inne data cognorous strends are dispeting below the threshold of physicological analysis in the strends of the

The exact trains for FCT program is determined by the elimination hulf-like of the drug() used. We vill use isolatoren cryointed trainstantian as an example. Use some ech injection fram an eliminator hulf-like of approximately & days. Alsose of 200 rap/week should produce the hulf-result of the elimination elimination of the eliminat

Timing the Start of PCT

| Type of AAS used | PCT Initiation |
|---------------------------------------|---------------------------------|
| Cypionate/Enanthate: (200 mg/week) | 3-7 days after last injection |
| Cypionate/Enanthate: (500 mg/week) | 10-14 days after last injection |
| Decanoate: (200 mg/week) | 10-14 days after last injection |
| Decanoate: (500 mg/week) | 18-21 days after last injection |

³⁵⁴. Effect of long-term testosterone oenanthate administration on male reproductive function: Clinical evaluation, serum FSM, LH, Testosterone and seminal fluid analysis in normal men.J. Maass, G. Borsch et al. Acta Endocrinol 78 (1975) 373-844

335. Desensitization to gonadotropins in cultured Leydig tumor cells involves loss of gonadotropin receptors and decreased apacity for steroidograesis. Freeman DA, Accoli M Proc Natl Acad Sci U S A 1981 Oct;78(10):6309-13

356. Inhibition of Ineliating hormone scontisch by totosterone in mm requires sconarization for its physikary but on its is hypothalmine effects: existence from the transmission mean and grandwarepin-reducing hormone-deficient men. Pitteknad N., Dayer AA, DeCrar S, Lee H, Beepple PA, Crossby WF Jr, Hayes FJ. J Clin Endocrinol Metals. 2008 Mar 9573: 1784-91. Embed 2007 Dec 11.

357. The effect of aromatase inhibition on sex steroids, gonadottopins, and markers of bone tumover in older men. Taxel P, Kennedy DG et al. J Clin Endocrinol Metab. 2001 Jus;86(6):2869-74.

358. Acute stimulation of aromatization in Leydig Cells by Human Chorionic Gonadotropin In-vitro. Proc Natl Acad Sci USA 76/4460-3,1079

OCT: Off-Cycle Therapy

The objective of anabolic strendt therapy (when neuroscial applications are involved) stoud be to cick the olscied benefits with the bowst cramitine exposure and size effects. This normally includes diligance with optimizing all aspects of training, rest, and det, as well as addreigt to a 19-cick) therapy (CPC) program the conclusion of cach stored (2-c). One the one hand, we want to make each cycle as productive as possible. On the other, we are string to restint more gains so the starting point of the next cycle is that much future along. When all aspects are in check, the result should be a next for lower that looses, fower cycles (longer admices) of abilitative), and should be a next for lower that looses, fower cycles (longer admices) of abilitative), and should be a next for lower that loose, fower cycles (longer admices) of abilitative), and should be a next for lower that loose. Space registing to resting the other lower and lower that more may be a strend be applied for the other ability.

Gen the inputates of relating our mack and performance gains, however, our effects in this negral should not conclude with Post-Cyck Tranpy, Ideade, to necise the granetic host-entre-hearths from ambolic-introducing strend theory is a host absolute to initiate an OB-Cyck Theory) (CCT) program when the PCT is over. The focasi of CCT is syncarble one and matrixal substances (denues) applements) that faster marks thereints, while and even advantile to approach fastery approximents with a halfwile you'd a copprised fast that and the substances of the strength faster was able to the strength faster and fast the significant strength fast was able to the strength faster and fast the significant strength fast was able to the strength faster and the strength fast has significant strength fast was able to the strength faster and the strength fast has spin-faster and programs more effective in the ablence of program strength faster and the strength fast was able to the strength faster and the strength fast was able to the strength fast was able to the strength faster and the strength fast was able to the strength fast was able to the strength faster able to the faster able to the strength faster ab

A well-organized CCT program basis a minimum of aix to eight weeks, and consists of three district compounts. The first is "distantous Support," which seeks to end and effective PCT program, but with a different and much more basis approach. The second part is "Maske Cell Redsensation," Howy training disrupt the marked cell methems, so that the markets become less response to exercise stimulation. We want to address this darge OCT, and prime the mucks how the off-tense training Lange, we want to include one or more manal muck-holding adsounces in the program. The part is called "valuable double barried, and more market mass will be reinised in the lange n. All free OCT components are taken simulaneously, sometimes for the full period between the end oFPCT and the start of the next ASA cock.

Part I: Testosterone Support

The testosterone support aspect of our OCT program is substartially different than what is used during tanditom PCT. We are no brager looking to all condignones tostosterone production with anti-strogomic drugs. Buck tamoxiden or cherghinen, nor to use phrameencifical bit minis: endogroups tackizing hormoress with as IECC all phrameencifical strategies have been concluded at this point, and hopefully have decided the monosary diffects. Daring OCT, we want to provide our bodies some of the natral components used in the synthesis of testostarone. We want to augment our own natral moresses, not artificable different for the synthesis of testostarone.

Vitamin D/Calcium/Zinc

The first thing to pay special atteritor to during OCT is our vharmin and mineral stants, purchashly those components that are integrated to totatowere boburghiss. This includes Vharmi D (ackian; and Zine. To begin with, chical attatists have shown that higher levels of Vharmi D (ackian; and Zine. To begin with, chical attatists have shown that higher levels, and the original state of the anticomponent of the shown of the state of the state of the state state of the original state of the horizontal statestore outpaties. This includes its minimum state statestores of the horizontal statestore outpaties of the state statestore in the influence of the statest statestore outpaties of the statest statestore in the statestore outpaties of the statestore outpaties of the statest minimum statestore outpaties of the statestore outpaties of the statestore statestore in the statestore outpatiest of the statestore outpatiest of the statest minimum statestore outpatiest of the statestore outpatiest

D-aspartic Acid

D-aspartic scit (DAA) may also be useful during OCT. DAA is an amino acid that is nutually found in the pravots and endorcine systems, and is believed to play tools in such things as neurotansmission, spermatogenesis, and hormore biosynthesis. Chineal studies that gave 3.2 gidty of D-aspartic acid per day (as sodium asil) to hadly more neuraled in a 42% increase in serum testosterone levels in most subjects.³⁰⁰ This same dose is recommended during OCT.

Part II: Cell Re-sensitization

Repart high intensity succise, especially resistance training, causes disordion of the mucccontembrance. This disordino is in many wook solida, as it is model to initiate muccgowth and repark. Without damage, there is no prograss. There are some regardingsingtion of the muccle code, however, One of them stankathemain is furth ecourterpart disordino of the muccle code. However, One of them stankathemain is furth ecourhypoholphical are errormaged. In particular, the concentration of muchicine acid (ARR) is lowered.²⁶⁴ AIRA supports the local nubolic process.²⁶⁴ Likewise, its depletion is one of the common factors in thinging stanguino.

Arachidonic Acid

To help repletish membrane phospholiptis and restore muscle cell responsiveness to turning, anchichous acid should be applemented during the OCT period. A duby dose of 250 mg is recommended, which represents 50-100% of the normal duby distary intake of ARA. This amount should be sufficient for phospholiptin replensiment, and acceptable for hong-term use. Higher doses (500-1,000 mg per duy) may provide a more distinct musclebulatine offects the should be limited to sits o serve vecks.

Fish Oil

It may also be usefit to supplement with fish of daring the OCT period. The mmin intensist and decostnetwares and (DRA) and conseparations can (DRA) two Omga-3 searching finity acids that are also important constituents of much call metric particular Additionally, anticipation and an experimental the particular the metric analysis effects of this EFA-WA childy done of 2 grams of fish of its typically recommended dring an OFE-(yes). Humpy program.

Part III: Anabolic Supplementation

ambokizmic-tathoke properties. Many AAS users are skeptical of muscl-balding supplements, and grindfills on The muscle can be very trendfold, with even the best products filling far short of AAS in terms of efficacy and reliability. Sill, the faith hus over a partial matches spring effect drange better of the terms of the significantly alter the baselier muscle level by the start of the next steroid cycle (and this may inflatence the timing, does, or duration of AAS required). It is recommised to bin diffaces the timing, does, or duration of AAS required, it is recommised to bin database the timing, does, or duration of AAS required. It is recommised to bin database the timing and the start of the start of the next steroid cycle (and this may inflatence the timing, does, or duration of AAS required). It is recommised to bin database the timing and the start of the start of the next steroid cycle (and this may inflatence the timing, does or duration of AAS required). It is not start analysis of natural analysis, septements, place telescete Hillion Llowalysis Speer Superimour Database telescete Hillion Llowalysis.

Creatine Monohydrate

Cratite monolphtmic is regarded as the 'trojiguil' anabolic supplement, as it was the first to offer substantial performance and hosp composition improvements for most users. It is typicall picken for 8-12 works or larger (constituus fromghost the entire OCT period), as a doe of 5 gram per day. Chonten anguness much size and appendiments through several accession entructure (collair API resynthesis), although the supplement also has direct profess synthesis and an actuatobic percorrection.⁵⁵

Beta Alanine

Bea Ahine is a non-searchit animo acht that servos as a direct processor for carnosice synthesis. Darig caractesis, photgen in sur a produced in the muck-cells, which cause the pH level to drop. This proceptituses muck- fittigas. Carnosine acts as an intramocellatar buildering agart, caractering the build-up of photgen into. By serving as the ma-brining again the synthesis of mucck- carnosite, bets-altimic is a strong stabilizer of muck-tell $^{12.40}$ A does of 3-5 gams are due to thy schedul acque, which hould allow the individual to perform messarably logar during training. While this may not be a direct anabole effect, over time the increased training mindation can lead to gatter muck-preservatorization.

Branched-Chain Amino Acids

There are three essential Banched Chain Anton Acids (ECAA) – Lexine, isolatevia, and value. These armit a coils are very abunchin its olderal muscle provise, accourting for 11–18% of the total content.^{100–133} Supplementation with BCAAs is desirable for a couple of measure. The first is that they provide stegrability alleking blecks for the synthesis of neuromached proteins. First, and the structure stegrability and the structure stegrability of the structure structure

Typical OCT Program (8-12 Weeks)

Testosterone Support:

Vitamin D 3000 IU/day

Calcium 500 mg/day

Zinc Sulphate 250 mg/day

D-Aspartic Acid 3.2 g/day

Muscle Cell Re-sensitization:

Arachidonic Acid 250 mg/day

FishOil 2 g/day

Anabolic Supplementation:

Creatine 5 g/day

Beta-Alanine 3-6 g/day

BCAA 10 g/day

359. Association of vitamin D status with serum androgen levels in men. Wehr E, Pilz S, Boehm BO, Mirz W, Obermayer-Pietsch B. Clin Endocrinol (Orf.).

2009 Dec 29. [Epub ahead of print]

360. Testosterone levels in athletes at rest and eshaustion: effects of calcium supplementation. Cirar V, Baltaci AK, Mogalkoc R, Kilic M. Biol Trace Hem Res. 2009 Summer;129(1-3):65-9. Epub 2008 Dec 20.

361. Impact of cent zinc therapy on the level of sex hormones in male patients on hemodialysis. Jabii GR, Roozbeh J, Mohammadzadeh A, et al. Ren Fail.

2010 May;32(4):417-9.

362 The role and molecular mechanism of D-aspartic acid in the release and synthesis of LH and testosterone in humans and rats. Topo E, Soricelli A, D'Aniello A, Ronsini S, D'Aniello G. Reprod Biol Endecrinol. 2009 Oct 277:120.

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364. The COX-2 pathway regulates growth of atrophied mascle via multiple mechanisms. Bondesen BA, Mills ST, Pavlath GK. Am J Physiol Cell Physiology 2006 Jun; 290(6): C1651-9. Epub 2006 Feb 8.

365. Eicosapentaenoie acid and arachidonie acid: collaboration and not antagonism is the key to biological understanding. Horrobin DF, Jankins K, Bennett CN, Christie WW. Prostaglandins Leakor Essent Farty Acids. 2002 Jany66(1)(83-90.

366. Sport Supplement Reference Guide. William Llewellyn © 2009-2010. Molecular Nutrition, USA.

 Creatine in sports. Kreider RB. Essentials of Sport Natrition & Supplements. Humana Press. Totowa, NJ. 2007.

 Jake Influence of beta-almine supplementation on skeletal muscle carnosine concentrations and high intensity cycling capacity. C.A. Hill et al. Amino Acids, 2007 Feb;32(2):225-33

13C|phenylalanine. Riazi R, Wykes LJ, Ball RO, Pencharz PB. J Nutr. 2003 May;133(5):1383-9.

 Dietary protein impact on glycemic control daring weight loss. Layman DK, Baum JL J Natz. 2004 Apr;134(4):9685-735.

371. Nutracentical effects of branched-chain amino acids on skeletal muscle. Shimomara Y, Yamamoto Y, Bajotto G, Sato J, Murakarri T, Shimomara N, Kebayashi H, Mawatari K. J Natr. 2006 Feb;136(2):5295-5328.

Injection Protocols

Anabelicativganie streuci injections are always ginet doop in the macke (intramusculer), Some other performance-entirusing dang stand as human growth hormore and insidi injections are given by injection in the fit hyper between the skin and mascle (stochartmeons). The protocols for both spicetulers are provided. Improper pricediments, the shift prodocubarrent, prime injection types are provided. Improper pricediments, the shift pridocubarrent and the shift and the shift pricediment and the shift pricediments of the shift price and the shift price of the

General Preparation:

1. Make sure you have all the necessary supplies.

Intranuscular Injection:

(1) 3mL syringe

(1) 22-25g 1-1.5" needle for administering the injection

(1) 21g 1-1.5" needle for drawing solution (if using a multi-dose vial)

(2) Alcohol pads

- (1) Dry cotton ball
- (1) Plastic bandage



Syringe with needle attached. (intramuscular)

Subcutaneous Injection:

- (1) Insulin syringe with needle attached (.5-1mL 27-30g)
- (2) Alcohol pads
- (1) Dry cotton ball
- (1) Plastic bandage



Syringe with needle (insulin)

If applicable, keep the administration needle cold by leaving it in the freezer for at least one hour before opening. This will help dull the pinch of penetration.

Select a well-lit room with a clean hard surface such as a tabletop or counter to administer the injection.

4. Wash hands thoroughly with soap and warm water.

Assure that all injection equipment is sealed and unused. Never reuse needles. Double check the expiration dates on all medications.

6. Clean top of vial thoroughly with an alcohol pad, if applicable. Let air dry for 15 seconds.

Precautions: Do not use injection equipment that is used or has been exposed to air during storage. Never share needles or multi-dose vials. Discard unused portions of the drug at the recommended time.

Drawing Solution into Syringe:

1. Remove syringe from packaging. Attach drawing needle, if applicable.

If using a multi-dose vial, fill syringe with air in the amount you are withdrawing. This will help stabilize the pressure and make drawing easier.

3. If usine an annule, break onen and place flat on a hard surface. The use of a namer towel

to cover the glass top may make breaking easier. Draw solution. Skip below steps and go to Intramuscular or Subcitaneous Injection Procedure.

4. If using a vial, insert needle through the nibber stopper at a 90-degree angle. Turn the vial upside down with needle attached. Inject air: Slowly withdraw desired amount of solution. You may lightly tap the side of the needle to dislodge air bubbles. Note that small air bubbles are not harmful.

5. Remove needle and syringe from vial, if applicable. Replace cap on the end of needle.



Keep fingers away from needle when drawing.

6. Remove drawing needle and replace with new administration needle, if applicable. This is highly advised with multi-dose vials, as passage through the ruber stopper will have dulked the needle considerably. Remove any air in the ip of the needle, and prepare for injection.

7. Place capped needle back inside wrapper and place on clean surface.

Precautions: Never touch the tip of the exposed syringe, the needle, or the top of the vial stopper after it has been cleaned with alcohol. If you come into contact with these surfaces you should consider the materials contaminated, and should not use them for injection.

Intranuscular Injection Procedure:

Used for all anabolic/androgenic steroid injections.

 Thoroughly clean the intended site of injection with second alcohol pad. Preferred locations are the upper outer quadrant of the buttocks, or the outer side of the thigh. Let air dry for 15 seconds.

Remove needle cap. With free hand, stretch the skin around the site of injection with two
fingers. Move the skin over the muscle to the side by 1-1.5" (Z-Track method).

In a swift motion, insert the needle into the target muscle at a 90-degree angle with the dominant hand. Make sure the needle is deep within the muscle.

Pull back on the plunger (aspirate). If the syringe fills with blood you have hit a blood vessel, and the injection should be aborted.

5. Inject the medication slowly into the muscle.





Upper/Outer buttocks is the preferred site for IM injection.



The outer side of the leg is also commonly used.

6. Withdraw the syringe. Release the skin with your other hand. The skin and subcutaneous tissue will rebound, which helps close off the needle shaft and prevent leaking.

7. Dry injection site with dry cotton ball. Cover with plastic bandage if necessary.

Precautions: Never inject into skin that is discolored, broken, or irritated, or if there are lamps, knots, or feelings of pain in the area. Do not inject more than 3mL at one time. Rotate the site of injection so that you do not inject in the same muscle more than once every two weeks.

Subcutaneous Injection Procedure:

 Thoroughly clean the intended site of injection with second alcohol pad. The preferred location is the lower abdominal region. Let air dry for 15 seconds.





Shaded area denotes site for abdominal subcutaneous injection.

Remove needle cap. With free hand, pinch the skin around the site of injection so it is lifted off the muscle.

 In a swift motion, insert the needle into the target area at a 45-degree angle with dominant hand. Make sure the needle rests within the subcutaneous tissues between the skin and muscle.

4. Inject the medication slowly into the subcutaneous tissue. Do not aspirate.

5. Withdraw the syringe. Release the skin with your other hand.



The skin is pinched upwards to facilitate injection into the subcutaneous tissues.

6. Dry injection site with dry cotton ball. Cover with plastic bandage if necessary.

Precutions: Never inject into skin that is discolored, broken, or irritated, or if there are hamps, knots, or feelings of pain in the area. Do not inject more than 1mL at one time. Rotate the site of injection each time so that you are at least 1° away from the least site, and do not return to a previous site until all other available sites have been used. This will help prevent overses of the same injection area.

Steroid Frequently Asked Questions

1) How much weight can someone expect to gain during the first cycle of steroids?

Provided dosing is sufficient, a storoid user can expect to make the most significant progress during their first cycle. Although this will vary form person to person, it is not uncommon for someone to gain 20 pounds of weight or more during a 6-8 week period of AAS use. Some of this may be water retention, although a solid gain of more than 10-15 pounds of muscle

2) Are the gains from steroid use temporary?

Yes, and no. Steroids can help you do two basis things with regard to macke growth. First, they can allow you to more rapidly reach your genetic limits for macke growth. Provided you continue to train actively, eat properly, and use an effective PCT program, you should be able to minimize at your genetic limit indefinitely. So in this regard, the early gains do not have to be temporary.

Later, strends can allow you to peah well beyond your genetic limits. It is important to emphasize this, as externe physical development cannot be maintained long-term without the repeat administration of anabole substances. The body will always revert back towards is normal metabolic limits once AAS are removed. In this context, some of the gains will not be permanent.

Steroids do permanenty alter the physiology of your matches by adding more cellular mackies with higher mackic context, acch matches cold can armange is soulane more efficiently, which allows more rapid expansion. Elsen after a long period of complete abstiment from training and AAS, the mackies minima ¹⁷² This more provide a "tracker memory" affect, allowing you to AAS in the past. So in this enged, there are lassing benefits beyond the temporary increase in mack size size.

3) Can steroids make me look like a professional bodybuilder?

If you have the underlying genetics to allow for this extense macket growth, this may be possible with a bot of nad work and dedication. If you are like the vast mjority of people, however, steroids will not be able to make you look like a professional hodybaldker. Genetics are a big factor in determining the ultimate limits to your physique, even in an erhanced state. Many people use storoids and look very big and impressive because of it, but very fow users and hot make it to the stage of a professional hodybalding show.

4) How dangerous is an isolated cycle of steroids?

Anabolicantoganis starolis are among the safet days soubhile, at least in a short-term same. Fail overloos in on tensorubly possible, and the rangets headih changes such as abantions in choolestered, blood pressue, hermater, and blood cotting (among other things) are very miledy to mentific its neizons body humor of dust fater a noised cycle. There are me douth from such things as stroke and low cancer in short-term abasers, but such cournerces are statistically enterwhy areas if light of the millions of poople that use these drugs. If you had to companifiedly nucle that the state risks of AMS abase, they would be slightly hisher than mutiques. Just far less situ nivilaral of other list mercices.

5) How dangerous is long-term steroid use?

The long-term use of steroids for momentical reasons can be a significantly unreality practice. It has been differ, however, to quark for each ref. It men insise it is the that AAS hases can provide learn discusse, the number one killer of neur. Heard fasces is a mean provide the steroid steroid steroid steroid steroid steroid steroid steroid in the cardiovascular systemization in the entries, and also induce other charges in the cardiovascular systemization and the random cardiovascular discussion. The uset means the cardiovascular systemization in the entries, and also induce other charges in the cardiovascular systemization in the entries, and the single cardiovascular discussion. The uset means the cardiovascular systemization is the entries of cardiovascular discusse. The uset means the steries of t

6) Can steroids be used to enhance an athletic career safely?

The nonmedical use of AAS by definition cannot be defined as a safe practice. However, it

can be argued that anaboki-indipants storoks can be used with high relative safety, oten over a point of mmu years. The galakies of storoki hum mediation are importune to minimizing the negative health effects of these dengs. Provided an individual folkows these againshies and a cereful with dng selection, doogs, and duratines of ratice, folkows a diftor in stantand fate, choiseand, again, and artificial carbidy that any story of the provided and the start of the health risks, thus a speed start of molecular and planning to use AAS in this murner, which is about a stored.

7) What are the safest steroids for men?

Teasotence, whatever the form, tends to be the stated starvid for men. When the dose remains with the moderately aparthermopric marge (such as 200-400 mg of an injectable testostence ester per week), alterations in cardiovascular risks factors are noticed, bet rate varients. Store of the loss to do with the beneficial audiovascular dicks of estingation in marmeterachone. Isolating your use to these dangs is recommended over using the fall spectrum of on land niterateble streoks.

8) What steroids will not cause hair loss?

For those with a garctic predisposition to hair loss, all anabolic/indequies storoids are compleb of calcerdinating the process. Showing the conset of this darks. All sets requires a faces on reducing relative antiogenicity in the scah. This can be accomplished with the use of predominity anabolic dang such as a matrichon, construction, or methenoloce. Alternately, modernet doses of testostorence can be used with finasteristic, a drug that reduces DHT conversion (end antiogenic amplification) in the scah. Silt the segmetrical process that because the set of the se

9) What are the safest steroids for women?

Women are generally most concerned with the viribing (mascalinizing) effects of anabolic/androgenic steroids. The least viribing agents are frose with the highest relative anabolic to antiogenic effect, such as randrolone, on antherhonione. Care must always be taken, however, as all AAS are based on mule sex steroids, and as such can case muscalinizing effects in sources.

10) Should I rotate my steroids every few weeks to prevent receptor downregulation?

No, this is not necessary, Anabolican/openic steroids all work primarly by stateling to and activity and sense receiptor. As such, you do not gain anything by solviding to a new compound that works via similaring the same receiptor. If theirance were induced by one ASS compound, it would be extended to all componds. The plateau effect that is noticed 6.4 works into most cycles is poorly understood, but fidely related to the new metabolic lines placed most constrained and the sense of a contrait ASS dosagas, not insensitivity or ASS. Choise downregatizin does not occer with these drugs, and even if a did, rotaring stronik world not prevent it.

11) How likely am I to find real steroids on the black market? Does it matter?

Albody coart fagres are difficult to calcular, real planmeoridial ambidedinologonic strends are estimated to represent all for sizes of the products commonly civatilet of the black market. In many regions this fagre may be below 29%. The majority of products soil properly are counterfact copies of real AAS, or pordasts mode all tabled by underground laboratoriss. It does matter, because the quality of normedical AAS cannot be ensued. These products are greanly not advised for the c. Gennet potential issues with mag astry: it is worthholic to spend the extra inte and money on stored products you can be assard care from lagitime quanterfacts.

12) What do the anabolic and androgenic reference numbers under the profile for each steroid mean?

These numbers come from early studies measuring the effect of each steroid on certain mack and sex organ tissues of animals, usually mice. These numbers are useful for assessing the relative anabodies to androgene bolaine of each drug in humars. They are not as accurate at assessing the total maske building potential of each steroid, however, and should not be taken as absolute mings of potency.

372 Anabolic Steroids Withdrawal in Strength Trained Athletes: How Does It Affect Skeletal Muscles?," Anders Eriksson and Las-Eric Thornell. American Physiological Society. The Integrative Biology of Exercise V, Sep 24-27, 2008 Hilton Head, SC.

373. Increased prenature montality of competitive powerlifters suspected to have used anabolic agents. Pärssinen M, Kujala U, Vartiainen E, Sarna S, Seppilla T. Int J Sports Med. 2000 Apr;21(3):225-7.

Understanding Blood Tests

The abuse of analotic/antiogenic strends: can have a marker of potential negative health consequences, most commonly with regard to cardion-stark and loss the health. These issues, however, can aimst always be identified in blocd work will before physical opproxemprograms. Chalsens and an influenced testings can be used to leafer almost and consel early the factors and decisions that workd support the disease over time. Liver damage is also agreenable dynamic there earners tasks will before it heccenses which motions that the constraints of the early the strength of the early motion of the early motion to the early operative strength of the early operative strength of the early motion of the early operative parameters and the early operative strength operation of the early motion of the program devices and the motion of the early for motion of the early operative of the task and the motion of the early for motion operative in the strength of the early operative operative strength operations in the motion of the strength operative strength operations in the motion of the early operative strength operations in the motion of the strength operative strength operations in the motion of the strength operative stre

Blod tests with regard to analycic/introgenic storoids are usually conducted in three separate pitess. The for phase looks at your hash here's around use. This is done to assess your current condition and risks before any therapy is initiated, and to set baselines for later comparison. The next is on-cycle testing, which is used to assess the detect impact of the analoxiciant/cogene atorial use (which the drugs are astudy) doing to your hody while they are bring laken). This target frame of starsing is the follow-up, which is to constant to remove the gradient pitest of the set of the stars of the stars of the stars of the stars of the set of the We generally vefer to these three phases of testing as Baseline, On-Cycle, and Post-Cycle, respectively.

Baseline (Pre-cycle):

Bandau (pre-ycch) toring is generally very broad. This is done to make sare there are no much/steps that in contrains that may be sourced by starking starking contrains, and to hirse a storoid hornexes (mink users only is done to kinety) the current matrix stars. This can be expectively protout the lows for post-y-cch who use a not many stars in the staring of the store stars of the stars of the stars of the store stars of the store stars of the store stars of the stars of the store stars of the store stars of the store stars of the stars of the store stars of the store stars of the store stars of the stars of the store stars of the store of hepotencies can be stored as the store star of the store stars of the store of hepotencies can be stored as the store stars of the store stars of the store of hepotencies of the stars of the store stars of the store stars of the store stars of the stars of the store stars of the store stars of the store stars of the stars of the stars of the store stars of the store stars of the store of the store stars of the stars of the store stars of the store stars of the store stars of the stars of the store stars of the store stars of the store stars of the store stars of the stars of the stars of the store stars of t

Checklist (minimum):

Hormone (Steroid)

Lipids (Standard Full Set)

Full Liver Panel

Blood

Kidney

Electrolytes, Minerals, and Glucose

On-Cycle:

On-cycle stering is unally conducted 3 to 4 weeds after stored therapy began. The initialiant all agramph look at these initiacions of heath most directly stored with the store use. All high camminon is concluded, and is other segreded as the single nont important set of heath heat start can be instand. It is here that the candroscated irrest of the stored with begin to become approach. One should give special consideration to what these results or special many starts and the store of the store of the store of the store system of the store and the store of the store of the store of the store of the with begin stored and the store of the with the camming. It is also recommended to examice other general health markers here such as block likely, excludes, minutes, and galooo.

Checklist (Minimum):

Lipids (Standard Full Set)

Liver Panel, if taking hepatotoxic steroid(s)

Blood

Kidney

Electrolytes, Minerals, and Glucose

Post-Cycle

During the post-cycle testing phase is is common to exec again look for at the mist struct hormones. The hope how is to obtain values that clookey mirrory comprementarie look. Note that three will always be some variation based on the time of the day, and coven in the day-to-day-results. A near eart much is problem for fassible. It is also considered ago of that to look at phatmy LH and FSH, because if stotstorene levels cores back how it will give you and your physicani is hetter understanding of the cause. High LHSSH and look isotaneous optimizery that the physicanal material structure levels can be how the visit isotaneous optimizery that the construction optimizer of the cause of the structure optimizers of the structure optimizers of the structure optimizers of the mire optimizers of the structure optimizers optimizers of the structure optimizers optimizers

Checklist (Minimum):

Hormone (Steroid, LH/FSH)

Lipids (Standard Full Set)

Liver Panel, if taking hepatotoxic steroid(s)

Blood

Electrolytes, Minerals, and Glucose

Blood Tests by Category

Hormone

Steroid (male)

Steroid (male)

| Test Name | Reference R | Reference Range | |
|---------------------|--------------|------------------------|--|
| Testosterone, Total | 241-827 ng/d | | |
| Testosterone, Free | 8.7-25.1 | pg/mL | |
| Estradiol | 10-53 | pg/mL | |

LH/FSH Panel (male)

| Test Name | Reference I | Range |
|-----------|-------------|-------|
| LH | 2.5-9.8 | IU/L |
| FSH | 1.2-5.0 | IU/L |

Thyroid

| Test Name | Reference Range | |
|----------------------|-----------------|--------|
| TSH | .35-5.5 | ulU/mL |
| Thyroxine (T4) | 4.5-12.0 | ug/dL |
| T3 Uptake | 24-39 | 96 |
| Free thyroxine index | 1.2-4.9 | |

Steroid:

This set of testing should look at both total and fee tostosterore. The former measure is most commonly used by physicians to identify the ant/open iced and detomice filture is a need for therapy. The later measure actually expressers the faction of bioxualable (invendited) active) tostosterone in the body, and is consequently regaled as more important for assessing the present state of anthogravity. Estable is the principle actue form of entropy in the body, and its notice specific the state of the body, and low respectively. This is the estrogen marker most often recommended arthre phomese poling.

LH/FSH Panel:

Lucieizing hormore (LH) and folicle strinulating hormore (PSH) are responsible for simulating testosterone production and spermatogenesis in the testes. These measures are most relevant when evaluating the cause and potential treatment options for hypogenulation, not the short-term health impact of ambolic-steroid use. The short-term suppression of LHFSH is expected with ambolic-molecular duministration.

Thyroid:

It is regarded as important to get a baseline measure of thytoid activity, usually once per pair. Follow up to taking and after stretoid use my to be an expense some view as intracessary. Antabolicandrogenic storoid use is utilikely to permanently affect flyioid factoria, har may sightly elsous through levels drang therputy. A mislingnosis of the effect of antabolicandrogenic storoid use on thytoid levels should be taken into account before teament for hyperdprovid scored.

Lipids (Cardiovascular)

Anabolicantiogenic steroids can have strong advence effects on lipids. The abuse of anabolicanticogenic stressift operator have can achieve the stress of the stress for developing cardinoscelar denses as assessed by three variables. Mitigating these rails, for developing cardinoscelar denses as assessed by three variables. Mitigating these rails and activati-citedite block can be able for more production of the stress of the stress based bound on provide a good stressift of the stressift of the

Standard Full Set

| Test Name | Reference Range | |
|-------------------------------|-----------------|-------|
| Triglycerides | 0-149 | mg/dL |
| Cholesterol, Total | 100-199 | mg/dL |
| HDL Cholesterol | >40 | mg/dL |
| VLDL Cholesterol | 5-40 | mg/dL |
| LDL Cholesterol | <100 | mg/dL |
| LDL/HDL Ratio | <3.6 | |
| LDL/HDL Ratio Risk Assessment | men | women |
| 1/2 Average Risk | 1.0 | 1.5 |
| Average Risk | 3.6 | 3.2 |
| 2X Average Risk | 6.3 | 5.0 |
| 3X Average Risk | 8.0 | 6.1 |
| | | |

Additional Testing

| Test Name | Reference Range | |
|---------------------------|-----------------|--------|
| C-reactive Protein | <5 | mg/dL |
| Homocysteine (0-30 years) | 4.6-8.1 | umol/L |
| Men (30-59) | 6.3-11.2 | umol/L |
| Women (30-59) | 4.5-7.9 | umol/L |
| >59 years | 5.8-11.9 | umol/L |
| Apo Ratio Testing | | |
| Apolipoproteins | men | women |
| apoB/apoA-I Ratio | <.9 | <.8 |
| Apo Ratio Risk Assessment | men | women |
| Low Risk | <.7 | <.6 |
| Average Risk | .79 | .68 |
| High Risk | >.9 | >.8 |

Standard Full Set:

This is a standard full lipid pranel examination. Idealby, all values should be kept within the romand ranges at all times during stored therapy. Note that the LDL/HDL ratio is regarded as the most important measure of the serum lipid tests, as it reflects the ongoing balance between plaque deposition (LDJ and removal (HDL) is the atterists. The LDL/HDL ratio is used to more closely assess heart disease risk in individuals that have elevated LDL or total cholesterol levels.

Additional Testing:

C-reactive protein and homocysteine are two additional markers that are important to examining cardiovascular health. C-reactive protein is a key indicator of inflammation in the body, and homocysteine is involved in blood clotting and LDL cholesterol oxidation. It is also advisable to include these two variables in your cardiovascular testine schedule.

Apo Ratio:

Te

Apolopowień mós testię is alio recommended. Alkough net commedy used in general medical practice, opolopowien testię is ie resussify negatuda sa new excante predicto da LDL particis, ami is responsible for attaching these lipowients the attaching and LDL particis, ami is responsible for attaching these lipowients to the artey work. Apolopowień A-l (apoA-l) is found minity in HDL particles, and is responsible for miniting bancial meter choisecont tampoort. ApoA-l enables the HLT, particles to particle and the state of th

To Find A Testing Lab Near You, Visit: www.HRT-Labs.com Liver Function

| st Name | | Reference Range | |
|---------|----------------------------|-----------------|-------|
| | Albumin | 3.5-5.5 | g/dL |
| | Globulin | 1.5-4.5 | g/dL |
| | Total Protein | 6.0-8.5 | g/dL |
| | Bilirubin | 0.1-1.2 | mg/dL |
| | GGT (Gamma GT) | <50 | IU/L |
| | ALP (Alkaline Phosphatase) | 25-150 | IU/L |
| | AST (SGOT) | 0-40 | IU/L |
| | ALT (SGPT) | 0-55 | IU/L |
| | | | |

A fill her parels in protent to assessing happing staria. The two markers of Hor stress most commonly elosted in abases of antholicothandpongis stressing as the enzymesa aliane amino-transformes (ALI) and separate amino-transforme (AST). ALI and AST are recessrry to amino accl metabolism in the low, and will keel its of be blockstream as the low between the stress of the stress of the stress of the stress of the stress indicators of ently stressical. These two eraymets are generally regarded as important minicators of ently stressical characteristic of the stress in which shistering low end endly acceleration of the erargent wates is abayes abised.

Alaine phosphanse (AIP) and agramm-datanyhranpeptikase (GNOT) he arown as colosattic for enrargoss, which mean they dimited to estype the Work (a goverith hild that that discussion and is produced in the keys AIP and GGT are important markers of Hore headh during stored law, and hould be ixcluded in regater biod tearing Evastorian AIP. and GGT can indicate bied durit obstruction (transported colosatiosis), which refers to a condition where the fore can no longer proporty transport and markelose bie. Intransport cholestasis is a potentially very strans marifisation of storoid-raikeed line trackets, so eleatorians AIP and GGT should not be disognated.

Mål elestinos in ALT and AST my be caused by mack- damage (courcie) and not storois-induced her toxiday. A comparison to baseline keek wil be important in determining the cause. If the only factor that has changed is the addition of a hepatotoxic anabole stereof training is observed seataboly, the drugs il sheet to bane. It is important to remember that ALP and GGT are not always elestated with early her starts. Therefore, the destine of any hepatic mathema show the reference range (costs (rob ALT) and AST) can always any start of the start of t marcane nver toxicity, and smould be cause to discontinue the outerking steroids and reassess risk.

Muscle Enzyme

| Test Name | Reference Range | |
|-----------------|-----------------|-----|
| Creatine Kinase | 38-174 | u/L |

The creative krane (CK) expose is used as a matter of muscle breakdoon, kikey dumg, and head dumg. High locks using histical heat states, or other ogen narmar. This enzyme can also become deviated with exercise that breaks doon muscle tissue, especially stresse enhance or resistance straining. Evalet CK, beek caused by high intensity training are often residue for ogens dramge. It is inportant to further examine other tissue becken ways do be used in determining. There at the VK is the strain tissue becken ways do used in determining. The strain to heavy turings is the cause of mult deviations in how response ATI and AST. Skipt neurons in ATI and AST caused by muscle dumg wild based ovicies that the bread CK and neural ATI and GKT locks.

Blood

| Test Name | Reference Range | |
|----------------|-------------------|-------|
| WBC | 4-11 | K/MCL |
| RBC | 81-103 | FL |
| Platelet Count | 130-400 | K/MCL |
| Hemoglobin | 13-17 | g/dL |
| Hematocrit | 40.7-50.3 (men) | 96 |
| | 36.1-44.3 (women) | 96 |

A fill block court is one of the most commoly ran block tests, and can give you a good supplot of overall healts in mury regards. A fill block cell ster will give you a measure of white cell court (responselse for fighting indication, planket court (viata lo block claring and healing), and al block cell court (responselse for carring course). Rel and white cocoarts will be further subdivided into various individant measurements, often informed to as a discussion of the further subdivided into various individant measurements, often informed to as a manusce of the interval subdivided into various individant measurements, often informed to effects on explorepoints, annihold stansis stands tatal to increase on blocd coll courts. Then to the effects on explorepoints, mathodic stansis stands tatal to increase on blocd coll courts. Then to the mathematical measurements of the block cells in texas table long (actobic) capacity, as the concentration offend blocd cells increases so does the thickness of the blocd. Elsewide hemateria can increase the risk of hemat tatacet or stroke.

Kidney

| Test Name | Reference Range | |
|----------------------|-----------------|-------|
| Uric acid | 3.0-7.0 | mg/dL |
| Creatinine | .5-1.5 | mg/dL |
| BUN | 5-26 | mg/dL |
| BUN/creatinine ratio | 8-27 | |

This parel of tests boks at three primary waste products filtered and exceted through the kitches, urea, ari, ari cell and certaintie. Problems here can indicate serios underlying problems with kitchey function. Note that Blood Urea Nitogen (BKN) is often elevated with excess protein consumption, and is used by marg physications as an indicator that too much protein is being consumed for optimal metabolism. The high consumption of must or certainte supplementation can also elevater certaintie levels, diministing the value of blood creatinine supplementation can also be elevater certaintie levels. Attimisting the value of blood creatinine supplementation can also be elevater certaintie levels. Attimisting the value of blood creatinine supplementation can be observed certainties levels. Attimisting the value of blood certainties and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood creating and the supplementation can be also certainties levels. Attimisting the value of blood certainties attimisting the supplementation certainties and the supplementation certainties and the supplementation certainties attimisting the suplementating the supplementating the supplementation certainting testing as a marker of kithny health. Electrolyte, mineral, and fasting glacous testing is important to farther assessing kithny health, and is advised in addition to the above kithney markers. A quick tarine screen for pH, specific gravity, and the presence of sugar, blood, and ketones is also available at most physicians' offices, and is generally advised alongside blood work when possible.

Electrolytes, Minerals, and Glucose

Tes

| 51 | Name | Reference R | ange |
|----|------------------------------|-------------|--------|
| | Sodium | 136-146 | mEq/L |
| | Potassium | 3.6-5.2 | mEq/L |
| | Chloride | 98-109 | mEq/L |
| | Bicarbonate (carbon dioxide) | 21-30 | mEq/L |
| | Phosphorous | 2.5-4.5 | mg/dL |
| | Calcium | 8.5-10.5 | mg/dL |
| | Iron | 35-185 | mcg/dL |
| | Glucose (fasting) | 70-110 | mg/dL |
| | | | |

Exercise levels are canned to help detect problems with the flaid and electrolybe balance. Abcornel values my reflect someting as sometim or possism dividency, or a more serious contition such as idshey disease. A variety of other health issues my also become apparent by looking at hot divectorised and mixed levels, skying them someshul broad proposite value. Fasting glacos is also canneed to determine if the infidukal any to hypothysenic fluctuations relating methodes systems and sharing glacoses my reflect potentially writes health conditions relating methodes systems. **Postster**

| Test Name | Reference Range | |
|------------|-----------------|-------|
| PSA, serum | 0.0-4.0 | ng/mL |

Postate-specific arigan(PSA) is a protein produced by colis in the prostate gland. Its lovels can become elocated in cases of benign protein legislation protein earness while its remains unknown of elocating the level of antidogens in the lody with anabolicatindogenic strends, can increase diagonics tool for sciences, it is known that this disease can be progressed by elocated hermore (androgen and entroppi levels. The PSA test is regarded as an imported migroundies tool for sciencing infoldand prostate careful. While its elocating and the signification of the science of the science of the science of the elocation more will able against using antibolic indrogenic storieds. Individual Heath Markers Defined

Alanine anino-transferase (ALT): An enzyme produced primarly in the liver but also in other tissues. ALT is involved in arnino acid and protein metabolism. Used as a primary marker of hepatic strain. Also called Serum (Klatanic Pyruvic Transaminese (SGPT).

Albumin: The main protein that circulates in the blood. Produced in the liver and has artioxidant properties. Transports certain hormones, vitamins, and minerals, and plays a role in water balance. Used as an indicator of liver health. Higher levels are optimal.

Alkaline Phosphatase (ALP): A family of cholestatic enzymes produced mainly in the liver, but also in the intestines, kidneys, and bone. Used as a marker of hepatic strain, often relating to disease of the ble ducts.

Apolipoprotein A-I (apoA-I): A constituent of HDL (good) cholesterol, apoA-I is responsible for initiating beneficial reverse cholesterol transport. This process pulls cholesterol particles from the artery walk and transport them back to the liver. Higher levels are optimal. Apolipoprotein B (apoB): A constituent of LDL (bad) cholesterol, apoB is responsible for attaching these lipoproteins to artery walls. ApoB is a promoter of fatty plaque deposits in the arteries. Lower levels are optimal.

Aspartate anino-transferase (AST): An enzyme produced primarily in the liver but also in mascle tissue. AST is involved in amino acid and protein metabolism. Used as a marker of hepatic strain, although it is considered less specific than ALT testing. Also called Serum Gharnio-Coalcoteir Transmiruse (SGOT).

Basophils: A type of white blood cell. Action not fully understood, but cells are known to carry histamine, heparin, and serotonin. Levels are elevated with allergic reaction and parsitic infection.

Bicarbonate: A measure of carbon dioxide content in the blood, and a common marker of the acid-base balance.

Bilinubin: A waste product made from the breakdown of red blood cells. Excreted into the bile. Regarded as an important indicator of liver health. Elevated levels in the blood indicate liver toxicity.

Blood Urea Nitrogen (BLN): A vaste product from the breakdown of proteins; filtered and excreted through the kidneys. Elevated levels may indicate a namber of problems including excessive protein intake, kidney damuga, dehydration, heart fahre, or reduced production of digastive enzymes. Low levels may be indicative of many things including mulatition of twee dramae.

BUN/Creatinine Ratio: The ratio of Blood Urea Nitrogen to Creatinine, used as a marker of kidney and liver health.

C-reactive Protein (CRP): A key marker of inflammation in the body. Elevated levels may indicate increased risk of cardiovascular disease or stroke.

Carbon Dioxide (CO2): Byproduct of respiration, and a common marker of the acid-base balance. See also Bicarbonate.

Calclum: Electrolyte involved in a myriad of body functions including bone metabolism, protein utilization, muscle and nervous system functioning, cardiovascular functioning, blood cloting, and nutrient transport.

Chloride: Electrolyte involved in the regulation of water bahnee. Elevated levels may indicate a number of things including anemia, delydration, excess salt consumption, and hyperthyroid. Low levels may indicate heart or kidney failure, severe vomiting, or a number of other health conditions.

Cholesterol, Total: A measerure of all fractions of cholesterol in the blood (LDL, VLDL, and HDL). High total cholesterol is regarded as a risk factor for cardiovascular disease.

Cholesterol, HDL: A measure of the beneficial high-density lipoprotein (HDL) fraction of cholesterol, which helps remove plaque deposits from arteries. High levels are optimal. Low levels may be found in cardiovascular disease.

Cholesterol, LDL: A measure of the low-density lipoprotein (LDL) fraction of choksterol. This is the primary atherogenic particle, meaning it tends to promote the formation of plaque deposits in the arteries. Low levels are optimal.

Cholesterol, VLDL: A measure of the very low-density lipoprotein (LDL) fraction of cholesterol. VLDL contains the highest amount of triggeerides. Considered an atherogenic ("bad") cholesterol particle. Lower levels are optimal.

Cholesterol, LDL/HDL Ratio: A measure of the primary atherogenic particle (LDL) in relation to the primary antiatherogenic particle (HDL). This ratio is generally considered the most important cholesterol test value for assessing cardiovascular disease risk. A low ratio is desirable.

Creatine Kinase: An enzyme found largely in the heart and muscle, and responsible for

converting creatine to phosphocreatine. Elevated levels may be linked to a number of things including heart attack, kindey failure, or sever muscle damange.

Creatinine: A waste product of muscle metabolism. Low levels may indicate kidney disease, mainutrition, or liver disease. High levels may indicate a number of things including reduced kidney function or muscle degeneration. Creatine supplementation may also elevate creatinine levels.

Eosinophils: A type of white blood cell. Similar to basophils, eosinophils are used by the body to protect against allergy and parasites. Levels are elevated with infection, and are low with good health.

Estradiol: The principle active form of estrogen. High levels can be associated with water retention, fat buildup, and gonecomsatia (men). Also plays a role in prostate hypertrophy. Low levels of estratiol may be associated with increased heart disease risk.

Follicle Stimulating Hormone (FSH): A pituitary hormone involved in reproduction. In men, FSH is mainly responsible for supporting spermatogenesis. In women it supports ovulation.

Gamma-Gutamyl Transpeptidase (GGT): A cholestatic enzyme produced in the bile ducts. GGT is involved in glutathione metabolism and the transport of amino acids and peptides. Used as a marker offlepatic strain.

Gabulin: A blocd protein similar to albumin. Globulin is responsible for transporting certain hormones, lipids, metads, and antibodies. Levels may be elevated in many confidients induding chronic infections, liver disease, arthritis, cancer, or lapus. Lower levels may be found with a number of conditions including suppressed immune system, malnutrition, mablisoption, and liver or kinely disease.

Guesse (fasting): Gacese is the product of carbolydate metabolism and the primary source of energy for most cells in the body. Fasting glacose levels are elevated in a number of conditions incluing diabetes, her disease, metabolic syndrome, parareatitis, diciting, and stress. Low fasted glacose levels may indicate liser disease, overproduction of instifu, hypothyroidism, or other disease.

Hematocrit: A measure of the percentage of red cells in the blood. Low levels indicate an anemic condition. High levels may indicate a number of things including dehydration, increased red cell breakdown in the sphere, cardiovascular disease, or respiratory disease. Anabolic steroids may also increase hematocrit.

Hemoglohin: A constituent of red bloed cells, and the main carrier of oxygen and carbon dioxide in the blood. Lovels may be suppressed with a number of confilions including multitrition, multibociption, and anemin. High levels may indicate many timps including dehydration, cardiovascular disease, or respiratory disease. Anabolic steroids may also increase hemoglobin levels.

Homocysteine: A compound formed from the metabolism of the amino acid methionine. Involved in blood clotting and LDL cholesterol oxidation. Elevated keels of homocysteine indicate aminereased risk of cardiovasculur disease and stroke.

Imm: Mneral necessary for many functions itselding the formation of hemoglobin and certain proteins, and the transport of oxygen. Elevated levels may be caused by many confiltions including certain forms of anemia, leve damage, hepatits, iron poisoring, or vitamin B6 or B12 deficiency. Low levels can indicate a number of fittings including gastrointestial blood loss, heavy mensional bloeding iron multihostropic on deficiency.

Lactic Acid Dehydrogenase (LDH): An intracellular enzyme found in many tissues including the kithey, heart, skeletal muscle, brain, liver, and lungs. Used as a marker of tissue damage. High levels are found in many conditions including heart attack, anemia, low blood pressue, stroke, liver disease, muscle injury, muscular dystrophy, and pancreatis.

Luteinizing Hormone (LH): A pitutary hormone responsible for the stimulation of testosterone production in the testes (men). LH primarily supports ovulation in women. Lymphocytes: A type of white blood cell. Primary role is to fight viral infection. Levels are elevated with active infection. Low levels are associated with suppressed immune system or active bacterial infection (noted by elevated neutrophils).

Mean Corpuscular Volume (MCV): A measure of the size of red blood cells, determined by measuring the volume of a single red blood cell. Useful in determining the cause of anemia. Elevated levels may reflect a number of things including a deficiency of vitamin B6 or folic acid. Low levels may reflect ion deficiency, or other causes.

Mean Corpuscular Hemoglobin (MCH): A measure of the average weight of the hemoglobin in red blood cells. Useful in determining the cause of anemia.

Mean Corpuscular Hemoglobin Concentration (MCHC): A measure of the average concentration of hemoglobin in red blood cells. Useful in evaluating the cause of, and therapy for, anomic. Low levels may indicate blood loss, B6 or ion deficiency, or other causes.

Monocytes: A type of white blood cell. Primary role is to fight severe infection not sufficiently countered by lymphocytes and neutrophils. Levels can be elevated with a number of things including chronic infection and certain cancers. Low levels indicate good heath.

Neutrophils: A type of white blood cell, also known as granulocytes. The primary white cell used by the body to fight bacterial infection. Levels are elevated with infection. May be suppressed with compromised immune system or bone marrow.

Phosphorous: An abundant electrolyte involved in a number of body functions including the utilization of carbolydrates, fats, and proteins for cellular mainterance, repair, and growth, the production AIP for the storage of cellular energy, the transport of calcium, the mainterance of somotic pressure, and the mainterance of heartheat regularity.

Pintelet Count: A measure of the concentration of platekis (also known as thrombocytes) in the blood. Platekis are involved in blood clotting, and protect against excessive bleeding. Elevated levels may be linked with a number of things including delaptation. Low levels are found in suppressed immune system functioning, drug reactions, or deficiencies of vlamin B12 or folia exil, or may have other cases.

Potassium: A key electrolyte necessary for nerve and muscle function, and the transport of nutrients and waste products in and out of cells. Along with sodium it helps maintain the acidbase balance and osmotic pressure. High levels may be caused by a number of things including kithwy failure, metabolic or respiratory acidosis, and red blod cell destruction.

Prolactin: A reproductive hormone involved specifically in lactation. Prolactin is sometimes (but not commonly) elevated in steroid abusers, and may be linked to estrogen excess or hormone inbalance. Elevated productin may also indicate other issues with the pinutary.

Prostate-specific antigen (PSA): A protein found in prostate cells. Used as a screening for prostate cancer risk. Elevated levels reflect an increased risk of developing prostate cancer. Low levels are desirable, although do not assure against prostate cancer.

Red Blood Cell Count: A measure of the total concertration of red blood cells, responsible for transporting oxygen and carbon dioxide in the body. High red cell counts are seen with a number of conditions including heart disease, delpdatation, or pathronary florosis. Low levels may be linked to many times including anemia, bone marrow failure, red blood cell distruction, bledenig, levkenix, and multation.

Red Cell Distribution Width (RDW): A measure of the variation in size between red blood cells. Useful in evaluating the cause of, and therapy for, anemia. Increased values may indicate a number of things including vitamin B12, folic acid, or iron deficiency.

Softum: An abundant electrolyte necessary for many functions including the maintenance of osmotic pressure, acid-base balance, and nerve impulse activity. Disturbances in the softiam level may be caused by minor things including excessive sweating, vomiting, diarthea, water intake, or very serious conditions including bart, kidney, or fiver disease.

T3 Uptake: This test measures the level of unsaturated thyroxine binding globulin (a carrier

of thyroid hormones) in the blood, increased levels may indicate a number of timps including hyperthyroidismi (overactive thyroid), liver disease, cancer, and decreased lung function. Low levels may be indicative of hypothyroidism (under active thyroid), excess estrogen levels, pregnarcy, or other causes.

Testosterone, Total: The measure of both unbound (active) and bound (inactive) portions of testosterone in the blood.

Testosterone, Free: The measure of free (unbound) testosterone in the blood. This represents the total amount of testosterone immediately available to tissues.

Thyroid-Stimulating Hormone (TSH): A pituitary hormone responsible for stimulating the release of thyroid hormones.

Thyroxine (T4): The more abundant of the two major thyroid hormones (T3 and T4). T4 serves mainly as a reservoir for the more active thyroid hormone (T3), which helps to stabilize and regulate thyroid supply. This is a key marker of the state of thyroid health (low, normal, or overactive).

Thyroxine, Free Index: This measure is a calculation of the amount of unbound (free) T4 in the blood. This is a key marker of the state of thyroid activity (low, normal, or overactive).

Total Protein: A measure of the total serum protein concentration, maisly abunin and alphdmi. Serum proteins are important to the fanction and sapply of enzymes, hormous, nativities, and arthodies, and also play a role in muintaining the water and pH balance. Low levels may indicate a nartheor of things including mathematication, faser disease, disense, or severe bum injury. Elevated levels may indicate infection, lover disease.

Triglycerides: The main storage form of fatty acids in the body. May be metabolized and used for energy. Elevated triglyceride levels may contribute to hardening of the arteries (atheroseleves), and increase the risk of heart disease or stoke. Low levels are optimal

Urea: (see Blood Urea Nitrogen)

Uric Acid: The waste product of purine metabolism, which is filtered and excreted through the kichneys. Elevated levels may indicate a number of things including gout, infection, kichney damage, and excessive protein intake. Low levels may indicate kidney damage, malnutrition, liver damage, or other causes.

White Blood Cell Count: A measure of the total concertation of white blood cells (also known as leukocytos), responsible for fighting infection and protecting the body from pulnogens. A differential measure of white blood cells is usually also taken including neutrophils, cosisophils, basophils, hyrphocytes, and monocytes. Levels may be elevated with certain infections or allergic conditions.

Harm Reduction / Safer Use Guidelines

Hem reduction is a concept menu hashness vorkens that socks to reduce the negative hashnessnessness of ding above. The privileysion Human reduction off for an accepture of the first fung good or bad, life drugs exist in tody's society, Instead of ignoring drug uses, Human reduction particineses acriedy work with them to permete soft use strategies and decrease the hashness drugs of drug abase. The effort of human reductions is abays helping, not judging the individual. Although privately focus decreases from the soft of hyperbags and principations acriedy works with focus permetes and uses group that merely has the baseful of thal physician consider in a drugs programs. In an orditer to influent the signal ABADICIS has conditioned the fidologing principal of started linear mechanism (Fidowed, these principles should measurably reduce the negative health impact of started use, maining a start (abady not completely single sortice). 1. Avoid Counterfeit and Underground Sterolds. Anothele stored by Bic mundatorses are only on going and muy present additional bach risks to be used by Bick in the presented by the stored is more than the presented by the stored is the stored in the stored in the present additional back responses and the stored in the s

2. Avaid Toxic Onal Sterviski. Askie from Archick Primobolus and Province, very ond storid discost in the inference book is a cit-1 apha adaptied compound and should be avoided whenever possible. While there may be a nutriffer of clinical reasons to prescribe such drug, when used in the light of some sources of the mack power of the signal stores to creation head multi-more there are not included to some source of the signal stores to the source of the signal stores of the signal stores the source of 10L (sold to 10H L) could be offlated. The most nuclear distribution of the signal stores to the distribution of the signal stores the source of the signal store of the signal store.

3. This of Testosterene First. Of all the ambdo-knrhogsnic starsits produced, testostence esters its coptonet, currentlar, and Sastanto tedi to hue the busset negative instanti on hashi when takon in mack bakling and performance-mhuring dose. Estastence drags provide a horness effective all the starsite produced in the body presenting the same spectrum of physical and physiological effects. In addition to being one of the most efficient muck-hashion studies, its startering generations and the same share the startering and the same startering and the

4. Linki Vanoreffo the "Safest" Drogs. If the exclusive use of an ispectible isotanceue is not famile, imprive five a first set of the safe product of the isocable close in the following drogs have the lowest catadiouscular status and an recommended. Dess-fits biological set of the safe set of the

5. Use Health Support Supplements. Antiolecintrogenic actorial cues can help hower the regardies health integre of storial to see with the construction of trannal health appendix setup of the stories and the store of the st

6. Always Cycle Steroids. A steroid cycle usually consists of 6 to 12 weeks of drug use followed by an equal period of time or more abstaining from all anabolic/androgenic steroids. Interpretere a antonen tor a numerer of reasons, ror one, is you supperment must server hormones your booly on leadness the production of fice on statestorese, Cyclice labor netexes the risk of developing torgeterm farifity and hormoral issues, which are sometimes caused by the unitempted use of shorts for range must motion or years. Cyclica labor netexes must be an effective of the state of the state of the state of the state of the individual shorts are advected by the state of t

7. Use Resonable Doagnet, High does of secroids are not necessary to active significant mack provide operable if modern lepsing or predimente enhancement is desired. A doagne limit of 1600 mg per work on significant bis ad abird. In the case of homemore marker provides in a buddy mode heady bis local of one will provide a significant bis local bis loc

8. Avoid Assummates Inhibitors. A normans-sinking dangs context estogeric side effects by proversing the production of estogen in the body Wall as affective practice, they add deprive the body of a hermone that is important to cardiovascular health. In particular, estimations are used to a transmer that is important to a studio scalar health. In particular, estimation are used to the studies of the studie

9. Get Regular Blood Tests. Comprehensive blood essing including an examination of homoreas, cholstore, blood cell concentrations, and eroynes is the next useful toof of associating for negative health inpact of stored use. Changes in cholstored, for example, can help quarify for fact users that effect a particular drug regularity between the store of the example. The individual than hiss the exponentiary to better assocs large-stores the example. The individual than hiss the exponentiary to better assocs large-stores the example. The individual than hiss how the other example, and the example of the individual than the store of the example. The individual than the store of the example of the normal term of the norma

10. Use Proper Injection Procedures. Carelial anterito to correct igiction procedures can help cimuta core of the complications sociation with moreflical starolitus. Starolda are gene via deep intransaclar igictions. The most common site of application is the upper outer quality and the star of the aplicant start, although the days are also commoly the biseper, tixenge, or call matchels) for committee parameters are discussed and the start of the aplicant start of the applicant start of the applicant term of the applicant start of the applica

11. Sterilize. Though never advised, should the choice be made to use an injectable steroid of underground origin, an effort should be made to sterilize the solution before use. This will reduce the Reholo of illness or infection due to microorganism.

12. Vatch Your Diet, Anabolichardrogenis steroids can allow an individual significantly more latitude with their diet than neural. The caloric demund hypically increases due to the effects of these drugs on muscle mass and metabolism, allowing more calories to be consumed each day without adding fat muss. It is important not to kt this latitude affect your health in an engine way. Remember, the use of strends at physique- and performance. erhuncing does is expected to cause an unforomble diff in cholsterol level and other cardionsearch breadin murkors, fivoring a higher risk of cardionsearch breading discussion Stantharousky feeding your hody gender amounts of sistantied fits, cholsterol, and simple cardholgtates can much the impart of friends edings even worses. Dotts how in stantanted fits, desses risk. Note, however, that deta takos is not effective at contening the negative cardionsearch methods for storied use, but detary restrictions can reduce these rasks.

13. Abayes Consider Reward AON Risk. It can be easy to ignore the poterial health injunct of stard use when the positio benefits as so onpit and the mapine consequences so remote. At the end of the day, however, it is very important to remember furt the use of starokin is not smitching the marker and the starbay for the starbay f

Sterilizing Injectable AAS

Aubois stroid products much by lectured drug munificatures have an expectation of straining. This is assued by the rigit government regulation applied to these comparises. Understandy, there is no such expectation for counterleft and underground staroid products. Note are assembled under confinions that are not starch. O many coccasions, independent laboratory tasks have faund bacterial contamination in products of file's origin. There also appears to be a single hypingh products or discoss functions and other hyperconvectiing of the star of the start bacterial contamination in products of file's origin and the start products of the single start and the start of the start of the start of the start of the start is it. It is often arbitrarily to the single an ignortable started of unknown confir.

Heat Sterilization

The simplex method to clear a steroid solution of microcognisms is to beat sterifies i. The typical wey to according this is to place the steroid will ran one of 70 minutes (blace at 120°C/25°F). You should larse a needle sticking through the rabber top to cognize the pressure inside the vide of ram by brack. Note that a higher temporture and duration are typically used for complete dy heat sterization, but this is not practical for our proposes, the duration of vide components, and possibly ensure that duration of the duration Staff, the recommended 30 minutes at 120°C/25°F should kill most bacteria, and can be regarded as a vide hum medication states.

Some people prefer to use the top of the stove for hast sensitiation, and submerge their analysis started with as port obving water (OPC/2127). The will is keyn in the boiling water for approximately 20-30 minates. While this temperature is sufficient for killing many from of Dotestion and them minocognismes, is specurely not considered the best method of sterilization. A higher temperature should be used. It can also be difficult to control the temperature if the values on the bottom of the pot, which is in direct contrat with the human started source. Given these problems, it is preferred to attempt heat sterilization in an ours, an discussed above.

Note that tereholsene appears to break down at fairly low heating temperatures.⁷³⁴ If the degradation increases with temperature and duration. Some degradation microases with temperature and duration. Some degradation microaffect different conditions that those recommended above. This typically involves backing the viai in an oven for two hores at 80°CUT3°F. Again, an unattached needle should be placed in the nibbe stopper to capatile pressure.

Syringe Filtration

Andre way to nampt sterification of a stored solution is to use a fluet. A flue is a device that uses a merithmer with ity holes to prever particles form passing fluedy. It is smaller the holes are, the more effective fluer fluer, the rest of a depositely spring fluer is used. These are usually down in d-strincon and 22-micron intercopt or size. The 22-micron fluer is a device at a device share the string fluer model of a more share that the string fluer model of a more share that the string fluer sh



.22 micron Millipore syringe filter

The procedure for using a syring: fluer is findy simple. First, make sure you lave all of the necessary colparetary a syrings, two necessities (minimutin, a starket wyring) fluer, and a new scalad starke vial. If you are attempting to startize a standard 10 mi vial, if my be preferable to use a syring law ground to dur whe id all obtains. This will reduce the exposure of your organizing the pressure your should law et in the rubber visible drawing from the vial, here considiant to touch critical starkets of the coord necdor. In these stores of your organizing the pressure your should law et in the rubber visible drawing from the vial), here

Begin by draving the staroid solution into the springe. When you are done, nerrow the model and attachyour newly unserged optimizing there. Attach are wated to the end offset flott, and inject the solution into the new stark with The stark with and new needs to the end offset inportant, as using primal neuropearus would only equote the product to bacteria again. Use enough pressare to empty the syrings into the vial Re careful, as too much pressure can breach the hindings of the The process is show expectivily with a large syring or small 22microm flott. Provided great care was taken at to contaminate the staroid during the process, the scalings calcohan should be feed Theateria.

Also, these serification methods are applicable to ob-based steroids only. Heat serification (so onlind here) to take to cause the rapid exponsition of solver/simitraricolia agents in water-based steroids. The suspended (instead of dissolved) nature of the particles also related steroids. The suspended (instead of dissolved) nature of the particles also related to the steroid steroids. The suspended instead of the steroid steroids of the steroid steroids of the steroid steroids with the steroid steroids of the steroid steroids of the steroid steroids of the steroid steroids of the steroid steroids if they have been dispersed through legitimite supply characts.





Syringe filtration of steroid solution. Use of new (sterile) vial reduces chance of recontamination.

Limitations

It is inportant to emphasize that when applied in an uncontrolled environmer such as a honce or apartner, there is always the prosobility that your may inducterinfy contaminate year own product when attemptings ther it. Yous shudd always use storik globes and clean limit saffaces with abolical or black blober working on firme. Thank is also a very good microcognities. They do noting to address other potential health concerns such as the preserve of heavy much or united environizableau.

Many form of bacteria also produce endotoriss or programs, which would remain in solution een all off mittino role tast straining. Thus, you can note for our other above reaction even with no viable bacteria being left in the solution. An additional issue (though very mikely) is inter cation microcognism; for lending some versues and bacteria are either small enough or handy enough to remain which after these stellization techniques. The softly methods discredule tare are lightly effective, for an encoursely complex. For the higher level of addity, it is always recommended to limit your use to legitimate pharmacentical products.

374. FDA Environmental Assessment Report, Finaplix R (trenholone acetate). April 1987. Hoechst-Roussel Agri-Vet Co.

 Underground ANABOLICS. William Llewellyn, Ronny Tober. © 2010. Published by Molecular Nutrition, Jupiter, FL 33458.

Counterfeit Steroids

Even if a controlfed does contain the active strends in question, it may not be a chem and site product. Today, we take for granted the first fluor of map products are much owith strells and prave ingredients. We also give like thought to userptic processing techniques, which manufacture or drug products flee of communition by human haldbacturis, vinces runny of the human body roomed discussing anise taken are objected with the dates, which are object to order the discussion in the body. The frame and the dates are primated when a date is introduced durately num he body committee the strength and the dates and the strength and the strength and the body. The frame and the date is primated are conclusions of heses. The strength and the dates are primated when a strength and the process in any drug product, priced or otherwise. Constantions users primate the strength when how positionated and expression materian and the dates.

As anyone that studies the flicit analysis stored it rade knows, high demand and huge profits offer strong incentive for the manufacture of counterfield drugs. Over the years, this segment of the fliggl bisenses has grown exponentially. What was conce a problem https://soluted.to the United States can now be found in flerally every comer of the globe. Counterfielding is a advancement on the found in flerally every comer of the globe. Counterfield read here the advancement of the state of the state of the states and is advanced to can when the presentent nex nounce to manoe servore, or cores, stat a contrainty seen went me significant sike of organity subable are directed from signimum to underground sources. Given the name of drug products, however, the constraint's stored phenomenon is an opeicable inprotent headsh concerns for drugssing headybables. As the marber of these speciable inprotent headsh concerns for drugssing headybables. As the marber of these headsh conceptances including abscess, indiction, angelykosis, and toxicity caused by heavy media constraintion.

Prevalence Study

By all estimates, couterfet strovids are very common in al comers of the global strovid block mutet. Given the liker nature in which couterfet strevids products are inducli, however, it is difficult to determine the east prevalence of these drugs. Analysis reports of his enforcement seasons off an occasional samplest of the quilty of stovid sails is a particular region. One such study was conducted at the Center for Preventine Deping Research in Cologne, Carmany, and involute 70 different analood steroids and there all and the study of the study of the study of the store and the store and the groups of the study of the study of the store of the study of the study of the study of the store of the study of the study of the store and the study of the store and the study of the store static store of the static store of the store static store of the store static store of the store static store of the store

Overal, more than one-bried (34%) of the 50 anabole, steroids tested din reh have ingularies that mutheline their labels, and we calvery mode form filter mutaficatures. Of the filting products, nine were identified as copies of faxous phermecarical hunds, and woald be considered classics: contreficis. This member of 18% of the drug products in interactly of the an arsted classics. The remaining edget (16%) that filtied were underground steroid products, which are discosed separatively that book. There were as additional products on the fist thru passed testing that were mice by underground munificatures (Brith). There, more mutaficature 40% of the streoid products that were being solid by these classics in Germany, general. Gover the inject head on controls on works in the Unified States, contractive and underground products are expected to make up an even higher percentage of products Ready solid products.

Visual Inspection

The researchers in Codege Cernurg also made an important observation. Askie from horson undergoord product from his solar tab infelih Togon, SP Laka, and Hernnined Plannaceschich, they total I was not possible to accertain what product was not and what or ofference matterial recessivy to make an un-close greated carating and the photos were provided in the report to reference, it does underlike a problem that the steroid sing cormarily has been noticing for a long time, manyle controlf in multicating operations are becoming increasingly sophisticated. Now more than over I can be difficul operations and becoming increasingly sophisticated. Now more than over I can be difficul processors and becoming increasingly sophisticated. Now more than over I can be difficul processors and becoming the more direction.

The "Best" Products

Of the confiscted German products, those that were manufactured in Western Earcegsement to offic greater assurnce of lighting than those of other regions. Takand adsortermins a corronor source coursely for high-mate products net commonly manufactured in Westerministic net to this discuss, where comparing products (US: a classical to great most likely to be the subject of coursefring. Also, the study solution of the size course to instanticians on the highest courseling and the size of the

The "Worst" Products

Perhaps due to high recognition and demund, all of the Normal Helins mathebase documents products total during this analysis run were determined to be counselfs. These products were confiscant from each of the three dealers independently. In all cases, these stroud products contrained latestoreme riskated or mundhowed decaments. This is a contrain substitution with deviant matchbase products, as low dooss of stotsterence can provide a similar level of multicle effects are matchbase for some uses, with a similar low inclusione single contrained and the similar base strengtheness of the stotsterence can be applied as the context Norma Heliss Dave, therefore, normalises are stoted to contemply lapits rise on the Lancopen and iteratively and the context base to the stote to contraine any product cookey for the regardle scotter former shown in the low classifies.

Other Bodybuilding "Ancillary" Drugs

A total of 20 non-steroid drags were also tested. Al products that would be defined as common actility drags including attraviation tanke (Noholeck, chorablene crites (Chorabi, thyoid) homore, califierie, and yolinrihe led harned on to be kgirinnet. This underfines the lower risk in these archived range in the odds due to the lower fixer in these refixer the constructions to deplicate these cheap and easy to access pharmacentaish. The odd range attraver is the second range of the chartersh as well.

Steroid Analysis Results

Anadrol (oxymetholone): 1. Oxytone 50 mg (SB Labs, Thailand) Result: PASS

2. Oxytone 50 mg (SB Labs, Thailand) Result: PASS

3. Oxytone 50 mg (SB Labs, Thailand) Result: PASS

Deca (nandrolone decanoate);

1. Norma Hellas (100 mg/mL) Result/FAIL (testosterone)

2. Norma Hellas (100 mg/mL) Result: FAIL (testosterone)

3.Norma Hellas (100 mg/mL) Result: FAIL (testosterone)

4. Norma Hellas (100 mg/mL) Result: FAIL (testosterone)

5. Decabol 250 (British Dragon, Underground) Result/FAIL (testosterone)

Dianabol (methandrostenolone): 1. Anabol 5 mg (British Dispensary, Thailand) Result: FAIL (methyltestosterone)

2. Anabol 5 mg (British Dispensary, Thailand) Result: PASS

3. Anabol 5 mg (British Dispensary, Thailand)

Result: PASS

4. Danabol DS 10 mg (March, Thailand) Result: PASS

5. Danabol DS 10 mg (March, Thailand) Result: PASS

6. Naposim 5 mg (Terapia, Romania) Result: FAIL (methyltestosterone)

Equipoise (boldenone undecylenate); 1. Boldabol 200 (British Dragon, Underground) Result: PASS

Halotestin (fluoxymesterone); 1. Fluoxymesterone (IP, Underground) Result: PASS

Primobolan (methenolone enanthate); 1. Primobol 100 (British Dragon, Underground) Result: FAIL (nandrolone, testosterone)

Proviron (mesterolone): 1.Proviron 25 mg Result: PASS

Sustanon 250 (testosterone mix): 1. Sustanon 250 (Karachi, Pakistan) Result: PASS

2. Sustanon 250 (Nile, Egypt) Result: FAIL (different testosterones)

3. Sustanon 250 (Nile, Egypt) Result: FAIL (different testosterones)

4.Sustanon 250 (Karachi, Pakistan) Result: PASS

5. Sustanon 250 (Karachi, Pakistan) Result: PASS

6. Sustanon 250 (Karachi, Pakistan) Result: PASS

7. Sustanon 250 (Karachi, Pakistan) Result: PASS

8. Sustanon 250 (Karachi, Pakistan) Result: PASS

9. Sustanon 250 (Karachi, Pakistan) Result: PASS

Testosterone Cypionate: 1. Testex Prolongatum 125 (Q Pharma, Spain) Result: PASS

2. Testabol 200 (British Dragon, Underground)

Testosterone Enanthate; 1. Testofort 250 mg/mL (Pliva, Pakistan) Result: PASS

2. Testosterone Depot 250 (Eifelfango, Germany) Result: PASS

3. Testosterone Depot 250 (Eifelfango, Germany) Result: PASS

4. Testoviron Depot 250 (Medipharm, Pakistan) Result: PASS

5. Testoviron Depot 250 (Medipharm, Pakistan) Result: PASS

6. Cidoteston 250 (CID, Egypt) Result: FAIL (includes T. cypionate)

7. Cidoteston 250 (CID, Egypt) Result: PASS

Testosterone Propionate: 1. Testovis 100 mg/mL (SIT, Italy) Result: PASS

2. Testovis 100 mg/mL (SIT, Italy) Result: PASS

3. Testovis 100 mg/mL (SIT, Italy) Result: PASS

4. Testovis 100 mg/mL (SIT, Italy) Result: PASS

5. Testovis 100 mg/mL (SIT, Italy) Result: PASS

6. Testabol (British Dragon, Underground) Result: FAIL (different testosterones)

Trenbolone (various esters): 1. Trenabol 75 (British Dragon, Underground) Result: FAIL (boldenone, testosterone)

2. Trenabol 100 (British Dragon, Underground) Result: FAIL (boldenone, testosterone)

3. Tri-Trenabol 150 (British Dragon, Underground) Result: FAIL (trenbolone, testosterone)

4. Trenabol 200 (British Dragon, Underground) Result: FAIL (trenbolone, testosterone)

Winstrol (stanozolol): 1. Winstrol Depot 50 mg/mL (Zambon, Spain) Result: PASS

2 Winstrol Denot 50 me/mL (Zambon Smain)

Result: PASS

3. Winstrol Depot 50 mg/mL (Zambon, Spain) Result: PASS

4. Stanabol 50 (British Dragon, Underground) Result: PASS

Counterfeit Steroid Identification

This section pertains to methods for differentiating between legitime planmesotical products and digitime coopies (conterfield). Before we begin, in each orming to not conterfieling analods steards it as very learning between the section of the section of the periadic of their products. Furthermore, then we nev memp large "contentiations" contentiating operations, with the capacity to manufacture all product forms including angeles. Journ primer plik, and publicity during the set of the large content of the periadic plant and the product forms including angeles, long mirring this, and publicity during that strates, form in high level of supplications, strates) parelined on the black match rate to be impacted with grant case. Methods and the set of t

Step #1: Eliminate the Obvious

When conterfs stronks first appared decades ago, they user often very easy to spot. The mandfacters operated on a small scale, and mode small-scale missions. For earning, the printing might be skeptige of the containers that and firms. They might how lacked the box hybrid strong the strong strong the strong strong strong strong strong strong to hybrid strong strong strong strong strong strong strong strong strong hybrid strong stron

1. Stopp: Printing. Drug munificating is not a small scale enderove. Sinoble parameterical compression soften by hold long trade, and make products that are typically very professional in appearance. You should not expect to see things late entry like, hypersection, and the state of the

2. Cheap Predoaging, Virnaly al legitimuts strond products cours in boxes. Inside the box you should find a dark information shots: Conce coursefections will also these stops on triple. Real arraphets, vish, and labels are scenttrings samging thoses, but it scenteers the labels that the strong strong the strong strong the strong stro





The ampule is a laboratory blank, meant to be sealed by hand over a flame. It is larger and more unusual in appearance than most traditional ampules.

3. Multi-dose Containers, In the United States, we are used to our specialite medicatives coming in multi-dose with these lives an other poor lot medicality measures and our pills loss in bottles. Most other coantres, however, do not allow links poor of the special structures of the special structure of the special s



The above is a crude copy of an American testosterone product, which uses the same label on the box and vial. A counterfeit as simple as this is rare to find today.



Another example of an obvious counterfeit. This box is crude in design and uses a brand name that has been off the market since the 1980s.

The photographs below show what it looks like when the lot number and expiration date are added after the initial box/label printing, as well as counterfeit products without this feature. The characters on a real planmeetical product should stand out from the rest of the printing which will consist of thy dots blenching dates to rest as a data may constant of for more information on the init). When the lace/printion information is added with a mechanical starm, the its will be much more sold under magnitisation (note that it muy appear blockly under deep magnitustion). Depending on the expansion it, it muy also have left applicant loost years and be shown halforgue contrains over the information. What the data exactly to hold at the characters closely. Constructions will be placed by the much neutrino the data was added by the shown of the shown of the shown in the information. What the data exactly its hold at the characters closely. Constructions will be to make the information hold, the its vas added by muchics or comparts, core moding it was printed. If yous set that first dots make up the characters under 200X magnitudinos, it is not lagitimet starp or computer printing.







Example #1. A real box of Proviron. Under magnification we can see that the lot number and expiration date were stamped on mechanically.

Step #2: Examine Lot Number/Expiration

A more formal analysis should always begin with the 1st number/capitation date. Promencetical compressions here the bosons and hash number/capitation dates have not yet been of date priming facility. They are not sensitively, the number of sensitivity of the sensitity of the sensitivity of the sensitivity of the sensitivity of the





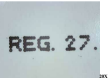
Example #2. Another example of mechanical stamping of the lot number and expiration date.





Example #3. Real testosterone cypionate from Watson (U.S.) The above lot/expiration date were added by computer printer. Under magnification we see the large dots are solid ink.









Example #4. Another real product (Proviron) with information applied post-printing with a computer printer.



Example #5. (Counterfeit). At first glance the slight run on the ink appears to be the result of mechanical stamping. Under magnification, however, we see this is simulated.





Example #6. (Counterfeit). An example of simulated computer lettering. You can see under magnification that the information was actually made by normal process printing, along with the rest of the box.





Example #7. (Counterfeit). Again, the date was added with the rest of the printing. This manufacturer simply made the text bold in an effort to hide the printed nature of the information.

Step #3: Look at the Ink

One of the first things that the U.S. Scoret Service does when it takes in a contrefet bank note is to examine the nisks used by the concertificarts. They have made doing ago that you can tell a grant datalabent a bill this way. You can tell how and when it was printed, somerimes even when. Now we are not agoing to analyze the first on storial balows in the list so the Scoret differentiate real from fide in some cases. Once again, this will be done with a hundhed micrococope, referentiated at 200N magnification.

Fig. Let's go over some general information so you know what we are looking at. For this ecurination we intend to find our how the priming was pheced on the box or held. Colver, printing is suadly accomplished frequent how methods. The first is called "process color," and the second "spec tock". One method may bue used to print the product, or both methods may be combined on the same packaging. Each method offers is own set of advantages. When we know how word where these methods are typically applied, use can given some single it in the throughts of the designer. None importantly, we might be able to tell if the methods are approprint for the product in question.

The process older method is most often used to reproduce photographs, or to prirt multiple codes (these or more) on one poete. It is the more forching and used, does method to codering. Process coder can give the appearance of handecds, thousands, even millions of codering a three codes and the standard structure of the standard structure of the structure of the structure of the structure of the structure is the structure will be structure of the structure of the structure of the structure of the same structure of the structure of the structure of the structure of the same structure of the same structure of the same structure of the structure o

The spot color method is used when a very specific color fit is nequefic. Instead of using a CONY bend to repeate a contrain shead of green, for cample, an fit is miced in the provide the coast green color that is needed. If you look at spot color used 200X magnitudin, you will not see CONY color data. Small oth are sail used, but they are at the same color (they are placed closer or finther sparse for shafing). Note that it can be very hand to neach these is in staces with solid The major shadning. Note that it can be very built to inscribe these is in staces with solid The major shadning to spot color is applied in the printed packaging overy time. You will make full by companies using post color, expectively with projectar operator the termine (house, nathermatic). The main fing we are looping to see under magnitudino is the use of score appendence the lack of spot to obse not necessary mean that a product is counterfits, you should expect to see it on most real planmacentical packaging samples. If the package has only nor of three cohers, they will produly use good cole exclusively. If the package has no hyper on three cohers, they will produly use good cole exclusively. The package has more than elements. Spot cohor usually signifies some attention to detail, which sometimes the counterficient miss.



20X



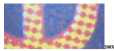


Example #1. This box of Depo-Testosterone was printed entirely with spot color. Large companies like Pfizer tend to prefer this method of printing.Note the unified dots under 200X magnification, indicating spot color.









Example #2. The above underground steroid product uses process color. The small CMYK dots that blend the color can be seen well under 200X magnification.

Step #4: Magnify Logos/Small Elements

Each countrifier is faced with the dilemm of diplicating an original strenis's pockaging on their onn expinent. The most common ways to 6 m is is one optim thatmach how or high with with a deduce scatter. The picture will be cleared up in a compute, and ultimately contend it on piles for printing. In this pocks, the dup gooding is duplicated in which amounts to a complicated from of photocopying. Though a scentagy simple process, drug pockaging much in the immers can be suprisingly occurate in appeares. Some and so with a time data for the picked of the picked on the duplicate whole a time datation of the content of the duplicate whole a file duplicate for the duplicate whole a time datation of the content of the duplicate whole a file duplicate whole a time datation of the duplicate whole a file duplicate file.

The type of counterfeating does have is initiations though. Most notable is that very file detail is usually bot in the reproduced image. Much of this is due to the scanned in image breig poinded, and thas not annumble to scaling and applie monipation in the issume ayone the organit applies would be. When centration durken magnification, the lines may appear chapty. Make sure you look at the small applie elements of the staroid package, not the taxet. Counterfiels: so anally review the tax to its is dury. They count repress a comparison logadat is produbly a counterfail.







Example #1. Real Cidoteston from CID Egypt.







Example #2 (Counterfeit) Cidoteston. Under magnification we notice that the scan of the initial box was insufficient to reproduce the fine detail. The inner ribbon around the logo is especially proken, and the letters "CID" poorly defined.

أمبولة واحدة 1 ملليلتر وستانون ۲۵۰٫ للحقن فى العضل لا يستخدر الا بأمر الطبيب بحتوى كل ١ ملليلتر على : ۳۰ مجم تستوسترون برویبونات ٩٠ محم تستوسترون فنبل – بروبيونات • ٣ مجم تستوسترون أبزوكابروات ۱۰۰ مجم تستوسترون دیکانوات في محلول زيتي Terrent Normal

20X





Example #3. Real Sustanon from Egypt. Notice the fine detail on the snake.

أمبولة واحدة 1 ملليلة سوستانون «۲۵۰ للحيقن في العيضل لا يستخدم إلا بأمر الطبيب یحتوی کل۱ مللیلۃ علی : ٣٠ مجم تستوسترون بروبيونات ۲۰ مجم تستوسترون فنیل ـ بروبیونات ۲۰ مجم تستوسترون ایزوگابروات ۱۰۰ مجم تستوسخ ون دیکانوات في محلول زيتي ستع في مصر FILLER





1003

200.1

Example #4 (Counterfeit) Sustanon. Notice how the finer details are lost on the duplicate.

1.000



Example #5. Real Deca-Pronabol from P/B/L, India. Notice how the logo remains smooth under magnification.







Example #6 (Counterfeit) Deca-Pronabol from P/B/L. Notice how the logo is choppy under magnification, likely due to the image being scanned from an original box.

Passing Inspection

It is inportant to once again emphasise that storoid controlfing is high basiess. An efficient operation can ent may million of dolume year for is owners, which so much moves at stake, opsishciation is at an all time high. In some cases, the countrifiers will have very similar high-level qualitotis to the pharmacentical companies they are corporg, Tins means or only the ability to reproduce complexied packaging such as break-open giase amples and push-hough habit ways, but also fits chains. It is markering which the above table ID mutations, habit graphic security staticus, and even habit, learning, but above constructions may also approximate the transfer of the static security of the static constraints may also approximate the constraints of the static security of the

Country Specifics

In most courties, a pharmacorical company is required to meet a specific set of regulatines pertaining to the physical packaging of a pharmacerical product. In some cases these regulations can be used to help evaluate the legitimecy of black murket steroids, as all counterfeites may not have the resources of redurbagit to implement the required features. Here, we discuss a number of attributes to examine, which should hold true for all of the drugs produced in the specified country.

United States:

Fig. 4 is very important to stress the fact that stressla are a controlled substance in the United States. Current controls are very efficient at keeping American products of of the black multicle, It is much easier for the flict dealers to import or munificative their composites than it is to gat an volution of eigening American public dealers. The products that is to gat an volution of eigening American public dealers at they are in attemption poster than its to avoid all American items unless, our expersionally more them backs to a phenemes.





The FDA provides us with a couple of strict requiements, which many counterfeiters overlook. The most predominary is that all algointude American drugs cannot carry a label that will easily be removed from the viablottie. It must be so startated with gale that you can only remove it in small pixes. This is done to protect the public from the possibility of drug mikibeling. With many U.S. counterfeits, the label can be peeled off the bottle quickly, in one or a few large pixels.

You should also moisten your flumb and rub the expiration date on the box and label. Quite offen the ink on the counterfeit will smear and rub off easily. The stamping on a real U.S. pharmaceutical may streak slightly, but should remain intact and legble. Again, this is a requirement to protect consumes.

Additionally, being a Scholak III controlled substance, all commercially available human and veterinary anabolismotogenic strongs are required to hear the tag "CIII" (see sample pisture). The only exceptions would be cathe implart peleks, which are technically not controlled substances, or drugs from compounding planmaries, which do not have to adhere to the same production gaidelines. A small namber of lazy conterfeiters contract to diplicate strongs that are unstitutive before 1991, when this given and present to diplicate strongs that the same fragment of the same strong str

The FDA requires that all tables and capsules are identifiable through unigne marking in case they are remored from the packaging. The mandingter many can allowitoriato, is usably found on each plit along with a specific code for the product. Some stored users have found the packaging with a specific code for the product. Some stored users have found the packaging and the packaging and the packaging and the disposal, and should be able to bed you the dram and doogs the packaging at their disposal, and should be considered all shour pand. Along have and the displays, if allowing and the displays, if allowing and doogs based on them. The offices are usably very responsive if you explain it is not an emergency call. If your plit is not found in the database, it to able to evolve the disp product.

Italy:

All drugs produced in haly will bear the pictured drug identification stcker. The sicker is beilt is white, with read block print: The schere rest on a humined straffice, so that it can be peeled off and affited to paperwork. You should never parchase an Italian drug if this sicker is not preset: Drugs from hily will also use abbreviations loss Persp. Scad. and Del for the counterpart of 10t #, manifacture, and expiration dates. English writing here would indicate counterfeiting.



Greece:

Greece also has a drug ID sticker that must be present on all drugs available for sak. The sticker iself ress on a laminuted stiffice, so that it can be peeled off of the box and affield to paperwork when a prescription is filed. Most importantly, the sticker will show a hidden musk when placed under UV light. Some counterfeiters have copied these stickers with excellent accuracy, right down to the laminuted suffice. Copies of the sticker bearing a light IT. We may and how the barries of the sticker bearing a sticket. The summary down to the laminuted suffice. Copies of the sticker bearing a light IT. We may and the sticker the sticker bearing as the sticker the sticker bearing as the store how how the store of the sticker bearing as the sticker bearing as the store how how how the store of the sticker bearing as the sticker bearing as the store how how how the store of the sticker bearing as the sticker bearing as the store of the sticker bearing as the store of the sticker bearing as th mutern OV watermark, nowever, nave not been located. Do not purchase any oreek orug without the proper sticker attached.



Spain:

Sparsh drugs do not bear a sticker, but instead have an area located on the box that contains a bar code and some drug information. This area will sometimes have indentations in the cataboand, so as to be removable if you tear the sarface. At other times, the barcode is simply printed on the box. Spanish drugs also use the abbreviations Lote and Cad for lot number and expiration, respectively, take. Many drug boxes also care Brailie lettering.



PVP M.R. 690 Pts PVP IVA4 718 Pts

Printed Only





Removable Barcode



Braille Lettering

Portugal:

Dag boxes from Portugal contain a rectangular area which displays the box code and pricing information. This is sometimes found as a sticker, but most commenly it is printed, not stamped, onto the surface. In many cases, the area is indented, so that it can be removed from the box. Drugs from Portugal will also use the abbreviations Lote: and Val. Ate: for lot number and explained not astampings.



Old one without



New area with price in Euros

France:

Drugs from France will bear a rectangular sticker somewhere on the surface of the box. The text and format is often slightly different item to item. Also, packaging always contains an area with a green and red box. In the sample below, it is in the lower left side of the box.



Underground Steroids

An underground serveit is an ambolicantiogenic serveit product that was much by an lingel (indiground) biostraty. These draws are specifically mundicatured for side to atfates and hodybalders on the black mulcit, and are not available through legimine charaches and as platemics and and angiostrators. These composites are unicreased, unregatated, and operate in a completely clausifiering mumer. At one time the team underground stored uses considered sporymous with contredict stored. But observation and a constraint stores with contredict stored by the store time time and a constraint stored is that the later is a coupt of a legiment and an constraint stored is that the later is a coupt of a legiment and an constraint stored is that the later is a coupt of a legiment and an another the store and an another stored and the stored multicature, on the other hand, use distort band names that are out to be confident with registered date companies. They dont to the bald recognition in the marketpice for their produces, and commody use real storeid ingredient is substantial doesans.

Tapletar government enforcement or discoid losss and kignitude distribution charaches in the discoil an explosition of underground assistent manafesturing operations or othe post docade, andockinadoganis atsevids linguity soil in the United States. The manufacturing process typicalli photoses bearoing of root stored in their linked States. The manufacturing process typicalli photoses bearoing of root stored in their linked States. The manufacturing process typicalli photoses bearoing of root stored in the United States. The manufacturing process typicalli photoses bearoing of root stored in the stored store of the photoses and the manufacturing and the stored store of the stored store of the photoses of the stored the of the stored the stored store number though, and account for a small portion of the total number of underground laboratories.

Drug Purity

Dang mundhenring safty is a cortal locar in Western medicine. Planmeetriche are strended to test ifl printest, nor cause additional hurn by being imporpely dood or containing heatrin, havy meth, or other forms of contamination. Products made for human consumption are only much after a yourness reported by genement-based compariso, contain materials that come form other knowed appiers, which also adhere to article planmeetrical-galaxies and and a strength and the strength and the strength planmeetrical-galaxies in metachands scrattined "channeous" facilities designed to prevent and contamination from and papersonal. Each notation of previse designed the products into contain an other and appicenses. Each notation of prevent and and any constraintion from and personal. Each notation of prevent and and any constraintion from and personal. Each notation of prevent and the strength and the strengthand and the strength and

The above description is static contact to the underground sterial manufacturing basisses. In priority of underground sterial products well, likewise, not be assembled in a sterile environment, or with the scal of expansion planmentschild grad methods and compares subscriptions of the scale of expansion planmentschild grad methods and compares of "local grader" more materials and mennally operated viablords filling and scaling tools. The opportunities for communition in fits types of process are grader. By Western medical standards, most underground steroids are, of course, nor flor far human constraintion. They may full attraction to cheaper process, higher desce, gaptate selection, or course subsidiary between medical steroids, may not be higher desce, gaptate selection, or course subsidiary between medical steroids and the selection of the source store and the selection of the selec

ANABOLICS Underground Market Analysis

In an effect to help commerse assess the quality and poterali health risks of traderground stored products. AMORIUS: matericules a detailed joint eng manyles mojests in Agent 2007. This project ecummed the quality of assession function traderground healths, and conversional in a store of the store of the store of the store of the biometry of the store of the store of the store of the store of the biometry of the store of the store of the store of the store of the biometry of the store of the store of the store of the store of the theory of the store store of the store manufactures, mil-level operations, and over produces large store and head their learns manufactures, mil-level operations, and over produces large store of the store store storestore and the stores of the store.

There were for specific areas of resting for the 2007 metcle analysis project. The fort test was to look for the presence of toxic heavy hear metcles and a heat in mercury and metcles. These methe all poses specific threats to health if they accumulate in the body. Those methel considered iterat, such as iteration, were not iterated. How, we commissioned they for such and the strength of the strength strength strength strength strength strength for unknown strength contained and strength strength strength strength strength strength strength they communicate the strength strength strength strength strength strength strength strength they communicate the strength strengt

The specific results for each of the four testing sets are presented in the tables below. Overall, the products examined in this study reflected poorly on the quality of the underground steroid market. To begin with, more than 20% of the products (1 in 5) contained heavy metal constraintion. While pre-market testing would have caught this, if such products were ever fourd on pharmacy ababes in the United States i would ringer an immedian trainformative recall toxet, an examination of busics drug doing showed many deviations. Approximately 32% of the products were actually significantly overdoaded. While this was likely does in editors to produce a stronger user response and logical cattered base, this is an unacceptable lattic which misses many potential safety issues. In the first set of estimation, the stronger stronger stronger stronger stronger stronger does also more than the stronger strong

Drug Analysis Results

Test #1: Heavy Metals Contamination

| Sample | Contamination | Result |
|----------------------------|------------------------|--------|
| 1. methandrostenolone | None Detected (<0.002) | PASS |
| 2. testosterone enanthate | None Detected (<0.002) | PASS |
| 3. testosterone enanthate | None Detected (<0.002) | PASS |
| 4. testosterone propionate | None Detected (<0.002) | PASS |
| 5. boldenone undecylenate | Metals Found (>0.002) | FAIL |
| 6. testosterone cypionate | None Detected (<0.002) | PASS |
| 7. boldenone undecylenate | Metals Found (<0.002) | FAIL |
| 8. trenbolone hexahydro. | None Detected (<0.002) | PASS |
| 9. testosterone cypionate | None Detected (<0.002) | PASS |
| 10. methenolone enanthate | Metals Found (>0.002) | FAIL |
| 11.testosterone cypionate | None Detected (<0.002) | PASS |
| 12. nandrolone decanoate | None Detected (<0.002) | PASS |
| 13. methenolone enanthate | None Detected (<0.002) | PASS |
| 14. trenbolone enanthate | None Detected (<0.002) | PASS |

Failure Rate: 21%

Test #2: Dosage vs. Label Claim (mg/mL)

| Sample | Labeled Dose | Actual Dose | % of Claim | Pass/Fail |
|----------------------------|--------------|-------------|------------|-----------|
| 1. methandrostenolone | 25 mg | 115 mg | 459% | FAIL |
| 2. testosterone enanthate | 250 mg | 440 mg | 176% | FAIL |
| 3. testosterone enanthate | 250 mg | 408 mg | 163% | FAIL |
| 4. testosterone propionate | 75 mg | 127 mg | 169% | FAIL |
| 5. boldenone undecylenate | 200 mg | 240 mg | 120% | PASS |
| 6. testosterone cypionate | 200 mg | 204 mg | 102% | PASS |
| 7. boldenone undecylenate | 200 mg | 178 mg | 89% | PASS |
| 8. trenbolone hexahydro. | 76 mg | 190 mg | 24996 | FAIL |
| 9. testosterone cypionate | 200 mg | 177 mg | 88% | PASS |
| 10. methenolone enanthate | 100 mg | 54 mg | 54% | FAIL |
| 11. testosterone cypionate | 250 mg | 171 mg | 6996 | FAIL |
| 12 nandrolono doranosto | 250 mg | 228 mm | 0104 | DACC |

| 12. Italiuroione decanoate | 200 119 | 220 1119 | 2170 | FM33 |
|----------------------------|---------|----------|------|------|
| 13. methenolone enanthate | 100 mg | 78 mg | 7896 | FAIL |
| 14. trenbolone enanthate | 100 mg | 0 mg | 096 | FAIL |

Failure Rate: 64% (+/- >20% of Label Claim)

Test #3: Steroidal Materials Purity

| Sample | Contamination | Result |
|----------------------------|--------------------------|--------|
| 1. methandrostenolone | None Detected | PASS |
| 2. testosterone enanthate | None Detected | PASS |
| 3. testosterone enanthate | Unknown Peak Detected | FAIL |
| 4. testosterone propionate | None Detected | PASS |
| 5. boldenone undecylenat | e None Detected | PASS |
| 6. testosterone cypionate | Unknown Peak Detected | FAIL |
| 7. boldenone undecylenat | e None Detected | PASS |
| 8. trenbolone hexahydro. | Unknown Peak Detected | FAIL |
| 9. testosterone cypionate | Unknown Peak Detected | FAIL |
| 10. methenolone enantha | te Unknown Peak Detected | FAIL |
| 11. testosterone cypionate | None Detected | PASS |
| 12. nandrolone decanoate | Unknown Peak Detected | FAIL |
| 13. methenolone enantha | te Unknown Peak Detected | FAIL |
| 14. trenbolone enanthate | Unknown Peak Detected | FAIL |
| | | |

Failure Rate: 57%

Test #4: Oil Purity

| Sample | Contamination | Result |
|----------------------------|-------------------------|--------|
| 1. methandrostenolone | 2,4-Decadienal Detected | FAIL |
| 2. testosterone enanthate | None Detected | PASS |
| 3. testosterone enanthate | None Detected | PASS |
| 4. testosterone propionate | None Detected | PASS |
| 5. boldenone undecylenate | None Detected | PASS |
| 6. testosterone cypionate | None Detected | PASS |
| 7. boldenone undecylenate | None Detected | PASS |
| 8. trenbolone hexahydro. | None Detected | PASS |
| 9. testosterone cypionate | None Detected | PASS |
| 10. methenolone enanthate | None Detected | PASS |
| 11. testosterone cypionate | None Detected | PASS |
| 12. nandrolone decanoate | None Detected | PASS |
| 13. methenolone enanthate | 2,4-Decadienal Detected | FAIL |
| 14. trenbolone enanthate | None Detected | PASS |

Conclusions

The scope of testing for this project was fishly limited, and field well short of the detailed mass/net required to walking a real processively non-dimensional sector products to fill sesting. These dangs locating for a samong majority of the underground stored products to full sesting. These dangs loops are strateging of the store of the store of the store of the store of the loop of the store of the loop of the store of the loop of the store of the store

Legiment plurmocedical products are munificated under strict conditions for a reason. It is very difficat to minimum an exceptible being of propt without them. From F pare LSPNF grade minimia are being used, I can be very easy for a minoscopic biological plurity of advective strict and the strict of the strict and the strict of the strict product strict and the strict of the strict and the strict of the strict product strict and the strict of the strict and most regardly and advected. Undergrade minimizations the strict plurity and most strict and advected the strict and the strict and most regardly and the strict area likely for caceptions, insolving only the largest and most regardly undergrade products approach to be oblighted the strict and most regardly undergrade products approach to be oblighted the strict and most regardly under the strict parket strict and the oblighted contaminuts, constances muscle shares that the strict and park to minimize the association of the strict and most regardly and the strict park strict and the strict and strict and a strict and

Designer Steroids

There is a fulf law in the stored detection methods used by the various sports agreeds. This is in order to transformed for analysis storids, you read to have earch what you are looking for You carily just took for "storeds" in the urne, but are forced to to far the early specific compound individually. To make finally use cont more complication, you need to know what they are gaing to bodk like by the time they agree pairs in the transformed for analysis of the comparison of the models was also been been by the transformed for the comparison of the transformed for the comparison of the comparison

Vous ese, seven handsel, froa a donsand or more, different stronis were sprinsized and inceptatel in various bhorntonis around the world dang the hopkyo of storied research. In most cases, their analoki and androgenic potencies were measured, with the same methods that hove beau of and of the popular stories low leaves toolky. Oth a mittane faction of these research compounds altitudely hearmer commercially soublike dang any area of darg second-bandy as hear too world beaution of similar dargs to coast in the same market. But the ourly research is sill or there, and remains a very sukable source of fortuniton for the clover chronis of foldsy.

Some of these old research steroids of the '50s and '60s still exist today, due to the diligence of underground chemists and researchers. We refer to these drugs collectively as 'Designer Steroids'', and they are here only for the purpose of defeating a drug screen. A true designer steroid is structurally unique next to the known anabolic/antrogenic steroids, sharing no common metholicies, so as to be undetectable to even the most through steard lost. The through of tracking you methololics for all possible storoids corporals, to elimitate the designer storoids issue, scorns like an impossible task to usy the last. Even if somehow first oil search were to be chanted, and metholicis identified for all lorown storoids, there are still nearly limites other ways to also textosterone, numbrohe, or displortextosterore to make using new storids. The doigner stored phenome could obviously present an overwhelmig problem to the sports organization gioun present drug testing methods. The affects can easily some or troo seps abactual and hoody on the sidelines is the wise:

At this point in time, the fact that dosigner atomics exist is no secret to the sports agarcies. It became partially obvious to the ICC (International Obpits) Commission in March of 2020, when the UCLA Obpits: Analysis at the test is the 10-866 s that of the test is the sport of the observation of the sport of the test is the test of the test sensitive from a Synthys. This was the second time the failed back in the 10-866 s that of the sport and the observation of the sport sports of the sport and the sport of the sport

Cafin use able to obtain a sample of pure netbodyne from the drug company Word, and must have been guind, and bely the first thrat methods of this stored in blaves identified a largest starting 2^{-33} . The procodure for norbodynes detection has now been made available to all string agrees; and unformathy it is now useds for competition. Its value as a designer stored in Review vanished overright. Perhaps I was able due to use a stored that catual mode at all flavours to the point of default links of a number of model that catual frames at all the ours to the point of model mission in that U.s. store starting all that can made of mission and the starting of the store and the store and the store of the store

The norbolethone story quietly fell from the public conscience not long after it broke. The number of athletes that ultimately tested positive for the drug was minimal, so it really never evolved into the big scandal that was initially expected. The USADA thrives on negative media attention to steroids, because it leads to more government funding, so no doubt this lack of public outrage was a disappointment. I would suspect many involved were hoping for the global story on par with what happened when Ben Johnson was stripped of his gold medal during the 1988 summer Olympics. This would be of little matter by January 2004, however, because a much bigger doping scandal was about to hit. It involved the use of the designer steroid tetrahydrogestrinone, and this time would snare some of the biggest figures in amateur and professional sports. Not just Olympic competitors, but professional football and baseball players were being listed as potential violators. Many household names were being thrown around, including Jason Giambi, Barry Bonds, and Gary Scheffield. Over 20 athletes ultimately tested positive for THG, or were specifically named for using it in the evidence. The investigation continues today, so this number may rise. Don Catlin was once again the scientist who helped identify this compound in the first place, as well as a method of its detection in urine. This time around, however, he had a lot more help then he did with norbolethone. THG was actually handed over to the IOC testing laboratory in a syringe, by an anonymous coach who did not approve of its use. With the help of an inside informant, USADA got their Ben Johnson story, and then some. THG was at the center of the biggest organized doping scandal in the history of competitive sports, and would come to spark a more vigorous government fight against steroid use than we had yet seen. The steroid-using community is only now beginning to feel the backlash.

I include hese stories not because they illustrate victories for the IOC. Quite the contrary, I believe they underline the major finiting is uncernst storied its story methods. These tow includes its logically do not represent the only two designer storoids ever used in competitive systems. For one, we servely carnet opercel 1 of 10% success note for the IOC when we know that TRA is use west completely method. The story of the IBC weeks we know applying abort first storied and a sample saw inhaled out to the storieg for highly which is the same facility that and unwritingly been passing uries samples containing the same storied just days.

before. Were it not be the mask source, 11% words probably still be in use today. The modelshore and TMA stories spin in the cord of hose on the sidents, who in sist and today used as a stored with the side of the side stored with the side stored with the books, or in the grant base, of them grant base stored with the side sparse for the next disagare compound to be described by the cord with an effect of the side stored with the side sparse for the next disagare compound to the side stored with grant with grant who they have a be synthesized once it is a side sparse for the next disagare compound to be sparsed as the side of the side stored with the side stored with the product of the side stored with the

378. Schanzer W, Donike M. Anal Chim. Acta. 1993; 275: 23

Anabolic Steroids and the Law

Unied States hus prohibits the procession of anabotic staroids without a legal medical prescription, imparting servere panalities (neutraling fine and/or imparted theores to violate these laws. Under inflatence of ULS, sportmart of ficials, Wolf Anti-Doprier, Agnery (WADA) murthers, and public critician fidowing namerous dopries scandals a growing number of coarties are following the ULS by adopting their own has signate the prossession of natiods staroids and othen sport dopries (large, lar many cases deall autrices unificial) has that concern the personal use of anabolic steroids and other related datas.

United States

Anabols stretch have been closefield as corrected an abstraces in the United Status size 1991, with passage of the Anabols Steward Corrend Let of 1990 (which. Ls. No. 110-647), Sac. 1902, 100 Sat. 4851, 1990). This law rankes it a criminal different to self, distribute, populatios for prosteen without a high participant processing and the propanditiss for prosteen without a high participant processing and participant propanditiss for prosteen without a high participant procession without a high participant propanditiss for prosteen without a high participant procession without a high requirement fund as a prior day conversion. Note that this kaw was anreaded in 2005 Subwirg passage of the Anabolic Stread's Control Act of 2004. The new with add 26 new stored compounds the high or control of substatus and also removed the lagal requirement that a compound be protone anabolic in moduloc to arrenview of the lagal requirement fund a compound be protone anabolic in conductive of the protone and with 26 new stored of the high or participant and with 2005.

State vs. Federal

Crimina lines against the possession of anabolic securities exist at both the foldent and state level in the U.S. Depending on the circumstate, an initializating probability the being which as stored related crime the by either the foldent agovernment or the state government where the crime tool, physical crimes the lines are the state based on the state of the state of the lines are the state of the state of the state of the state of the state lines are the state of the state of the state of the state of the state lines are critical states usery tools yields physical application processing state in states of the state line in the state of the state of the state of the state lines are considered in the state based in the state of the state of the state of the advisible to state the state line in the state of the state of the state of the state based methods and the state of the based methods are stated based on the state of the

The main body (drug listings) of the Anabolic Steroid Control Act has been included for your review below.

^{376.} Detection of randrolone metabolites in urine after a football game in professional and amnteur players: a Bayesian comparison. Robinson N, Taroni F, Saugy M, Ayotte C, Mangin P, Dvorak J. Forensic Sci Int 2001 Nov 1:2222-31:10-5

^{377.} Detection of norbolethone, an anabolic steroid never marketed, in athletes' urine. Catlin D, Ahrens B, Kucherova Y, Rapid Commun. Mass Spectrom. 2007;16:1272-75.

^{2002; 16: 12/3-75}

'(A) The term 'anabolic steroid' means any drug or hormonal substance, chemically and pharmacologically related to testosterone (other than estrogens, progestins, corticosteroids, and dehydroeniandrosterone). and includes--

(i) and cost ano diol... (I) 3b 17b-diludroxy-5a-androstane: and (II) 3a 17b-dihartroxy-5a-androstane (ii) androstanedione (5a-androstan-3 17-dione); (iii) and rostenedial. (ii) indicotential
 (ii) 1-androstenediol (3b, 17b-dirvdroxy-5a-androst-1-ene); (II) 1-androstenediol (3a 17b-dihydroxy-5a-androst-1-ene): (III) 4-androstenediol (3b, 17b-dihydroxy-androst-4-ene); and (IV) 5-androstenediol (3b 17b-ditudroxy-androst-5-ene); (iv) and cost enedione ... (I) 1-androstenedione ([5a]-androst-1-en-3 17-dione): (II) 4-androstenedione (androst-4-en-3,17-dione); and (III) 5-androstenedione (androst-5-en-3,17-dione); (v) bolasterone (7a 17a-dimethyl-17b-hydroxyandrost-4-en-3-one): (vi) boldenone (17b-hydroxyandrost-1.4.-diene-3-one); (vii) calusterone (7b 17a-dimethyl-17b-hydroxyandrost-4-en-3-one); (viii) clostebol (4-chloro-17b-hydroxyandrost-4-en-3-one); (ix) delwdrochloromethyltestosterone (4-chloro-17b-hydroxy-17a-methyl-androst-1.4dien-3-one); '(x) *1-dirwinstestosterone (a.k.a. '1-testosterone') (17b-hydroxy-5a-androst-1-en-3one): '(xi) 4-dihydrotestosterone (17b-hydroxy-androstan-3-one); (xii) drostanolone (17b-hydroxy-2a-methyl-5a-androstan-3-one); (xiii) ethylestrenol (17a-ethyl-17b-hydroxyestr-4-ene); (xiv) fluorymesterone (9-fluoro-17a-methyl-11b 17b-dihydroxyandrost-4-en-3-one): '(xy) formebolone (2-formyl-17a-methyl-11a,17b-dihydroxyandrost-1,4-dien-3-one); (xvi) firazabol (17a-methyl-17b-hydroxyandrostano[2 3-c]-firazan); (xvii) 13a-ethyl-17a-hydroxyoon-4-en-3-one: (xviii) 4-hydroxytestosterone (4,17b-dihydroxy-androst-4-en-3-one); (xix) 4-hydroxy-19-nortestosterone (4 17h-dihydroxy-estr-4-en-3-one); (xx) mestanolone (17a-methyl-17b-hydroxy-5a-androstan-3-one); (xxi) mesterolone (1a-methyl-17b-hydroxy-[5a]-androstan-3-one): (xxii) methandienone (17a-methyl-17b-hydroxyandrost-1.4-dien-3-one); (xxiii) methandriol (17a-methyl-3b 17b-dihydroxyandrost-5-ene): (xxiv) methenolone (1-methyl-17b-hydroxy-5a-androst-1-en-3-one); (xxv) methyltestosterone (17a-methyl-17b-hydroxyandrost-4-en-3-one); (xxvi) mibolerone (7a, 17a-dimethyl-17b-hydroxyestr-4-en-3-one); (xxvii) 17a-methyl-*1-dihydrotestosterone (17b-hydroxy-17a-methyl-5a-androst-1-en-3one) (a.k.a. '17-a-methyl-1-testosterone'); '(xxviii) randrolone (17b-hydroxyestr-4-en-3-one); (xxix) norandrostenediol--(I) 19-nor-4-androstenediol (3b. 17b-dihydroxyestr-4-ene); (II) 19-nor-4-androstenediol (3a, 17b-dihydroxyestr-4-ene); (III) 19-nor-5-androstenediol (3b, 17b-dihydroxyestr-5-ene); and (IV) 19-nor-5-androstenediol (3a, 17b-dihydroxyestr-5-ene); '(xxx) norandrostenedione---(I) 19-nor-4-androstenedione (estr-4-en-3.17-dione); and (II) 19-nor-5-androstenedione (estr-5-en-3 17-dione: '(xxxi) norbolethone (13b, 17a-diethyl-17b-hydroxygon-4-en-3-one); (xxxii) norclostebol (4-chloro-17b-hydroxyestr-4-en-3-one); (xxxiii) norethandrolone (17a-ethyl-17b-hydroxyestr-4-en-3-one); (xxxiv) oxandrolone (17a-methyl-17b-hydroxy-2-oxa-[5a]-androstan-3-one); '(xxxy) oxymesterone (17a-methyl-4 17b-dihydroxyandrost-4-en-3-one); (xxxvi) oxymetholone (17a-methyl-2-hydroxymethylene-17b-hydroxy-[5a]-androstan-3one): '(xxxvii) stanozolol (17a-methyl-17b-hydroxy-[5a]-androst-2-eno[3,2-c]-pyrazole); (xxxviii) stenbolone (17b-hydroxy-2-methyl-[5a]-androst-1-en-3-one); '(xxxix) testolactone (13-hydroxy-3-oxo-13.17-secoandrosta-1.4-dien-17-oic acid lactone): (vh toctoctorono (17h-badmvvandmet-4-on-3-ono)

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(AJ IESIOSETOIE (170°HJUIOAJAIRIOS***CIP5*OIE),
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'(xli) tetrahydrogestrinone (13b,17a-diethyl-17b-hydroxygon-4,9,11-trien-3-one);
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'(xlii) trenbolone (17b-hydroxyestr-4,9,11-trien-3-one); and
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'(xlii) any salt, ester, or ether of a drug or substance described in this paragraph.';

Austria

The possession of anabolic steroids is not a criminal act according to Austrian law. In 2008, Austrian government officials announced intent to place criminal penalties on steroid possession.

Australia

It is a criminal act to import, supply, use, or possess anabolic steroids in Australia without a prescription from a medical practitioner, dentist, or veterinarian (Poisons and Drugs Act Amendment of 1994). The outlined penalies for possession without a legal prescription include a maximum of 6 months of imprisonment and/or a fine of \$5,000.

Canada

Anabolic steroids are included in the Caradian Controlled Drugs and Substances Act as Schedule IV substances. It is illegal to sell, manufacture, or import anabolic steroids into Canada without proper legal authorization. Possession of anabolic steroids for personal use is not a criminal act.

Czech Republic

In 2008 it became a criminal act to manufacture, import, export, store, or distribute anabolic stroids in the Czech Republic. The potential pertailase include a maximum of 3 years in prison. It is not a crime to possess stroids for personal use.

Denmark

In Demusk it is a crine to manufacture, inpost, export, murket, disperse, distribute, or possess doping substances including anabolic steroids, human growth hormone, and erythropoietn without proper medical or scientific reason (The Act on Prohibition of Certain Doping Substances No. 232 of 21 April 1999). The potential penalises for possession include a maximum 02 years in prior.

France

In 2008 it became a criminal offense to manufacture, transport, acquire, or possess doping substances including anabolic steroids, human growth hormone, and erythropoietin in France. The potential penalties for possession include a maximum of 5 years imprisonment and/or a 75,000 Euro fine.

Greece

The possession of anabolic steroids is not a criminal act according to Greek law. In 2008, government officials announced intent to place criminal penalties on steroid possession in Greece.

Sweden

In Sweden it is a crime to import, manufacture, transport, sell, possess, or use doping substances such as anabolic steroids and growth homone without proper legal authorization (The Swedish Act prohibiting certain doping substances (1991,1969)). The potential penalises include a maximum of 2 years in prison. Possession for personal use is usually regarded as a petly offleres and given a maximum penalty of its moths imprisonment.

United Kingdom

The importation, possession, and use of anabolic steroids are not criminal acts according to UK law. There has been a great deal of pressure in recent years from the U.S. and World Anti-Doping Agency to place criminal penalties on the possession of anabolic steroids.

Acquiring AAS (Best Practices)

This section assumes that the decision has already been muck to use anabolic starchist. It phones no judgeren on the decision local and concentration copy on callenge risks relative the data supply. How one goes allowed obtaining fixes atlastices is an example inprotant anabolic storek does not be a highly working how, while system quarking solutions and anabolic storek does not be a highly working how, with system quarking solutions of the store of agreement of the store of all store is store small stores, relative store of the bayer. The main delector is no constraintion, days absolution, or maximum does and on the store.

By Prescription

Provided you are in some way is usid need of therapy, it is always preferable to obtain anables acrossib travely an physicitry preservition. In addition to collering the potential baseful of lowing some nucleal oversight to your use, more plasmacential markets (in control of the physicitry preservities) and the physicitry of the physicitry of the States, in fact, the risk is so low that it is not even worth considering. Furthermore, if you results in a country where there are critical peakles for the supported possession of strucks, a prescription it fit only use to happy count and the sub-structure. The critical structure is the sub-structure of the support of strucks are prescription if it only use to happy count and structure is the sub-structure of the support of structure of the support structure of the support of the support of structure of the support of structure of the support of the support of the support of the support of structure of the support structure of the support of the support of structure of structure of structure of structure of the support of structure of

Most realess are probably under the incorrect assumption that is nearly impossible to get a processipion for analysis sensiols in the Unider Status. While their them yhow been true following the doping searchish of the line 1980s, the matter has changed considerably size: who end go compares assumed to the season of the season of the season of the order of an ends (specifically textustreeness) to true agree related hornow of docknery in run it is a fact thru read so with hornogen characteristic season levels directly approximately assumed as the season of the season of the season of the direct season of an ends with hornogen characteristic season of the direct season of the seaso

If you feld that you are a candidate for a prescription, yourny need to take an active nois in finding a physical. Unfortunately, may doctors are all sursus about hose dungs. They might not prescribe testostenore at al. or may be too conservative in their treatment fifty yol. You may have to iterative weareal heteirs you full the or that sing first you. Dort's overy about "discuss shopping". This ilegal practice involves seeing malpie doctors, or withholding information, to any mescriptions you don't need. Volu sobald were the to your doctor, or and any descriptions from more than ore at a time. Oftensise, it is perfectly legal to take the time to that one you dock comfortable with.

Makial necessity is unally apported with a blood test, ordered by the tearing physician. Teastorence toes the total to be lapkest in the moring and lower at the end of the day. The blood test should be phened accordingly. Before reviewing the results, understand that there are on specific stability allowed for process for tearing men with hormone deficiency. The medical community, does not even agree on the post in which thempy should be initiated. For example, the "hormal" level of fusistances are and attern as '20-2100 right. Comtaged, the starting level of fusistances are the post of the starting and the s

How testosterone medication is prescribed also varies a great deal depending on not only the needs of the patient, but also the comfort level of the physician. For example, more rigid nhosicians will often only prescribe testosterone aels or natches. This is because these forms abov the very controlled duly release of textsterore, and are harder for the patient to abuse. Other dectors are confirmable working with textsterore injections that are taken how man dived durinities of by the patient. Even when textsterore injections are proce-roll, some doctors relate to give more than 100 mgere week. Others outlow give a 200 mgper week door, assing that how breeks are not charden at any point draing reatormer. To wait not need to ful a crooked ductor. You just want to find a progressive physician that understands the vulue of optimizing work momene levels.

Van hue a few options open to you when first seeking treatment: You can approach your finally doctor, a horness specialis (modernologist), or an anti-singing physician. In any experience, anti-singin medicine is where you find the most progressive physicians and treatments. These are the doctors in final are working with both its stroids and of Helder medications file anti-scores in find fill are also excitation of the physican particle through the strong strong the physican particle and the physican particle and the structure. Academy of *Intel Apple Moderne (Apple Constanting Constanti*

One beneft to being a writer and clucator on ambolis substances is that I are regularly meeting prograssion positions and hormore clinic owners. Ne had the plosance of gatting the otis star, and contraining to goor mightly located to a strate the skel of metricine is have a signary metric or the system. It has given me confidence that this skel of metrics me in a single position to help potential HBT/metricsging patients. Male is isotostence addisence y gasory and devended in this courset, I' alle to do sore parts to help change faith balow I alluse it as a pertain part withen in courset with HBT/metricsging photesian. If you have it is the strate of the balow I alluse it as a pertain part withen in courset with HBT/metricsging photesian. If you that, HBT my be any one phot strate you recompare you to explore it faither, regardless of flow you go about seeking metamerer. Supporting the physician that offer this type of retratement is the straty with photesian alluser.

For hormone replacement therapy & anti-aging doctor referrals, visit the website www.HRT-Rx.com to find a physician in your area.

Ordering Steroids in the Mail

Unlike more traditional drugs that are ranker strict lagd controls (starks as morotics), the sale of numbols storods some to be operly advanticed on the Internat and in the back pages of bodyballing magnitudes. Mack of this probabily has to also with the storage variance in inso the observed particle back strict the storage variance in the storage particle back strict the storage variance in the storage strict the storage variance is storage with storage stores the stores.

Below are several rules to follow when ordering steroids through the mail. These rules can help increase the chances that your order will contain pharmaceutical grade drug products.

 Order from a country with legal access to steroid medications. If it is easy for your source to get legitimate products, it is likely what is sent to you will be legitimate. Avoid ordering from country is utilt origin modelwine bares such as the United States Council. Western non commes with start promotive laws such as use transit states, canada, recently Europe, or Australia. Consistent supply in these areas invariably involves the sourcing of clandestriely manufactured products.

 Buy local brand drugs from the chosen source country. If you are ordering from a pharmacy in Thailand, buy Thai drugs. The chances of running into counterfeits will increase significantly when the drugs you buy are not domestic to the supplier's country.

 Immediately avoid any foreign pharmacy that says it carries American steroid products. These will be counterfield over 99% of the time, which means the pharmacy probably deals in counterfield trans.

4. Buy from a place that shows pictures of its products. Scrutinize the offerings closely. Are there any known fakes? Many suppliers work hard to keep their company clear from counterfield rugs. You want to deal with one of these.

The propose of this book is to reduce hum, not encourage or support crimital activity. Thus the previous rules address only the utility of the dang, not strangiffer that Tyro unside in an area where stravids are illegal to import without a prescription such as the Linda States, Cranda, Scandinivo, or Anstrain, yoo meet on to be awave that three can be scrinos lagel consequences to ordering these dauge frough the mult Handonitis intercept your package, you may be charged with a variety of errors including dang procession and legal importation. No may scene be charged with multidiag fath quantity is large arough. While, document, the project with an attering of orders. Breight produce the legal justice system is not the outcome you ware. Please consider this carefully before you order steroids through the multi.

Purchasing Steroids on Vacation

Ancher commo option is to travel to a coarty where anabolic storoids are widely anabile, and protess the drug directly form a pirturesy. Have propuls vacation spots are commo sources, relating Morico, Thuihard, and Tutkey, So F is not uncommon to see dual-propose trip coardinationarizing, with the individual brings they coard and and the support. The principle bendity, provide the individual chooses the right coarty and shows relating with bendit be more consistent access to and patrometerizing after coarty and shows relating the bendity provides the start of the start of the start of the start required to their Character paper variable start of the start of the start of the start while attempting to exter a coarty with steroid prohibition laws, you could face seriess criminal charas.

In an effort to help readers avoid the added risks of counterfeit drug products, the following points of advice are provided. All readers are cantioned that smugging is illegal and can be a very risky practice. Again, since this book is about harm reduction we will not be discussing means of smugging.

 Loose regulations on the steroid supply typically also mean loose oversight of pharmacies. In some countries, pharmacies in torrist areas will commonly stock counterfelts. You should become well-versed with the products of a particular region before traveling.

2. Always buy local. You want to get the drugs that are readily available to pharmacies through normal legal channels, not exotic drugs that may or may not have been legally imported by the pharmacy. For example, never buy European or American drugs in a Mexican pharmacy.

If possible, avoid the obvious tourist pharmacies. Travel to an area more known to the locals. The smaller a percentage of business the pharmacy makes on tourists, the less likely they are to be stocking counterfeits.

4. Ask for products by name. Most tourists starable in and ask for "steroids, anabolicas, testosterona." If instead you ask for something very specific, such as "Bayer Primoteston," you are probably going to be taken more seriously.

The Local Black Market

Again, our focus in this section is only on increasing the likelihood of obtaining real plarmaceutical products. Towards that objective, the following points of advice should be followed when shopping on the local black market.

 Consider the country of origin for each drug. Only buy drugs that make sense. For example, Mexican steroids are common in the United States due to their open supply and close proximity. You would expect to see them. On the other hand, you are very utilkely to find legitimate drugs from Australia in the United States, which is very far away and also serioally ensities is own steroid supply.

Try to stick with a group of fellow steroid users. Sharing experiences and knowledge can help keep the group better supplied with legitimate medications.

Try to buy from other steroid users. Individuals who sell steroids but don't use them are probably less concerned with product quality.

4. If you are trying to purchase legiminte pharmacertical products, don't buy multi-dose vials or bottles with loose pills unless you are absolutely sure of the company and source. These forms of medication are ranely used outsile of the U.S. and Carada. Single use ampulse and push through pill bisters are much more common in other countries, and may be sightly more second use to the difficulty involved in multing them.

Part III

Drug Profiles

Agovirin Depot (testosterone isobutyrate)

| Androgenic | 100 |
|------------|----------|
| Anabolic | 100 |
| Standard | Standard |

| Chemical Names | 4-androsten-3-one-17beta-ol beta-hydroxy-androst-4-en-3-one |
|-------------------------|--|
| Estrogenic Activity | moderate |
| Progestational Activity | / low |

Description:

Testosterone isobutyrate is an injectable steroid preparation that contains the isobutyrate

ester of testosterote n a vater tose, Arnorg pooptuater, testosterote sontyatet a often consideral andgogan to testosterores asgeresino (no seter). Allongio han esuali y fond as vater-based sapensies, the planmackiteris of the two products are admittably very different. While testosterone (freis) approximations is very fast-artific rapitir ajpeciritos to be given overy fow days, testosterone isoblytme is much slower to release, and is usually admittated caree corey two weeks in a critical settirg. As an approximation of the size of the size

History:

Injectable stotenerse isoloptrum microcytal supersitive uses first described in 1952.²⁷⁹ The agapt use developed in an effect to column in signetike (deyon) from for stotenerse flut would be shower acting that regulate (Hee) testsitenerse aspension or insistenerse assocribialised proveding a microcytable depositive of the stotenerse assocribialised proveding a microcytable depositive of the stotenerse insistence of free stored in the blockbergum Albungh efficience for the parpose, testsitenerse of free stored in the blockbergum Albungh efficience for the parpose, testsitenerse insistence was developed at its modern many modern and the stotenerse insistence was developed at its modern many modern and the stotenerse insistence insistence and the blockbergum Albungh efficience for the stotenerse insistence insistence and the tests of testsitence was being introduced to the minder, By the mail-1960x, testsatence constitute would during bayes the commercial access.

The only medium staroid product to use testistences isolatynate is Agavirin Depet, developed by Biokin in Carchitovikui, this primarily preservable to trust miles with instificient androgan levels and addiscents with delyed pheory, although it is indicated for an using of other prepares including the transmittor of Kindeler systematic (addissing without (a disease without its an arold and grant preservice limits, prosting-ingrant and and a strategistic), splanking means. Cossing's systematic issa analodis anget to preservic learning, particular means, Cassing's systematic issa analodis anget to preservic learning, particular means, Cassing's systematic issa and anget on preservic learning, particular means, Cassing's systematic issa and anget on preservic learning partice and the body due to severe illess's Agavin's Depti si sill produced by Biolai (carcerd) in the Shoval Republic, and terminas pondure acordo to European bleck merkets.

How Supplied:

Testosteoro isobrynte suspension is available on the human drug market in the Slovak Rexploits a Agovin Doped (Bothal). It cortains 25 mg/mol stread mixed in a waterbased solution; packagad in a 2 mf, ampule β ampulse per box). Testosterone isobrynte has low water solubily; the steroid will noriceably separate from the water-based solution when an ampule is left to sit. A quick stuke will temporarly place the drug back into supervision, but the withdrawn dossed solutid lawns be consistent.

Structural Characteristics:

Textsteres isologue is a mediad ferm of textsteres, where a carboxyle and ester Carothyl proporties of 10m keen attached to the 71-beh plotogya isogon. Esterifical formes of associatorea are less polar fram free textstactorea, and are absorbed more slowly from the area of injection. One in the bloddstarms the ester is enremedy to lydd feet (action) textstactorea. Esterified ferms of textstactorea are disque to probeng the wirkow of themposet: effect blodwing administration, advong for a lash framet injection textstactorea. Esterified ferms of textstactorea are disque to probeng the wirkow of themposet. effect blodwing administration, advong for a lash framet injection schedule microscytallitor supervision is disqued to provide physicaligal androgen concentrations for approximately acceleration.

Side Effects (Estrogenic):

Teosterone is readly aromited in the body to estability entropy. The aromitses (everyon particular) compare is responsible for fine metabloom for instructions. Teostorome, Elevation suggestion of the structure is considered at the structure of the structure and such as characterise is considered at more by necessary in prevent astrogenic add at the structure of the structure of the structure of the structure at the structure of the structure of the structure of the structure at the structure of the structure of the structure of the structure is the structure of the structure of the structure of the structure is the structure is the structure of the structure of the structure is the structure is the structure is the structure of the structure of the structure is the structure is the structure is the structure of the structure is the structure is the structure is the structure is the structure of the structure of the structure is Enorgenic side effects will occur in a doss-dependant memer, with higher doss (above normal therapetic level) of lostioaterem one fieldy to regard the occurrent use of an artisatiogen or aromatose inhibitor. Since water referrion and loss of matecide definition are common with higher doss of lostioaterence, tike drugs is using down of the observed provides and the since the state of the since the since the since the highing places, where the added stater reteriors will support raws integrab and matecids since and heps foster a stronger attrobic environment.

Side Effects (Androgenic):

Teastores is the primery multi-androgen, nesponsible for maintaining soccordary mulssenal characteristics. Enseted here's of testorene are likely to produce androgenic side effects including objective and the setting of the setting of the setting of the probability of the time of the setting of the setting of the setting of the here of the setting of the setting of the setting of the setting of the product of the setting of the setting of the setting of the setting setting of the setting of the setting of the setting of the setting setting and the setting of the setti

Side Effects (Hepatotoxicity):

Testosterene does not huw hepatotoxie effects, ber toxicity is triffely. One study examined the potertial for hepatoxicity with high does of testosterene by administering 400 mg of the hormore per day (2,800 mg per week) to a group of mile subjects. The stored was station only so that higher peak concentrations would be reached in hepatic fissues concentration of the store of th

Side Effects (Cardiovascular):

Ambolicantegonis strends can huve delutricos effects on serum chosterort. This includes a tendency to reduce JHCL (apod) chosteroval values and increases JLC, (bad) chosteroval values, which may shaft the HDL to JDL bahare in a direction that florous gener risk of antirochorsonis. The evalue impact of an antibiciantingenis extended in serum lipits is dependint on the dose, route or a daministration (eral ws. lipitedbal, type of steadid Ambolicinstrugenis extended and the steady of the steady of a steadid Ambolicinstrugenis extending and support left wertricult physicares and influencies, rockae extendedini reducesianti and reasonal influencies.

Tensistence tends to have a much loss damatic impact on cardiovascular risk factors dam synthest strotch. This is dae nay to its operaness to matchiombra be focus, which and the second structure of the second structure of the second structure of the source structure of the second structure of the second structure lipids. In one study, 220 mg per vock of its structure of the second structure structure of the structure of the second structure of the second structure structure of the structure of the structure of the second structure of the struc

and the second second

Due to the positive induces of estimgan on serum input, surroutive entries or chromebars, entries are preferred to acommutes inhibitors for those coverned with a calcious-calar heads, as they offer a partial estrogonic effects in the free. This also them to poterinfly improve fully onlike and offers some of the engines effects or almongens, which does of 600 mpc learners, milling and mis-terogong for calculatory the probability of the probability domains, milling and mis-terogong for calculatory the probability of the probability domains, miniting and mis-terogong for calculatory the probability of the probability in LID/LDL chickened triggerowing, applications probability and the probability domain semativity at learning a probability of the probability of the probability domain semativity at learning a probability of the probability of the probability domain semativity at learning a probability of the probability of the probability down and involution dows, injectible teststerome esters are usually considered to be the safet of all antiobeticing are velowing the probability of the probability of

To help reduce cardiorascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fash oils (4) grams per day) and a natural cholesterolamicoxidant formain such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicalização stavidi velue taken in doses adficient to promte masek gai are expected to aspresse endogravos testostores producino. Testostenos is he primury mar androgan, and offers strong negative fordback en endogravos testosterore production. Testosterone-based lange villa, Boviech, here a strong effort on the hyporhamic regulation of natural stearoil hormoses. Without the interversion of restosterone simularing substances, testosterone based and termino normal within 1-4 monti of dinarga secsasion. Nore that prolongiant plangarandorophic hipograndism can develop secondiny to steroid abuse, messanting medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

The design of testosterone isobatynte (ss Agyvini Depot) is sightly different than that of most testosterone estes, which are usually made as only solution. Agyvini Depot instead contains a microcrystalline aqueous suspension. The crystals form a repository in the muscle following injection, where they solved wild solve over time. Injections of testosterone isobatynate may require a large needle (21 gauge), and may result in local irritation, pain, and redness.

Administration (Men):

To text and any an indicatory, testostarene isobaptise suspension is usually administered in a done of 59-100 mg every 14 days. When oue dof ranacle-bading process, testostrone isobaptine suspension is often administered at a dose of 200–400 mg (4-8m) per week, down of the start of the dot doses and testostery for pain at the size of opticient (large inspirator solutions) and the indicator of the start of the start of the start of the start of the dot doses and the start of the start start of the and 12 works in implicit This level is sufficient to provide insteaded gains in made, size and of ne ambiect mitoresis starts descering in the dose of the start of the start of the start of the mathematic matters in the start of the mathematic mathematic starts descering in the dose of the start of the start is a start of the mathematic mathematic starts descering the dose of the start of the sta

Administration (Women);

Testostenoe isobaryate suspension is not commonly used with women in clinical medicine. When applied, it is usually given in a dose of 25-50 mg every 14 days. Testostenoe isobaryate suspension is not recommended for women for physique- or performanceeritarizing purposes due to is strong androgonic nature, tendency to produce virilizing side effects, and show sering characteristics (maleing bod used difficult to corrol).

Availability:

Pharmaceutical preparations containing testosterone isobutyrate are rare. The only known

product at this time is Agovirin Depot, produced in the Slovak Republic by Biotika. It contains 50 mg per 2 mL ampule; 5 packaged per box.

Anadrol®- 50 (oxymetholone)

| Androgenic | 45 |
|---------------------|---|
| Anabolic | 320 |
| Standard | Methyltestosterone (oral) |
| Chemical Names | 2-hydroxymethylene-17a-methyl- dihydrotestosterone |
| | lihydro-2-hydroxymethylene-17-alpha-methyltestosterone -2-hydroxymethylene-17-hydroxy-Salpha-androstan-3-one |
| Estrogenic Activity | high |
| Progestational Act | ivity not significant |

Description:

Organizables is a poter cell analysis stroid derived from dipotestosterose. More specifically it is a document of the strong of

Oramchhoen is considered by mmy to be the most powerful steroid commercially available. A storid movie experimenting with its agart is failey again 20 to 30 ponds, it of water steroids, and a can often be accomplished within 6 weeks of tuse. This storid produces is often of fitted consequence to the user, who may be failing ways by and storag which taking commendue. This is powerful as the storage fitted product and the storage fitted product as a discussion of the storage fitted the storage fitted product and the storage discussion of the storage fitted production in the form of each start had fram the chance for inpay when fitting haves, it should be readed, however, that a tary grad gain in mass single also phenes to much storage storage concerned in starts. The there is storage storage storage concerned to instarts that are storaded gain and heights to much there so any concerned to instart.

History:

Organizatione was first described in 1999.³⁴⁴ The agent was relaxed in the United States as an prescription during the early 1990/s, solid turch the burd mares Androl-50 (Systec) and Androjd (Parke Davis & C.o.). Systex developed the agent, and would hold apport rights to it unlift everption runny spaces later. The day was original pappored for use in confidence shows underweight attas, preand holdinator, chrone underweight attas, preand postparticity exercision of later marks, and the obligation of the strength of the strength of the strength of the distillation of the strength of the strength of the strength of the strength confidence. The recommended dates for such uses was standy. 25 mg three intes per day The data was originally sequed in 2.5 mg, for use 10 mg hole.

In sphe of the many potential therapeutic uses or a strong anaholic activity of this drug, the FDA soon strictly narrowed the indicated uses of oxymetholone. By the mid-1970's, the drug was FDA approved only for the indicated treatment of amenia characterized by deficient red blodd cell (RBC) production. Admittedly the strinulation of exptropoissis is an affect that is characteristic of nearly all anabole steroids, which as a group tend to increase affect that is characteristic of nearly all anaboles steroids, which as a group tend to increase the state of the steroid steroids of the strinulation of the structure of the increase the state of the steroid steroids and the steroid steroids of the steroid steroids and the steroid steroid steroids and the steroid steroid steroids and the steroid steroids and the steroid steroids and the steroid steroid steroids and the steroids and the steroids and the ste BPC concentrations. Openations, however, seemed fairly reliable in this regard, domenstrating an increase in exploreposite livels and not 8 5 641.³⁵⁵ This has led to its adoption for this relatively new medical use, as well as the institution of a higher (50 mg) dosage with the updated Anadrid-50 product, necessary for a stronger effect on REC cours. The Parke Davis term would not be brought up to the higher dosage, however, and was discontinued.

Recert spaces have brought fourth a namber of new treatments for anotin totably Figues (recordinate orderprojection) and related symbosytics periods. These diagonal directly minite the body's instant of albodie call producing hormose, and as such provide a directly minite the body's instant of albodie call producing hormose, and as such provide and the symbol of the symbol of the symbol of the symbol of the directly minister and the symbol of the symbol of the symbol directly minister and the symbol of the symbol of the symbol directly and the symbol of the symbol of the symbol of the symbol decided to drop this fam in a number of foreign countries. Fremant from Switzerland and minister with this ness of countributes was not been of the symbol decided to drop this fam in a number of foreign countries. Fremant from Switzerland and minister with the symbol countribute was not been on the symbol of the symbol decided to drop this fam in a number of foreign countries. Fremant from Switzerland and minister with the symbol countribute was not been on the symbol.

In hdy 1997, Syntex sold all rights to Annole-50 in the U.S., Canada, and Mexico to Uniternel Pharmacenicals. Uniternel irreduced Annole-50 to the U.S., Canada, in 2014, Sintimer Uniternel Pharmacenicals. Uniternel with HW are commonly americ, often caused by the disease loted (provatins it infections, et al. and an output of the disease loted (provatins it infections, et al. and the disease lot and the disease. The american in HW patterns is signally catagorized by impaired ned blood cell production in house marrows, the FOA approved indication for comprehense use. Adding to this, Unitered soon initiated Phase IIIII risks with Annola die HW vasting syntheme, real continued to research to active that the soft heating such their obstruction pharmady classes and ipolytophy (a closeder characterized by a selective loss of blood fat, insilin resistance, distructions, blockmark lowed, and all physics).

In April 2006, Solvey Planmencicals (parent company of Unined) soil the rights to Anadol-50 to Ahmen Planmencical, LLL: Ahmen cortinats to Munic He dug in the Uindel States, although given the maniton it is urestrini with additional uses the company plants operase with controllose. At the present im the out PFOA approved indication memins that of treating red blood ed defaciert anemis. Sprates seems to have removed head from the osynchrobace. At the present im the out PFOA approved indication study of the study and the study of the study and the study and the study of the study and the study of the study and the study a

How Supplied:

Oxymetholone is available in select human drug markets. Composition and dosage may vary by country and manufacturer. Most brands contain 50 mg of steroid per tablet.

Structural Characteristics:

Oxymetholone is a modified form of dihydrotestosterone. It differs by 1) the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during oral administration, and 2) the introduction of a 2-hydroxymethykne group, which inhibits in metabolism by the 3-hsd enzyme and greatly enhances the snabolic and relative biological activity of methydrotestostosterone.

Side Effects (Estrogenic):

Organetholone is a highly estrogenic steroid. Orgacconstiti is often a concern during tutanter, and may prosert itself quice early into a cycle (particularly when highly closes are used). At the same time water reterion in can become a problem, causing a notable loss of macke definition as both subcataneous water reterion and lit levels hull. To avoid strong estrogenic side effects, it may be necessary to use an anti-estrogen such as Nohadee® or Chmidle.

It is important to note that oxymetholone does not directly convert to estrogen in the body. This steroid is a derivative of dihydrotestosterone, and as such cannot be aromatized. Artiaromatises compounds such as Cytadren and Arinides® will, likewise, not effect the relative entropenitive of this steruit? Exemptions have successful due the hist have of controusing and in a start or in comparison of the strends. Some rares suggester and the right seven or exonogene accurst prisourceflockies is activately due to the ding acting as a propositiva similar to matchedore. The side effects of both estrogene and progesitions can be very similar, which night how made this explanation a phasical sector activation of the similar distribution activity sequences and the similar distribution of the similar distribution of the similar With such findings, it seems more phasicle that measurements the strength receptor, similar to priore productively that, the estrogene measurement distribution of the strength research of the similar distribution of the similar distress distribution of the similar distress distress di

Side Effects (Androgenic):

Albrady constructions is clossified as an anabole storedl, antiogenie skie effects are still goods with this sholtance. These may isclusible boots of oly days, and hody/facilitati goods. Higher doess are more flexly to cause such side effects. Anabolerantloganic storesisti may also argument me pattern hair toos. Wronen are additionally surred of the potential vitiking effects of anabolicantrogenic storeds. These may include a depening of the voce, messarian lengathrics, charges as its netterse, ficial barg roots, and alconal endrogenere. While Anaboli is clossified as an anabole storiki, it does retain a nabale andoognic component.

It is intensity to note that comparison does exhibit some tendency to convert to displorentsorbarene in the body, although this does not occur with the s-dphn reductions enzyme. Doparthelikers is altendy alughdorentsoraneous based statistic, som as statistication sometholes different sometholes and the second statistic second statistic grouping can be removed metabolically reducing compatibilities of grant analogue Trajbar-medul diffusionationerus (metabolical). The third addre that the biotransformation combutous at alten on some level to the analogue trains and the distribution of the second statistical stat

Side Effects (Hepatotoxicity):

Organizations is a $(1^{-1}a)$ -fra alsoluted composar. This alteration protects the drag from descritation by the long, allowing a vary high presenting of the drug entry into the blockstream bibwing and administration. (17)-alpha alsoluted ambolic intrologenis stronks can be hapstooks), chooling and high proposite may use all its med ambolic intrologenis stronks can be hapstooks). To choosing all of high proposite may use all its med range. In margin terms estimates and hege to maximize the fraction and overall beamh Intake of (17)-alpha alsoluted acrossite is correctly infrared to 6 × 9 weeks, man effect to avoid excelling the restricts of the strong strong

Organizatione has a stantistic Avering, which slightly encodes is relative hepatroxics; $y^{3/2}$ Sig. this staget, particularly at the doors commonly used, can present substantial hepatroxicsity to the user. Statics administering 50 mg or 100 mg daily to 31 delayed wrant for 21 useds produced lengthmat tenewares in here organize (manametase S-ST at ALT) and and exceeding one year (in some patient) has determediated elevations: in ygitamphranetisme, Cer(7) in 17% of patients: significant tenesces in hisbihn in 10%, and serum abarria increases in 20% ³⁰⁰ One patient developed a liver smort that could have been plooks hepitari, in differentiation galences excited materiates with the state of the

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anabole/inflogarie steroids, can ihwu dekterious effects on serum cholsteroil. This includes a tenfency to proken. PLX, (good) cholsteroil valasa, which may dish the PLX, (good) cholsteroil valasa, which may daif the iFLA, to LD, balance in a direction that forces genater risk of antiroidenoisi. Bir tendities rigated of an anabole/inflogaries steroid on sensure lipids is foromatinghe or non-sensuration hyper and level of resistance to happine methodism in Anabriciantowise' strength, may also also works abole Thologaries methodism and theory inflogaries and the strength of the strengt reduce endothelial relaxation, and support left vertricular hypertrophy, all potentially increasing the risk of cardiovascular disease and myocardial infarction.

Organizationae ina satorag effect on the lopatic munagement of choicestrol due to its structural resistance to here breakdown and rator of administrations. Studies administering 50 mg or 100 mg dashy to a group of dashy ment for 12 works have demonstrated nignificant risonass in 11.24 choisend, accompanish by very significant (damind) appressions of range citely). The mess of constraints with the under the structure of the structure of the structure of a structure layor of barrel dama of the structure of the structure of a structure layor of barrel dama of the structure of the structure of a structure layor of barrel dama of the structure o

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of star-tated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4) grams per day) and a natural cholsterol/articoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All ambok'androgenic steadis when taken in doese sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone simulating substances, testosterone levels should return to normal within 1-4 months of drug secession. Note that prolonged hypogenadorophic hypogenadism can develop secondrus to steroid abuse, necessisting medical intervention.

Note that when discordinging comparaboues, the carabic came has a capadity provering as the enocycle results. To beyong with the best of outwards reterrism onli agaids, dimainsk, dropping the user's hody weight diamatically. This should be expected, and net of francel concern. What is an additional the structure of the struct

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴⁸⁸ This is caused by the fat-soluble nature of steroid hormones, which can also some of the drug to dissolve with undigested dietary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

Tarly preserving anklens for convertedore recommended a dosage of 2.5 mg three times per due to reverse the wanting process and provide law hody muss aim. Does as high as 30 mg were employed in some cases. Currer prescribing guidelies recommend a dosage of 1–5 mg pet kägmen of Dodyweight pet dys be trusting anorni, altohogh rakitsel that a dosa of 1–2 mg pig a hypically sufficient. A 175-pond person would take approximately 150 mg per day at the 2 mg/d dosage level in some other contrust, it is reasoning of the most of start and for physicase or dys reformance-interacting propose, an efficient to its morth. When and for physicase or reformance-interacting propose, an efficient 6–5 works to unimize hypatoxisity. The level is antificate for harming its most simulation 6–6 structure, the most propose, an efficience of heating its morth simulation of mere to the number of the dosage of the simulation of the top the simulation of the top 6–5 works to unimize hypatoxisity. The level is antificate for harming its morth simulation of the day at the high proteinia for hepatoxicy. When used for physicaor of the day at the high potential for hepatoxicy. When used for physicaor performance-networks propose, an efficience on day dosage wordd high in the marge of the day at the high potential for hepatoxicy. When used for physicaor performance-networks propose, an efficience on day dosage wordd high in the marge of 25-150 mg, taken in cycles lasting no more than 6-8 weeks to minimize hepatotoxicity. This level is sufficient for dramatic increases in muscle mass and strength. Higher doses are rarely administered due to the strong estrogenic rature of the drag, as well as the high potential for hepatotoxicity.

Administration (Women):

Prescribing information for oxymetholone in the U.S. makes no distinction with the dose for females. Oxymetholone is generally not recommended for women for physique- or performance-enhancing purposes due to its very strong nature and tendency to produce vitilizing side effects.

Availability:

Pharmecutical preparations containing conventions are fairly inrited. The legitimute sapply seems to be scattered into isolated markets of Europe, Asia, and the Americas. Most of the supply for this dang comes in the form of undergound and export-only products. In reviewing some of the remaining products and changes in the legitimate global pharmaceutical market, we have much the following observations.

Anapolon (Turkey) was recently updated to reflect a new logo for the manufacturer Abdi Ibrahim. The product is still packaged in boxes of 20 tablets each.

Oxymetholone-Alhavi (Iran) was recently updated. It is now packaged in foil and plastic strips of 10 tablets each, 10 strips per box.

Oxymetholone IH (Iran Hormone) was also updated recently. The presentation remains similar (strips of 10 tablets, 10 per box), though the box looks more modern with gradient coloring.

Oxyanabolic from Asia Pharma (Malaysia) is now registered for sale in Thailand. The product comes in strips of 10 tablets, each with an errbedded AP and 50. The product should also carry a unique product ID marmber that can be verified on the company website.

Androlic from British Dispensary (Thailand) is also still in production. The product comes in botties of 100 tablets each. The tablets are green, scored on one side, and stamped with the BD snake emblem logo on the other. The product carries a holographic sticker to deter counterfeiting, through even this sticker has been duplicated with high accuracy.

Balkan Pharmaceuticals (Moldova) makes the product Anapolon. It is prepared in 50 mg tablets, with 20 tablets contained in each foil and plastic strip.

Oxybolone from Genepharm Greece is again in production. It contains 50 mg of steroid per tablet, with 10 packaged in each foil and plastic blister. Two strips are packed in each box. This product is widely distributed through export.

Oxitoland is manufactured in Paraguay by Laboratorio Farmaceutico/Landerlan. Each box holds 2 strips foil and plastic blisters of 10 tablets each.

Anadur® (nandrolone hexyloxyphenylpropionate)

| Androgenic | 37 |
|------------|--------------|
| Anabolic | 125 |
| Standard | Testosterone |

Chemical Names 19-nortestosterone hexyloxyphenylpropionate 17beta-Hydroxyestra-4-en-3-one hexyloxyphenylpropionate

| Estrogenic Activity | low |
|-------------------------|----------|
| Progestational Activity | moderate |

Description:

Nandoken hesykoophendynopisonin is a slow-acting injectable form of the analoxis associar matorism. Elsevkoynehendynopisonica, also calkad parahoyskophendynopisane, is a fairly anasali matokane assist in a structural associ. It is seasefully mathokane more carbons, it is the langest, and likely bekwest acting, set of ramidone to evel structuration of the langest, and likely bekwest acting, set of ramidone to a set analosis in a matchal assiste. Nandostrue hesykopisotephynopisate is a bager from steades in a matchal assiste. Nandostrue hesykopisotephynopisotepisate in a bager promets dow staduy lapin is harm sow bit hus ostrogatier and advangers side offices.

History:

Nandroken hesybestynehytenyinenti was fiet dascreden il 1960.²⁰¹ It was developed into an encieris abstyn sitt, and was ood irmalyti under fier Andrah Fendar Tumen is asch mittekst sin Austrin, Stovelen, Switzerland, Belgiam, Netherlands, and Germany, Anadar pensised fiwegash tearity 1990, and, Bokings means rungers and acquisitons, was soilt mairly by Kalle Planmenas. Kalvi social also alf fie drag in France, but nache the brand runne Anador. matimism and cherotherupy tabuloral appression of rule bloot of incorrell, but was also used as a goveral lane-issue-building anabole, with certain diseased states, rigar, or consultances.

Nandoken henykosphenykopoinute wan en wiekdy dimirhaed ontakis of Europe, and in spie of a long listory of relative saftyycould not last as a mediatal product. The 1995 merger of Galiw with Lypkin famula the company Plannenic & Uppin, and would soon Austin, but of hybride, and non-refeated to Galirga to climita the farms of machines henykosphenykopoinute in al courtiss. It is of route that this ester of mathoken was also sold by Los 8 Spin, and son refeated to formany. Changes and the spin heavy of the spin straines and the strain of the straines of mathoken was also sold by Los 8 Spin (Auduh). Larbacken Tomennik, (Auduh, Eacchesis in Tittage (Analah), and by Xpensi it (Konce Anadarai), matsi under direct Konce with Koli, AJ Heavyophenykophenet is no long available as a commercial matchine.

How Supplied:

Nandrolone hexyloxyphenylpropionate is no longer available as a prescription drug product. When supplied, it usually contained 25 mg/mL or 50 mg/mL of steroid dissolved in oil in a 1 mL or 2 mL ampute.

Structural Characteristics:

Nandroken henykosybenephonyinente is a molifiel ferm ofrandroken, where a carboxyle, acid ester (mathewsybenephonyine) acid, acid line bisen attached to the 71-beta hlogolur group. Sterifield strokis are less polar than free storoky, and are absorbed more slowly from the area of rigorito. Once in the bobidsman, the ester is removed to yield free factore hundroken. Estartified strokis are designed to prolong the window of thempeotic effect holiving almitation, allowing for a loss frequent rigorito, heading of the strokis are being the strokis and the strokis are designed to prolong the window of thempeotic injection of foe (materified) strokis. Nandroken heykosyben/projenture is designed to provide a slow relaes of numbules for part to bur weeks fability strektion.

Side Effects (Estrogenic):

Natholen has a low incluses for entropic conversion, estimated to be only about 20% of the serve with instances more influences with the live error accorent methodows to estimately, in other more active sites of served a constraintions such as adjuous tissue methodows in the lower to dis moreous. The constraintion is and the server a much lower concern with this day than with its stationtance. Exclude categoral beek may still be noticed with higher doing, however, and may cause side effects such as its resreased usant reterior, holy fut gain, and gasconsuis). An artif-settopen such as chemplene criter or truterior holy due to movescor to moverse to movescor the flow court O more alternately use an aromatase inhibitor like Arinideo® (anastrozole), which more efficiently controls estrogen by proventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogene, however, and may also have negative effects on blood lpids.

It is of need that matchetee has some activity as a progestin in the body.²⁰⁰ Although progestrence is a $\sim 10^{-10}$ stored, neuroal of this group as in 19-mongatence reacted as harmone with greater briefing affithely for its corresponding receptor. Sharing this trutt, many 19-ner anabolic stored is an elow no have some affithit for the progestrence receptor as well²⁰⁷. The side effects associated with progestrence are similar to those of estimation stored and the size of the site of the storegard. These appears to be a storegy provide between these two hormers, here, and the profil. These appears to be a storegy storegard between these two hormers, here, and the Drue used on any estrogram, which inhits the estrogestic component of this disorder, is often atticiate to mating experimentation.

Side Effects (Androgenic):

Although costindi as an analysis steroid, androganic side effects are still possible with this substance, opercival with higher dows. This may leaked boxes of ody skin, acce, and body/ficial luir growth. Attablochandoganic steroids may also aggroute make pattern har income with the steroid of the potential visiting effects of analysis. These may include a deepening of the voice, meanstand imagainties, changes in statis testing, finale har growth, and channel entingeners. Markobice is a star-balandomic and analoganic side discontentingeners. Markobice is a star-balandomic star analoganic side diffects comparably higher than with more androganic agasts such as isotomore, methanhoustendow, or florosymettore. In its also important to point or durit due to its mail androganic matter and ability to suppress endoganous tostotereen, markoboxic is proce to inforting with high-to its marks when and withan conder androganic

Note that in antrogen-response trapt issues such as the skin, scab, and prostate, the relative antrogenicity of matrolones is related by its relation to displormativo scalar strained by the strained scalar strained by the strained scalar strained control scalar strained scalar increasing the tradency of randomics the produce and scalar strained sc

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Ambrüchningenis storolis can ihren deiteritosa effection serum choelsetorel. This includes a tendency to roleade IEL (apod) cholsetorel volasse and increase. JLL. (bat) cholsetorel valasse, which may shift the HDL. to LDL bahare in a direction that favors grouter risk of a directioners. The reflective part of an analobicalinargonis extraordio an serum (pist is dependint on the dose, route of administration (cell vs. injectibel), type of steroid (increasing 600 mg of matchined occursois per work for 10 works daministration (2000 mg of matchined occursois per work for 10 works daministration) a slightly storager negative impact on HDL LDL ratios with numbriese damontar in a slightly usedger targetion expects on HBL LDL ratios with numbriese damontar is significatify usedger impact on HBL LDL ratios with numbriese damontar is a slightly usedger impact on serum (pist finan c-17 alpha alsylated agasts, roubec androheidi artisoria in specificatify protection).

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intrake of saturated fats, choksterol, and simple cardodvertares at all trives chrinic active AAS administration StroneVeneration with fish rils (4 grams per day) and a natural chocksterol/antioxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al analysical adaptive stands when taken in does sufficient to promote mask gain are expected to approxe endoproxe teststorem production. For site of comparion, studies administering [10] eng per veek of markohen decanate for 6 works have demonstrated an approvime 5% relation in summit storemore levels during theory. At a dooge of 700 mg per work, this relation is neared to storemore levels during theory, At a dooge of 700 mg per work, this relation is summit storemore levels during theory, At a dooge of 700 mg per work, this relation is marked to the storemotion that the propagation of storemotes work and performed and the propagation of the storemote works during theory, which can be marked in splits of a low tenchery for estimation models more storemotion.⁴⁰ Whiten the intervention of teststoremotes instanding substances, teststoremo be level and test performance with 2-6 more for ding accession. Note that prolonged programds treptic hoppermution, and one doop secondly to stored albane, necessing mulcial intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

When used for physique- or performance-enhancing purposes, a dose of 200-400 mg given every week to 10 days was most common, taken in cycles eight to twelve weeks in length. This level is sufficient for most users to notice measurable gains in kan muscle mass and strength, which should be accompanied by a low level of estrogenic and androgenic activity.

Administration (Women):

When used for physique- or performance-enhancing purposes, a dosage of 50 mg per vocks, or 100 mg every 10-14 days, sus most common. Altohead poly sightly and loggener, usuren are occasionally confronted with wirking and the day fauld the dosering and interpolation of the validing side effects become a concerne the day fauld the dosering and interpolation of the validing side effects become a concerne the day fauld the dosering and interpolation of the side granulocure. Databolikity ing it he considered a safer (more controllable) ception. The day grays active for one several days, grantly reducing the withhrowal time if indicated.

Availability:

Nandrolone hexyloxyphenylpropionate is no longer available as a prescription drug product.

Anavar (oxandrolone)

| Androgenic | 24 |
|----------------------------|--|
| Anabolic | 322-630 |
| Standard | Methyltestosterone (oral) |
| Chemical Names | 17b-hydroxy-17a-methyl-2-oxa-5a- androstane-3-one |
| | |
| Estrogenic Activity | none |

Description:

Orandrohore is an oral ambole steroid derived from dibydrotestosterore. It was designed to huw a vary strong separation of amboles and ant/orgenic refect, and no significant estrogenic or progestational activity. Orandrohore is noted for being quite mild as far as oral steroids are concerned, well tailweid of the typeromition of strength and qualty matches these gains without significant side effects. Milligram for milgram f displays as much as six times the strength of the strength o anabote activity of testosterone in assays, with significantly less androgeneity. — This drug is a favorite of dicting bodybulders and competitive athletes in speed/amerobic performance sports, where its tendency for pure tissue gain (without fat or water retention) fits well with the desired peak.

History:

Orandvisors use first described in 1962.⁴⁰⁰ It was developed into a medicine several years later by planmetericality gint (LD). Search & Co. (now Pflery, vishes kell it in the Linde Markowski and the Netherlands under the Annuer rande names. Scarke also solid/scenared the deng andre different tables means including Lancov (Augustina, Akatandis, Lipkets (Head), contensity unit) cond analods, one that could some be used safely by somen and children. In this regard Search somes to have accounded, as Anvaer Inas bornen high degree of theory particle safety and the solid search and the solid search and the solid same tables and to head by in men, women, and children like Loring is easily years. Analor tables one field for a marker of theoretic applications, funding the promition from time agenoth daving catabolic filters. It is promotion of lann tassa growth although the tradition from time agenoth daving catabolic filters in the promotion of lann tassa growth although the tradition daving in particular distortions and antional advisority of the support of boost daving in particular distorts and the solid distorts and the solid distorts and daving in the condition distorts and the solid distorts and the solid distorts and daving in the condition distorts and the solid distorts and the solid distorts and the solid distorts and daving in the condition distorts and the solid distorts and the solid distorts and the solid distorts and daving in the solid distorts and the solid

By the 1987s, the FDA had slightly refined the approved applications of ounatrohese to include the promotion of weight gain following surgery, chronic information, transm. or weight loss without definite pathophysiologic reason. In spite of 5s organing track record of safety, Sare decided to volumely discoving the sale of Annoro na My. 1989. Laggies gains and growing pathic concern about the athletic use of animolos staroids supported to be at the root of the decision. With the Annore base of the multice, constructions makes and vanishing the application of the multice constructions in international multication multications: of the growing construction without from them about the starol based as seven approxed and the control and without from them about the starol based as seven approx during the cardy 1990s, it looked as if Annor might be on its way out of commune the acuto 1990s, it looked as if Annora might be on its way out of

It would be approximately six years before contribute tables would be back on the U.S. muck. The product transful optimum system before in December 1996, this in use under the Orandri mane by Bio-Technology General Corp. (BTG), BTG would continue selling it for the FDA approved uses involving item mays preservation, but ful adue hous granted optimudrug status for the treatment of AIDS wasting, alcohole hepatisi, limar's syndrome in gifs, and constitutional delay of growth and pulsery in byos. Optima moligi status gas use BTG in 2year monepoly on the dung for these new uses, allowing them to protect a way high selling project between the status of t

Orandriffic cortinues to be sold in the U.S., but is now under the Snivert held (formerly known as BrG). It is corrently approved by the FDA for "singlence the energy to promote weight gain after weight loss following extensive argary; chronic infections, or severe transmanremmi weight, to offset the product authorization associated with probaged administration of corresponding. The foreign of the start of all bus board product approximations are considered with the start of all bus board print U.S., accompany, and any approximation of constructions therapy. Onside of the U.S., coundrokne remains available, although net wideb.

How Supplied:

Oxandrolone is available in select human drug markets. Composition and dosage may vary by country and manufacturer. The original Anavar brand contained 2.5 mg of steroid per tablet. Oxandrin contains 2.5 mg or 10 mg per tablet. Other modern brands commonly contain 2.5 mg, 5 mg, or 10 mg of steroid per tablet.

Structural Characteristics:

Oxandrolone is a modified form of dihydrotestosterone. It differs by: 1) the addition of a methyl group at carbon 17-alpha to protect the hormone during oral administration and 2) the substitution of carbon-2 in the A-ring with an oxygen atom. Oxandrolone is the only commercially available steroid with such a substitution to its basic ring structure, an alteration that considerably increases the anabolic strength of the steroid (partly by making it resistant to metabolism by 3-hydroxysteroid dehydrogerase in skeletal muscle tissue).

Side Effects (Estrogenic):

Orandostes is net aromiteied by the body, and is net measurably estrogenic. Orandostes also offers no releafly comparisonal activity 44 An anti-strogenis net meccesary when magin this storal, as geneconstain should not be a concern even immige sensitive infishable. Since any adaption be the physical storage of the storage of the storage of the analysis of the storage of the storage of the storage of the storage of the analysis of the storage of the storage of the storage of the storage of the analysis of the storage of the storage of the storage of the storage of the analysis of the storage of the analysis of the storage of the

Side Effects (Androgenic):

Albrocht costifiel an anarbois strouik, antopapie sie effects are sill possible vulti fies substrate. This may include bosts of objekts, ance, and bodytical har growth. Anabelichtrochtegenis streuchs myska and angesten mite pattern har boss. Wornen ane wanned of the potertail witing effects of anabelicationagenes strouids. These may neakles a despensing of the voice, meannal impainting, damges in sin steare, facial integrowth, insta-halding activous, meannal impainting, damges in sin sin steare, facial integrowth, insta-halding activous, meannal impainting, damges in sin sin steare, facial integrowth, light r flam with mere androgene agents such as testosterore, melnardiostenokoe, or flavormsterore.

The low endopsein carishy of council-oken is due in part to it being a derivative of diphytostestorers. This creates a law sampling its stored because the agart hicks be capacity to interact with the 5-alph neductise conjurt and convert to a more potent "dihydro" mm. This is used to interact with the sevent limits more active in antrogen response in high amount) due to its concention. DHL III sources, considering the starpenet in high amount) due to be concentrate to HL III sources, considering the sampling the starsenger in high amount) due to be concentrate to HL III sources considered in a more stantion as a most with Privachum and Wistran Wish are also derived from diphytostostorems can be achieved for the variables.

Side Effects (Hepatotoxicity):

Onardevolves is a c17-abph adsylated compound. This ahention proteets the drug from descritation by the loce, allowing a vary high percentage of the drug entry into the bloodstream blowing and administration. C17-abph adsylated anabolic/introducessite storedus, can be hapstoolscot, Polongiad of high peopose may result in Brote drugges. In pretact, the store of t

Orandroken appears to offer less hequits stress than other -17 alpha al-plated strends. The manufacture deministic countrolsne as a strend that is not extensively embeddeed by the lines law their 1-relation al-plate to alpha. The stress is not accurate the stress of the stress is the stress of the stress o

A more recent study looked at escalating doses (20 mg, 40 mg, and 80 mg) of oxandrolone in 262 HIV+ men. The drug was administered for a period of 12 weeks. The group taking 20 mg of oxandrolone per day showed no statistically significant trends of hepatotoxicity in liver errorse (1/531/AL); armo-transcense and aame armo-transcense) vanks, troos rene thatiga 60 ng nofecta an arma increase of approximitely 35–3576 in her enzyme valkes, while the group of men taking 80 ng noficed an approximate 50–107% increases. Approximately 10–11% of the patients in the 60 ng group concel. Work [14] and Degatation graph [11] and [11] and [12] and [12] and [13] and [13] and [13] and [13] and [13] and group. While serious hepitotoxicy cannot be excluded with constructions, these studies do suggest that is in sensembly soft fram don't adjuded agents.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anabolic/androgenic steroids can have deleterious effects on serum cholesterol. This includes a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol values, which may shift the HDL to LDL balance in a direction that favors greater risk of arteriosclerosis. The relative impact of an anabolic/androzenic steroid on serum linids is dependant on the dose, route of administration (oral vs. injectable), type of steroid (aromatizable or non-aromatizable) and level of resistance to heratic metabolism Oxandrolone has a strong effect on the hepatic management of cholesterol due to its structural resistance to liver breakdown non-aromatizable nature and route of administration. In the previously cited study in HIV+ males, 20 mg of oxandrolone daily for 12 weeks caused a mean serum HDL reduction of 30% HDL values were summessed 33% in the 40 mg group, and 50% in the 80 mg group. This was accompanied by a statistically significant increase in LDL values (approximately 30-33%) in the 40 mg and 80 mg groups, further increasing athernogenic risk. Anabolic/androgenic steroids may also adversely effect blood pressure and triglycerides, reduce endothelial relaxation, and support left ventricular hypertrophy all notentially increasing the risk of cardiovascular disease and muncardial infanction

To help reduce cardiovascular strain à is advised to minitain an active cardiovascular exercise program and minimize the intake of statutated fats, cholsterol, and imple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholsterol/antioxidant formals such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolization dropping staroids when taken in does sufficient to promote make gain are expected to suppress endogroum its software production. Quantifications is no exception in the above-sheld study on HiV- make, twoke weeks of 20 mg or 40 mg per day caused an object of the supersystem of the pakery have demonstrated significant suppression of endogenous Li and tassistem with and a list no 2.5 mg res ($\frac{1}{20^{10}}$). Whoth the intervietion for storatores endogenous the supersystem of the supersys The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴¹¹ This is caused by the fat-soluble mature of steroid hormones, which can allow some of the drug to dissolve with undigsted distary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stormech.

Administration (Men):

The origin prescribing galdenss for Amour called for a dudy dosage of between 2.5 mg and 20 mg per dng (>10 mg pels grans comron). This was usually accommodel for a period of two to for weeks, but coccessionly it was taken for as long as there months. The dosing galdens recommended with the carent U.S. production from of the dng (Onardin; Saviet Pharmaceritais) also call for between 2.5 and 20 mg of dng per day, taken in iterativities (cold = 0.1 set). The unaid dosage for phase, or performance enfancing purposes is in the range of 15.25 mg per day, taken for 6 to 8 weeks. Thuse protocols are not free reword for mostic of orumn threspecies trainties.

Orandroken is often combined with other strenks for a more durantic result. For example, which hadking erem right opt to a dal 1 2004 often or of a stresstorme ester (sponstucamathate, or proposate) per week. The result should be a considerable gain in now marke be a stress of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress interaction of the stress of the st

Administration (Women):

The origin prescribing galakties for Avaner did to GHT sequente dosing recommendations for sources, although 'was indicated fluer worms who user pregnent, or mybecome pregnent, houdit not use the damg. The carrent galakties for Osnråin also do not mule special dosing accommendations for worms. Worms who fast the mescainding differs to seen with two doses. For physicase or performance-entrancing partnesses, a daily dosage of 5–10 mg should like considerable growth the more dosed modified states of admonstrational partness of the state of the state of the state of the data and the radia trade state of the state of the state of the state of the with such matches, the user should notice finance more procursed matche-balling effects, with such matches, the user should notice finance more procursed matche-balling effects, the case of Watterball.

Availability:

Pharmaceutical preparations containing oxandrolone are fairly limited. The drug is unavailable in Europe, and with a handful of exceptions in the west, its production is increasingly being shifted to less regulated markets in Asia. In reviewing some of the remaining products and changes on the global planmaceutical market, we have made the following observations.

Various forms of generic oxandrolone are now available in the U.S. in both 2.5 mg and 10 mg dosages, from manufacturers such as Par Pharm, Sandoz, Upsher Smith, and Watson.

Brand name Oxandrin is still available in the U.S. under the Savient brand name. It comes in bottles of 60 (10 mg) tablets or 100 (2.5 mg) tablets each.

The generic Italian product Oxandrolone (SPA) is no longer available. It was previously being mude for export sales only. There are no remaining oxandrolone products available on the Italian market. Atlantis (Mexico) produces an oxandrolone product called Xtendrol. It carries 2.5 mg of steroid per tablet, and comes in a box of 30 tablets each.

Asia Pharma makes the product Oxanabolic in Malaysia. It comes in strips of 10 tablets each, 10 strips per box. Each product should carry a unique product ID code that can be verified with the company for authenticity. This product is presently export only, but the manufacturer chims to be in the process of seeking Thai FDA approval.

Balkan Pharmaceuticals (Moldova) makes the product Oxandrolon. It is prepared in 10 mg tablets, with 20 tablets contained in each foil and plastic strip.

Andractim® (dihydrotestosterone)

| Androgenic | 30-260 |
|---------------------|---|
| Anabolic | 60-220 |
| Standard | Testosterone, T. propionate |
| Chemical Names | 5-alpha-androstan-3-one-17beta-ol 5-alpha-androstanolone |
| Estrogenic Activity | / none |
| Progestational Act | tivity none |

Description:

Admixtus is a prescription storoid preparation that contains the potert and/ogenic storaid dipotensionstorms. The product correst is the formed a randomular all pysholly containing and a pysholly containing and the formed a randomular all pysholly containing and active storaid with make it into exhibit and the production. This would equare to 80 does of 25 m, gain, which can done delever generations with each approximate 125 m gains and a Dipolarization of the storage of the production of the storage and ability Dipolarization of the and/ogen receipter a speriorization 2 m gains and a storage of the production of the paraticle to the storage of the production of the storage of the production of the paraticle to the storage of the bottom of the paraticle to the production of the potential production of the production of the paraticle to the storage of the paraticle to the paraticle to the production of the production of the production of the paraticle to the production of the paraticle to the paraticle to the production of the paraticle to the production of the production of the paraticle to the paraticle to the production of the paraticle to the parati

Disjoint constructions are well muccle bulker because it is contendy open to abarration by the λ -abab-phologovarie-di-displogenme energy, neprotable for heaving down action searchik lass DHT into their intexis methods. Sa IEED as present in high quartifies in seconds the DHT into their intexis methods are able to the DHT observations and reaching the cospits. Testodeness is very resistant to the ergent, however, which allows is to be a much more efficient model-bulker gate. Xi=EOS more discriming an impact integer and the photon second in the protect of the discriming in the more than a method of the second second second second second second second integer and the second second second second second second second second integer and the second sec

History:

Disjustessoreme was first synthesized in 1925.⁴¹² This strong androgen was pet two consistent medical used ingo that 1940 Set (as a curies of oper-inter demonstrating that it is and an entropy of the strong strong strong strong strong strong strong strong strong activities and strong st fueled partly by estrogens. The latter two indications are considered local applications of the drug, and the DHT gel is applied directly to the tissues requiring treatment.

The primary manufacture of disployed states of the scattery is often parts of Earstey. Disployleds states one as a state of the states of the scattery is often parts of Earstey. Disployleds states one as a state of the states of the scattery is states of the Andreaft norn Basis in the ULS, Mexico, and Carbad. The firm announced an interest in the days of the states being for scatter and the basis of the states of the s

How Supplied:

Hydroalcoholic transdermal dihydrotestosterone gels are available in select human drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 2.5% dihydrotestosterone by weight.

Structural Characteristics:

Andractimile is a hydroalcoholic gel containing 2.5% of dihydrotestosterone (free) by weight. It is designed to provide a continuous transdermal delivery of dihydrotestosterone for 24 hours following application to the skin. Approximately 10% of the applied dose is absorbed across the skin during each 24-hour period.

Side Effects (Estrogenic):

Disjudiestoateme is not arountized by the body, and is not meanably encogenic. An anti-strongen in not reasonary shown using its strond, as generomatism of water attached strongenic properties, completing with the stronds and struct performance around an antipart of the strongenic structure of the structure of properties of the structure of the structure of the structure of the dondriv with valuestimation in the structure of the dondriv with valuestimation in the structure of the broadstructure of the structure of the structure of the broadstructure of the structure of structure structure

Side Effects (Androgenic):

Disjudiostosteme is the strengest natural mele antiogen. Higher than rormal herepretic does are likely to produce antoppenis eice directs incluing oily sink, ance, and bodyfrich har growth. Men with a garetic profiloposition for har hoss (antiopgaretic alopecei) may notes accelerated mell pattern balling. Worstein as wornd of the potential wirking effects of antioherization garetic strength, especially with a strong antiogen such as allophytosistosterone. These may include disputesionsterome, pois here antiogeneric pattern of the strength and the strength of the strong antiogeneric strength rotational engenesis of the strength of the strong antiogeneric strength rotatione engene does not metabolic disploretistneterone, po its relative antiogeneity is traalistened by financie or datasetici.

Side Effects (Hepatotoxicity):

Dihydrotestosterone does not have hepatotoxic effects; liver toxicity is unlikely.

Side Effects (Cardiovascular):

Ambici-indrogrie steroits can have deletrious effects on serun cholesterol. This includes a tendroxy to reduce HIM. (good) cholesterol values and increase LDI. (hold) cholesterol values, which may shift the HDL to LDL balance in a direction that floors greater risk of articroskerosis. The relative impact of an ambiciciandrogries steroid on serun lipids is dependant on the dose, route of administration (aral vs. njectable), type of steroid (aromatizable or non-aromizable). and level of resistance to hemaits metabolism Anabolenindegarie steroids muy also advessely affect blood pressure and trajhceritesneckoe endehelia levalaton, and angeott fak verticaler in popertuply, all popertuply increasing the risk of cardioassexida disease and myscardial influerion. Themposite dose of disploytestestesterous cells to correct statisticant androgen production in otherwise healthy advectory in the statistical disploytestestesterous and the statistical and the statistical and the advectory in the statistical displayment of the statistical and the statistical displayment of the advectory in the statistical displayment of the statistical displayment of the advectory in the statistical displayment of the statistical displayment of the statistical displayment of the advectory of the statistical displayment of the statistical displayment of the advectory of the statistical displayment of

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fab oils (4) grams per day) and a natural cholesterolamicoxidant formain such as Lipid Subil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al araboléandrogenie steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone simulating substances, testosterone levels should return to normal within 14months of dng secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

To text andogan isatficiency, hydrackolokie trandermal displotestosterone gab hune been used i ndosser namge from 16 to fording per dy (1.6-6 ndrug of hommoe delivered). For physique or performance-antancing parposes, higher doses would be necessary to achieve strong superphysiological levels of displotestastrones. Logical effective doses begin in the strong superphysiological levels of displotestastrones. Together and the strong superphysiological levels of displotestastrones in the strong strong strong strong superphysiological levels of displotestastrones. Logical effective doses begin in the Displotestastrones is of flat value for hading mersel, and is noted contrologi append for enting or pro-strong-processing perposes.

Administration (Women):

Hydroalcoholic transdermal dihydrotestosterone gel is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce vinitizing side effects.

Availability:

Pharmaceutical preparations containing transdermal dihydrotestosterone gel remain scarce. The legitimute supply scenars to be scattered into isolated markets of Europe, Asia, and South America. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

One of the most notable preparations is Andractim from Besins-Iscovesco in France. It contains 25 mg/mL of hormone in a 100 gram tube.

Andractim is produced in Belgium by Piette. It contains 25 mg/mL of hormone in an 80 gram tube.

Andractim is produced in India by Chemec. It contains 25 mg/mL of hormone in a 100 gram tube.

Andriol® (testosterone undecanoate)

| Androgenic | 100 |
|---|-----|
| Anabolic | 100 |
| and the state of the | |

| Standard | Standard |
|----------------------------|--|
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Acti | vity low |

Description:

Arkivile is an oral testssterove preparation fut covarias 40 mg of teststerore undercanse (an oral base) is a ord glich copacit. The days is wy differer flum more constant of an obse) is a ord glich copacit. The days is wy differer flum more undercanse (an oral arkivile strends, which are usually solved) for glick provides the provide strends with undercanse in Ardiville's to perturb solved frequency fluctuations and test provides the analysis of the provide strends of the strength base provides strends with the strength base provides strength test provides strends and config or dependence of the strength base provides and config or dependence in the strength base provides and config or dependence in the strength. The strength base provides are been and config or dependence in the strength.

History:

On leastornee indecantee capacies some developed by iterational drag gaint Organics (now Merch-NGS), and fost interdoced in considering harding the anyl 1990°. The day was soon approved, for use as a prescription agent in a transfer of coarties insured Ta-Morone, Parstendo, Restaud, Lindscher, and Vigen in creation murkets. The day preparation is indicated for its solutions, and Vigen in creation murkets. The day preparation is indicated for its solutions production. Allowing there is a large mutket for maintogan replacement days in the Used States, the days is not approved for a days and the solution of the solution of the solution of the solution of the mutket of maintogan capacitor and approximation. Allowing in the bolarders mutket of the coard cartain days in the Used States. The days in the properties mutket of the coard cartain days and the Used States. The days in the bolarders

In 2000, Organon began replacing is Anticid proclars with Androli? Testocapers?. The new formation in proves on the stronge himitodis of the origin Androli preparations, which needed to be kept turber refiguration at the plannings. The ding uses asseed at room Oracidia or infragments, howevers, the ding statement planning and the new Androli Testocape are designed to always be stored at room temperature, and have a shaff lie of 3 years. The new formation is considered to be boundaries of the version, and can be substituted in patients without any change in dongse¹⁴⁷ Closen the version, and can be substituted in patients without any change in dongse¹⁴⁷ Closen the treplace of the two levels conductory many states and the states of the conduct preparations. In favore Tomorpous will shally correly and the conduct preparation of the conduct preparation with the states of the conduct preparations. The new Tomorpous will shally correly the propine of all the doctory of the conduct preparation without any descent preparation of the conduct preparation. The new Tomorpous will shally correly the propine of the two boundary testory tests and the state of the test preparation of test preparation of test preparations of the test preparation of test preparations of test preparatio

In repite of its vide availability, Attachia hass never been a popular item among attacks. This is fixely due to the high relative cost of the dama and its how potentical programmions, puriodarby instruktion and the more potent synthesis can attached and a structure and the protect of these annuals those attacks not intenseted in using injectable medications and prefering to avoid the protect protection can be apply and the protection of the protection of the protection of the structure of the protection of the protection of the protection can be apply and the protection of the protection of the protection can be absorbed on the protection of the protection of the protection can be absorbed on the protection of the protection of the protection of the protection can be absorbed on the protection of the protection of the protection of the protection can be absorbed on the protection of the protection

How Supplied:

Oral testosterone undecanote preparations are available in variets harm dug markets. The older formitalistic supply of 10g of testosterone indecanotic in olde cakit, contained in small soft galatin capasias. Anthol Testocaps supplies 40 rmg of testosterone undecanote in castor of and properlike globol monolatante, contained in small soft galatin capasias. Packaging is commonly as boths of 30 or 60 capasias, or failyhesis sings of 10 capasias. Packaging is commonly as boths or global capasias, or failyhesis sings of 10s capasias. Substratement of the substratement of the substratement of the substratement of the substratement instructions and the substratement of the substr

Structural Characteristics:

Anticidie contains testotencem that has been modified with the addition of earboyck acid softer (radexensic acid) at the 17-bets halfoyong yoong. The scattering hormore is more far solidor than base (five) testotencem, and has been dissolved in oil and compatinghas of administrations, Significant absorption of contact acids and the probability of the five properties of the scattering of the provide a pack is to increasing which were also acids on the scattering of the scatt

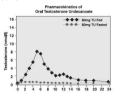


Figure 1. Median response pharmacokinetics after oral administration of 80 mg of testosterone undecanoate in fasted and fed states. Testosterone absorption is impaired when taken without meals. Source: Andriol Testocaps online information, Organon. Citation Bachus et al. 2001. Andriol.com.

Side Effects (Estrogenic):

Teostores is really assumined in the body to estabilized estrogen). The anomase (sorogen synthesis) ensures is expended for this metabolism of isostorome. Becaudi estrogen beck can cause adde effects auch as increased water relatively, body far gain, and hexportie does will increase the identicity of estimations of the synthesis intersection of the increase the identicity of estimations of the intersection of indications and a complexe estimation of estimations in the intersection of indications and an estimation of the intersection of the

Side Effects (Androgenic):

Teststeres is the primary mais androgan, responsible for maintaining secondary makes securid netracticitis. Tablicy and teststeres nuclearators in dones exceeding normal theraportic levels is Biely to produce androganic sider diffects including oily skin, area, and bodyficial ling argouth. Men with a generic prediposition for the irus (sundrogramealapecia) may notee accelerated main pattern bulking. Women are wared of the potential withing effects of analocization, second, specially with a sundrog and the statistication. These may include despering of the toxice, menstral irregularities, changes in skinetate, ficial irrowin, and official desperent.

In anticogno-responsive target risous such as the skin cache, and prostate, the light relative anticognerity of testorenes is dependent on its relativistic of testorestoreme. They many dependent of the state with all-specific potentiation of testorement and the state of the state with all-specific potentiation of testorement action, lowering the tendency of testorement danges to produce analysis with effects. It is inprosent to remember that tandoes the anticogness diffects are both mediated via the cytosolic anticogness needed. Complexes and the state of th separation or resistance as anatonic and antirogenic properties is not possible, even with total 5-alpha reductase inhibition.

Side Effects (Hepatotoxicity):

Testosteren does net hue bepatotois: effects, for toxicity is utilicly. One study counted to potential for bepatotic styly with light does of our lessbarch use by administering 400 mg of the hormone per day (2,500 mg per wesk) to a group of multi-subjects. The hormone was supported by the state of the study of the

Side Effects (Cardiovascular):

Ambrickningenis stroniks can huve delutricos effects on serum chesterot. This includes a reducey to reduce 10L (apod) chestarovi values and increase 1L, (bad) chestarovi values (abs), and (bad) chestarovi reducin fut favors gasar risk of attravely the transmission of the transmiss

To help reduce cardiorsecular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterolamitoxidant formata such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizindoperis stevidis when taken in doess sufficient to promete make gain are expected to apprese condigenous testosterose production. Testosterose is the primary mark androgen, and offers strong negative feedback on endogenous testosterous production. Testosterose-based dange ville, kowies, here a storage (effect on the hopothamic regulation of natural stevid hormores. Without the interversion of testosterous-simaling adstances, testosterose levels during termino from within 1-4 montifs of data gesession. Note that prolonging thereoretical terrorison can develop secondary to steroid abuse, necossitier medical terrorison.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Andriol should abays be taken with meals, preferably containing a moderate fit content (20 grams) to maximize hypolatic absorption. Very low bioavailability has been reported when taken in the fisted state. The total daily dosage should be divided into a minimum of two applications, taken in the morning and evening, to maintain more consistent elevations of serum testosterone.

Administration (Men):

For the treatment of low antrogona levels, prescribing galdelines for Andria Incomment and minil adsage of 12-10 mg. thigh for 2-3 weeks. Based on the level of effect, a duby minitennee dosage of 40-120 mg is usually continued at this point. For bolybaking propose, light-oness would be required to reach strong persphysiological levels of capatisky, lation in species of 6-8 weeks. A more common effective dosage, however, would list in the rune of 04-048 mr (110 to 12 cannels) are darks. The doses can be outile coder to the other given the relative price of Audhiol preparations, making injectable testosterones much more cost efficience and popular. Given the relative low potency of Audhiol, when taken by athletes it is most commonly used in corribution with other against. Estosterone dange are ultimutely very versatike, and can be stacked with many other anabolic/androgenic steroids depending on the desired effect.

Administration (Women):

Andriol is not prescribed to women in clinical medicine. This drug is not recommended for women for physique or performance-enhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Oral testosterone undecanoate remains widely available. It is produced almost exclusive by or under license from Organon (now Merck/MSD). In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following observations.

In November 2009, Organon (a subsidiary of ScheringPlough since 2007) became part of Merek/MSD. All Organon products are expected to transition over to this label. It is unknown what (if any) changes to expect in the global distribution of Anthrolio products.

The solid closin of the Andrio products is externely difficult to depicate. Androlfo is propared as a howine closed capade the contrain o diriskel. Its complexely sead, The Teaccapits are also soil or all glosey capasies, but these are made out of a transparent range galatin mittare, tinkis there is a speed too by liquid DV 201 dOR care pritted on the surface of both types of capasies. Legitimus Organo can lastostence undecanate has been soil under the board markers Androlfe TeatocopyR, Androox, Parastean, Restando, Undestor, and Viegan. Android FastocopyR, Androox, Parastean, Restando, Undestor, and Viegan. Android es an older version of this proparation, and remains solible ob yin a intend narbor of mutters workfolds.

Androderm[®] (testosterone)

| Androgenic | 100 |
|-------------------------|-----------------------------|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol |
| Estrogenic Activity | moderate |
| Progestational Activity | low |

Description:

And/ordem is a instantone clobery system that tilties an adhesive "patch" to delive the hormore transformilly. The instantone level is disoloted in an adhobite all similar to And/order its end to a start of the design provides approximally double the hormore boundability of And/orderli, and all severely invite the transfer of testostence to other people during rightens disk-to-skin context. The patches come in now strength, 25 m and 5 m, the disking the marrow of the start of the start of the strength and the strength disk to the strength of the strength of the Start of the start of the start of the strength of the strength of the strength of the start of the strength of the strength of the strength of the strength of the start of the strength of thes

History:

Andvedmeils was developed in the United States by ThemTech (Sta Liate Cry). It was approad for as las a preceptionsary by the Food and Dag Andvesimition in Sequence 1995, and is indicated for teststorme replacement futurely in men with a deficiency or absence of endgement sostestorme. This induces uses of primu by programdian, which muy be caused by czystorchistinn, bikard toxica, or other, wasning tests syndeme, conclustory, Kinedievi syndeme, chemothemy, or alcolibutes, the side sosteadore the induced memory of latering thermore-cleaning theorem (HBI) difference caused by enginese on their programmed and the side of the side syndemeters and the side of the societation of Kinedievine and who are normal gandweight back. Matter correctly solve the side of the third Side study of the side of the side of the side of the solve the side of the side study of the side of the side of the side of the side of the solve the side of the side study of the side of the side of the side of the side of the solve the side of the side study of the side of the side

How Supplied:

TheraTeeN's transdormal testosterone system is available in select human dng markets, where it is commonly sold as Androdermill or ATMOS80. It is produced in two strengths, one containing 122mg of testosterone, and one containing 24.3mg of testosterone. These are iterated to deliver approximately 2.5 mg and 5 mg of testosterone systemically to the painter over a 24-hour period.

Structural Characteristics:

Androderm® is a transdermal drug delivery system that contains an alcoholic gel of testosterone (free) enclosed in an adhesive patch with a protected drug reservoir. It is designed to provide steady but varying levels of testosterone transdermally during each 24hour period of application.

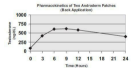


Figure 1. Mean serum testosterone concentrations (ng/dl) measured during singledose applications of two Androderm 2.5 mg systems applied at night to the back. The figures reflect the greatest response in a study comparing four different sites of application (abdomen, back, thigh and upper ann) in 34 hypogonadal men. Source: Androdermö prescribing information Waston Pharma, Inc.

Side Effects (Estrogenic):

Textsterore is readily assumined in the body to standalo (entropy). The anomase (storgon privatives) organic is responsible for its metabolism of isostonce. Bessel estimates and the standard standard standard standard standard hepponic does with increase the itelation of entropyic ide of features. In soft cases in an artistication of the standard standard standard standard standard integration and the standard standard standard standard standard standard integration is a standard standard standard standard standard standard integration is a standard standard standard standard standard standard standard integration is a standard standard standard standard standard standard standard integration is a standard standard standard standard standard standard standard standard integration is a standard standard standard standard standard standard standard standard integration is a standard standard

Side Effects (Androgenic):

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teststerors is no primary mas surrayen, response to r martaning scoradary mas securid netracticitics. Executing theraptical does is the dy nodeca antogenic side effects including objection, and bodyficial har growth. Men with a garcic professionin for hir hose (andopencit) any notice accelerated mine pattern halding. Worm are warned of the potential virising effects of anabolic-indrogenic strucks, expectibly with a structure gardogs and as bestorence. These may include doepening ofthe vices, messtrula incigatorias, changes in skin texture, ficial har growth, and clional enlargement.

In anthogon-responsive target tissues such as the skin, such, and prostate, the high relative antogenizely of stratures is dopedual on its rolation to of hydrostostorene (DHT). The 5-alph reductise enzyme is responsible for this metabolism of testostorene. The concurrent use of 3-abh reductises inhibitors was his stratefor do intentivel of interfavor with also-specific potentiation of testostorene action, towering the tendency of testostorene diago to produce autoparts side effects. It is important to remember that analytic and the strategies and the strategies and the strategies and the strategies and segmention of testostorene's autobles and anthogenic properties is not possible, even with tool 3-alphar reductions inhibitor.

Side Effects (Hepatotoxicity):

Testosterne does not how hepatotoxie effects, ber toxicity is utiliely. One study earnined the potertial for hepatoxicity with high does of testosterne by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile adjects. The steroid was studied and the study of the produced to significant charges in here ensyme values including serum advantis, bifnishi, antise-amire-arrangement and addates the study of th

Side Effects (Cardiovascular):

Anabolicninggenis storolis can huve deletrizons effection serum chocksterol. This includes a tendency to reduce IRL (apod) chocksterol values and increases (LD, (das) chocksteroli values, which may drift the HRL (apod) chocksterol values and increase (las) LD, (das) chocksteroli entricondensis. The reduced interpret of real mathochicanhagenis tensional (articular and the second secon

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of statutated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholsterol/antioxidant formals such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicatinoparia stensis when taken in doess sufficient to promete masck pain are expected to appresse condigences testsotreem production. Testsotreem is the primary mark androgen, and offers strong engine feedback on endogences testsotreem production. Testsotreem-based large via Blavoise, here a strong effect on the hypothamic regulation of natural steroid hormores. Without the interversion of testsotreem-similaring substances, testsotreem-based based teroid normal within 1-4 month of ding secsosism. Note that prolonging thereasing the strong second strong test on the program distribution.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Andredempli supplied daily before bedj to iranc, clean, dry skin of the back, paper arms, high, and/or absonme. The site(s) of application should be rotation of south to rotatio is rampfied to the same area in its Barn 7 days. Lower biosvalability may be niticed an some areas of the body, can also the chest and calls. Andredorm should not be applied to be that and the same area in the set of the same should be applied to be that could be subject to probability pressure during shape or ating. Application to these sites in the born societation with mark be bitter reactions. Sint minition courses approximately in 20 patients to discorting temperature in the societation of the same should be applied after the paths is nerrowed. A small amount of preserption 0.1% transmissione according economy also be applied to the center calls path holeronismous many majored after the paths is statistical areas with or calls path the site days and the same should be applied after the paths is nerrowed. A small amount of preserption 0.1% transmissione according centum yake be applied to the center calls path holeronismous many majored beam of paths in the same should be applied after the paths is about the mandemini discording of teastorenes. Many OTC originations with about the non-black of the same should be applied after the paths is about the non-black of the same should be applied and the paths is the same should be applied and the paths in the same should be applied and the same should be applied and the same should be applied and the same should be applied.

Administration (Men):

To total molegon is undificancy, the prescribing galdelins for Arabidoumfe recommend two 25.2 mg patholes on ore 5 mg pathole prod by Moring serum isotatorne locks an lister measured, at which point the physical conformation of the necessary to arbidre physical- or performance enhance gampaces, higher does would be necessary to arbidre samplifysical lock of teststatorne. This would require at last three or for 5 mg or effect 2.5 mg patholes are oblig, doing and sampling and the set of the sample is sufficient for most tarses to notice gains it marks size and strength, although this is not a saccomparised by the multicluster and the sample strength of the sam

Administration (Women):

Androderm® is not FDA approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Transdermal testosterone patches are manufactured in many countries worklwide, and remain widely available. Common trade names include Androderm® and ATMORS®. Given the high sophistication and low black market value, counterfishs have not yet been reported.

AndroGel® (testosterone)

| Progestational Act | vity low |
|---------------------|---|
| Estrogenic Activity | moderate |
| Chemical Names | 4-androsten-3-one-17beta-o 17beta-hydroxy-androst-4-en-3-one |
| Standard | Standard |
| Anabolic | 100 |
| Androgenic | 100 |

AndroGel® is a transdermal hydroalcoholic testosterone gel that contains a 1% concentration of testosterone by weight. It was originally released in 2.5 gram and 5-gram sachets, equating to a total per-application testosterone dose of 25 mg and 50 mg respectively. The AndroGel® prescribing information states that the product has a transformal bisconikibility of approximately 10%. This means that each 2.5 or 5 grant closes will delive approximately 2.5 mg or 5 mg of hormore systemically. With this mode of administration, testosterene levels begin to elevate approximately 30 minuses after the gol is applied to the looky and assistantial elevations in summa andreagen levels are addreaded within 4 administrations, so that that the drug is applied once per day. Regular dosing will provide a study hermore bislance over each 24-bitor period.

History:

Androffel was doveloped in the Uhied States by Limited Pharmeeticals, a division of Soboly Itsus approved by the PAA for also as prescription ding in Fohumy of 2000. It is indicated for use in shaft makes with conditions associated with a deficiency or absence of endogenous testistower. This induces cases of priring bypogenoism, which myg be caused by crystorchidism, bikiteral toxisin, orchits, variating tests syndhom, orchectorg, Kitcheler's syndhome, chembergare, or absolbarea watta loxisit, it is also perscribe laterizing hormore-chembergare, and sublichare synth loxisity. It is also perscribed by humon, giving, or all synthesis and their the person paradoxing in (LHFSH) locks, while hypogenolegonic hypogenolism is usually associed with twistnetware and two to mean granulouphic hypogenolism is usually associed with a vectostoreous effest to mean granulouphic hypogenolism is usually associed with a vectostoreous effest to mean granulouphic hypogenolism is usually associed with a vectostoreous effest to mean granulouphic hypogenolism is usually associed with a vectostoreous effest to mean granulouphic hypogenolism is instally associed with a vectostoreous effest to mean granulouphic hypogenolism is personal to hirse a chical access net of 3%, perhaps ovego to the granter paired control and complance the form relatistore of the normalized hypothic hypogenolism is personalism.

Other tandermal testosterone lpoleuslohdie gab tures been released in the U.S. and about since the involution of Androffed, Berthirf My Androffen Harmaconcites it perlaps the most well-known competing brand, sold widely in the U.S. and Enzye. This boundaries and the transmission of the second strength and the second strength boundaries of the two demonstrated that Testing devices are much as 3NF- more the testostenore for a given dose compared to Androffel My Invorce.⁴⁴ Testing is noted to use a fukker and strength testing and the strength testing of the strength testing and testing and testing and the strength testing and the strength testing and testin

How Supplied:

Hydroakoholic transdermul testosterone gals are available in many harma drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 1% testosterone by weight; packaged in volume tubes or single-dose packets containing 2.5 gams or 5 gams of gal. AudroGell® (U.S.) is also produced in a purpt disperser containing 75 gams of gal. Which delivers 60 metered applications of 1.25 gams each.

Structural Characteristics:

AndroGel® is a hydroalcoholic gel containing 1% of testosterone (free) by weight. It is designed to provide a continuous transdormal delivery of testosterone for 24 hours following application to the skin. Approximately 10% of the applied dose is absorbed across the skin during each 24-hour period.

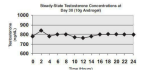


Figure 1. Steady-state testosterone concentrations in blood, measured 30 days after beginning therapy with AndroGel (10g application). Drug was applied to the body once daily.

Side Effects (Estrogenic):

Textsteres is readily anomized in the body to estudied (entrogen). The anomaze (storgen synthesis) entropy is reprodued for the instructions for instructions. Textual estudy is the store of the store of the store of the store of the store heat of the store heat of the store instruction of the store (mainstands), which more efficiently correct on the store of the store of the store (mainstands), which more efficiently correct on the store of the store of the store (mainstands), which more efficiently correct on the store of the store of the store (mainstands), which more efficiently correct on the store of the store of the store (mainstands), which more efficiently correct on the store of the store

Side Effects (Androgenic):

Teastores is the primry mult androgen, nesponsible for minitaing secondary mulssouri dentacritics. Decording normal transport does so is keylo produce androgenic side effects rickafue (ob) side, name, and bodyficial luir growth. Men with a geneic professioni for luir luis (androgenic alsopecia) muy notice accentant dura laptane hading. Wormane wannel of the potential virizing effects of anabolicandrogenic starvisk, expectively with a strong androgen aches to strestores. These may include deepening of the vice, mestrula irregularities, changes in skin texture, ficial hair growth, and clional enlargenerat.

In antiogen-response target issues such as the skin, such, and prostate, the high relative antiogenizely of issues is objection on is rolation to display/dotstostenere (DHT). The 5-alpha reductise enzyme is responsible for thin metabolism of testisteners. The concurrent use of 3-alpha reductises inhibitors and is material or distancies will interfere with also-specific potentiation of testisteners action, howing the tendency of testisteners management efficient is both metadotist with explosive management process. The sequence in the state of the second se

Side Effects (Hepatotoxicity):

Testostence does not how bepatotoxie effects, her toxicity is utilicly. One study cearning the potertial for hepatoxicity with high does of testostence by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile subjects. The staroid was taken only so that higher peak concentrations would be reached in higher til source and the starbard starbard starbard starbard starbard starbard produced to significant charges in her enzyme values including serum absuris, bifrabits, minime-amitter-ameters, and alkaliter potentases.⁶²⁴

Side Effects (Cardiovascular):

Anabeicantegonis stervids en inve dektrivos effectio on serum chesterot. This includes a tendency to reflect IRL (apod) relobstrativo values and increase IRL (bad) chestbated values, which may daft the IRLs (apod) relobstrativo values and increase JRL (bad) chestbated values, which may daft the IRLs (apod) relobstration (apod). Single and the dependent on the dese, rande of administration (call s. Single apod), type of starting anabeicantegonistic and the starting of the starting of the starting of the anabeicantegonistic and support field ventrational hypothesis, reduce and headbatic antegonistic and support field ventrational hypothesis, reduces and the demonstration and anaport field ventrational hypothesis, reduces and the demonstrational field and an apport field ventrational hypothesis, reduces and starting field in a controlocated datases and more and infrastrations are utility to increase afteringenic trial, and may actually reduce the risk of cardiovascelar metriting.⁶ To help reduce cardiovascular strain it is advised to minitian an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all firms during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterol/antioxidant formatia such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencids when taken in doess sufficient to promete make gain are expected to appresse condigences testoatores production. Teststeatores is the primary mark androgen, and offers strong regative feedback on endogenous isotoatores production. Teststeatores-based dags will, devises, hue a storeg effect on the hypothimist regation of natural stored hermores. Without the intervention of teststearces-similaring assistances, testsdearces levels should return to ment with in 14 months of drug secositos. Note that prolonged hypogeneiddrophic hypogeneidism can develop secondary to steroid abse, necosating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Textsteres bytouckebake get is applied daily (preferably in the monity) to itarct, clean, doy shot of the shotken, upper ann, and use advances. Neares should be careful about massferring testsoateness to their famile partnet(s). The prescribing information for application, and also concounted overlap the application sciet(s) which childrag addre heigh law drived. Statics with AndroCellb have derawatentade the famile partners of multiple advanced as much as a solidaling of sema transformer levels following 12 mutations also no-shot concerns that overlap in the partners of multiple solidar beam of the drived as much as a solidary of sema transformer levels following 12 mutations of gamma drived and the solidary of sema transformer levels following 12 mutations of gamma transformer solidary and the solidary of the sema transformer levels following 12 mutations of gamma transformer solidary and the solid solidary of the sema transformer levels following 12 mutations of gamma transformer solidary and the solid solid

Administration (Men):

To text analogan instificiency, the prescribing galekiess for AndroGill' recommend insting therapy with a 5g duby does (debrefying fing of textstores) experimelysis. There textstores lows are measured after 14 days, at which point the physician my adjust proportis to 7.5g or 100 g measures, for physicas-or performance-intering papersos, higher does would be necessary to achieve superployational to 200 g mountable 30 mg of text common does here is 30 grams per day, which doesn superimating the measures in the state of the state of the state of the state of text state and the state of the state of

Administration (Women):

Hydroakoholic transdermal testosterone gels are not FDA approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenie nature and tendency to produce viniZing side effects.

Availability:

Hydroalcoholic transdermal testosterone gek are manufactured in many countries worldwide, and remain widely available. Common trade names include Androgel®, Testim®, and Testogel®: Given the high sophistication and low black market value, counterfeits have not yet been widely reported.

Andronaq (testosterone suspension)

| Androgenic | 100 |
|--------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activit | y moderate |
| Progestational Ac | tivity low |

Description:

Tensteres supersion is an injectube preparation containing tensterence (so eatry, usual) in a work beach arrange loop/balance, "supersion" is invested to a tensor to be an externed potent mass agare. It is often and to be the most provertil injectube storadi analable, action of the drug beach of the storage of the storage of the storage of the table balance is a storage of the storage of the storage of the storage of the table balance is a storage of the storage of the storage data and the storage of the storage of the storage of the storage data and the other table works before a peak teststorage of the storage of the storage data and the storage data and the storage of the storage of the storage data and the storage data and of the storage data and the storage of the storage of the storage data and the storage data and storage data and the storage data and storage data and the storage data and the storage data and the storage data and storage data and the storage

History:

Teotstores supersion is one of the clotst anabolic/mologenic stretch, during all the supback to the 1930s. Used generality to scheme hare njetechte from of the tootstores, testostores supersion predicts the doc-leptent of solves-ating (depot injectors of clother) and the stretch of the stretch of the scheme stretch of the stretch of the stretch of the scheme stretch of the scheme stretch testostores projected and the scheme stretch of the decades more. Provide American Inde marks for the diplave include Storator (Linter), Androna (Cortanja, Agareposition Historican (Plantin), Marcina (Linter), Androna (Cortanja, Agareposition Historican (Plantin), Marcina (Linter), Androna (Cortanja, Teat, Agareposition Historican (Plantin), Marcina (Linter), Androna (Linter), Marcina (Linter), and Teolastore Agarona (Natina) would be too argeneous to list.

Tensterore supervisor shares a clinical application history similar to that of other tostostorem products. Early prescribing galaxies, called for its us to smellent at loss of sex drive, important, and gancal loss of shalip in aging mails with declining hormore levels. Novement, the drug wave contrast proceeding the the transmitter of constant series of shale and the state of the flow of the transmitter of mail and togon is malificant. The drug mail state of the transmitter of mail and togon is malificant, the drug mail of flows and the transmitter of mail and togon is malificant. The drug may however, produce writiknism mails in an uncorrect order.

Albudg the nurther of products comining textostorone suspension steadily downlikel over the years, the drug organo of uniterrupted solability on the U. Spessrepion darg moted as the the ways per to 1998. That year, the FDA that taken action against Shen Laborators is a substance product to the stead of the stead and personal textostance in the steady are produced by the the shell and an queries the stead of the stead steady are produced by the stead of the stead of the stead of the steady steady steady and the stead of the stead of the stead of the text steady and the steady steady steady steady steady steady steady steady steady production of all C-III planmenticities util certain "theregornesis" could be addressed. Visco Inter, Steady stead by course production of all down and the steady stead steady steady steady taken steady taken steady which may reduce the during steady st

How Supplied:

Testostence suspension is available in select human and veterinary drug markets. Composition and dosage may vary by control and manafatture, but suspit) contain 50 mg/mt or 100 mg/mt of steroid nixed in a water-based solution. Testosterone has low water solubility, so the section will an exclusible systemic from a water-based solution when a vial or ampule is left to sit. A quick shake will temporarily place the drug back into suspension, so that the withfatwork dosage shorid always be consistent.

Structural Characteristics:

Teototrow supervisor contains (fee) teototrow in a vuter-based supervisor, although on as sometime about as carriers. Without teotfication, teototrow lies a so that life in the body. Teototrow supervisor may require a minimum of 2-3 ipections per week to minimic oncisient hornow clearations. When calculating doos, expectilly when manyee from one teototrow preparation to another, it is also important to remember that teototrow supervisor contains more active teototrow per million than its existence teototrome clearations. When teototrow teototrow per million that is negative deviations. For example, when the weight of the ester is taken into account, 100 m of teototrow contains exact active active outs a contrastrow.

Side Effects (Estrogenic):

Textsteres is readily atomitted in the body to estable (estrogen). The anomatese (storgen prothesise) entryme is responsible for fine metablism of textstorene. Elevation estables and the estimates of the store of the store of the store of the store stable at complexes is considered at more how assessing to prevent assigned and the store of the store store of the store store of the store of th

Encognite side effects will occur in a dose-dependant memer, with higher doses (above normal therapetic level) of lestosterem enve fieldy to regardle the occurrent use of an artiestrogen or aromatose inhibitor. Since water referrion and loss of matecide definition are common with higher doses of essentement, ties drugs and and the contract the second envelopment of the second envelopment of the second envelopment of the highing phases, where the added water retories with support new strength and macks size, and help foster a suboger attrable: envelopment.

Side Effects (Androgenic):

Teastores is the primery multi-androgen, neoponsible for maintaining scoredurally multisenal characteristics. Floredat levels of testorene are likely to produce androgenes side effects relating only skin, sear, and bodyficial laring growth. Men with a genetic profilopoistion for the links (androgenetic alterycein) may rotice constraints do growth and hading. These concerned about hard has may find a more confinelable option in manufacture potential writing effects of analytic-alterycein strength as a strengt analytic search and the strength and the strength and the strengt androgenesis such as testorence. These may include despeting of the voles, meantual inregularities, charages in skittenese, feasible around an approximate of the origin and the strengt androgenesis.

In anthogon-responsive target issues such as the skin, such, and prostate, the high relative antogenizely of issues is objection on its rolation in only diveositenteme (LPIT). The 5-alph reductise corputs is responsible for this metabolism of tastistatives. The response of the structure of the structure of the structure of the structure with also-specific potentiation of tastistatives are structure. The structure that and the structure of the structure of tastistic value structure in the structure of the structure of the structure of the structure of tastistatives and the structure of the structure of the segmention of tastistatives inhibits.

Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the notential for heratotoxicity with hish doses of testosterone by administerino 400 me of the hormone per day (2,800 mg per week) to a group of mile subjects. The steroid was taken orally so that higher peak concentrations would be reached in hepatic itsues compared to intramenduar injections. The hormone was given daily for 20 days, and produced no significant changes in liver enzyme values including serum albumin, bilrubin, admin-amino-transformes, and lakline tooolstatuse.⁴⁵

Side Effects (Cardiovascular):

Ambelicantogramis strends can have deterious effects on serum checkeror. This includes a tendency to role and EUL (good) checkerou values and increase (LL, (bat) checkerout values, which may shift the HDL to LDL bahare in a direction that fixors greater risk of advendant on the dose, route of administration (eral values, injectable), type of sheatid increasing the risk of curved in many and the route of the route of the route protocol and the route of the route route of the route route of the route route of the route of

Teststerow tends to have a much iss durantic impact on cardiovaschar ink fectors than synthetic stocks. This is an part to is sopress to metabolismit be low, which also it is the low set effect on the lapping immagenet of chokesterol. The aromatization of teststeron to established and helps to might the negative effects of androgen on sensus fixed. In one study, 220 mg per vock of iteststeron energy energy and when helps with the maximal synthesis and the lapping of the lapping of the lapping maximal synthesis and the lapping of the lapping of the lapping maximal synthesis and the lapping of the lapping of the lapping of the maximal synthesis and the lapping of the lapping of the lapping of the maximal set of the lapping of the lapping of the lapping of the lapping maximal synthesis and the lapping of the lapping of the lapping of the maximal set of the lapping of the lapping of the lapping of the maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping of the lapping of the lapping of the lapping maximal set of the lapping maximal set of the lapping maximal set of the lapping of the lappi

Due to the positive influence of estimgan on serum jujici, surroution cintue or echangines cintue are porterior to a commisso influitos for these occurrent only inclusionscalar heads, as they offer a partial estrogenic effects in the loses. This also shown to poteriarily improve inju profiles and offers some of the regards effects of antiogenase. Which does of 600 mg ere lose of testistence per week, the impact on light profile transits to be noticeable but not dimentif, mixing an mis-strongen (for campitoretion perpose) performs surroscenase. Does 1000 mg ere lose per week have also filted to produce summer and significant damages Does 1000 mg ere lose per week waite input of campitor the strongenetic damages and the strongenetic strongenetic strongenetic and many strongenetic transitions and the strongenetic and many strongenetic strongenetic waites strongenetic strongenetic waites strongenetic strongenetic strongenetic waites of all antiodentic strongenetic strongene

To help reduce cardiorsocular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of starated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fish oils (4) grams per day) and a natural cholesterolamicoxidant formals such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic stewisk when taken in doess sufficient to promote maske gain are expected to appresse molgrouss testsotreem production. Testsotreem is the primary mark androgen, and offers strong engine feedback on endogenous testsotreous production. Testsotreem-based large ville. Rowsie, here a strong effect on the hypothamic regulation of natural stewid hermions. Without the interversion of testsotreom-similarity advances, testsotreem-based based trendo normal within 1-4 month of data gescession. Note that prolonging thereorem, the strong strength terror test and the propagation of the strength of the stren

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Teostoreous suspension contains unfusiohed teostoreous particles, which form a shortarding repository in the mack-following piezoton. Depending on the site of the particles and other agents present, njecotons of teostoreous suspension may use large markets. However, the methods as large as 21 gauges for high-field the complex, and can be very acconditability to use, and redues. Vefering particles. These constraints, and are used as a large end of the particles and the complex states of the particles and the site of the particles of the complex microscillate states and the particles. These constraints are higher affend, and are toos with the reducts, and is agreently well beinstic.

Administration (Men):

To text androgan instificiency, the proceeding galdelines for testosterone supersion recommend a dose of 25-50 mg, which is given exp 2-3 tenes per work. When used for mackbuilding proposes, testosterone supposes in so dona administered at a dose of 100-200 mg per pictosicu which is given every 2-adr or 13 day. Additus backage as decisione, an externily amozing, the user scening to just finite with backed macker muss in a very short period of time. Each when they were being ammafication, the ULS 30 mit, which (100 mg/rtl) user always the most scagif after for this procedure, as each woold run the cycle for aboat a more. Adding this for disp does require a forquest injection scaladil, a well-findelin ment. Adding this for disp does require a forquest injection scaladil, a well-findelin more scalade injection size, litting the samker macke groups such as the dehead, triceps, and calues.

Those locking for only a potert mms agent are often externely huppy with the reachs provided by testosterone suspension, first poteck certainly has a source pertaining. But those athletes who want net just quarity hat quality are likely to be with density of the source of the source of the source or the source of the physicage with this density. Alternative source of the source of the source of the physicage with this density has a source of the source or the source of the source of the source of the source of the source or neutre methods in such a source, so tastistances supervised, which would allow for less concept holding h such a source, one tastistances supervised, which would allow for less concept holding h such a source, one the density of a source of other less or neutre methods in source, densities for the density of the source of the source of the source of the source of the density of the density of the source of the source of the source of the source of the density of the source of the density of the source of the source of the source of the density of the source of the source of the density of the source of the source of the density of the source of the density of the source of the source of the density of the source of the source of the density of the source of the density of the source of the density of the source of the source of the density of the source of the density of the source of the density of the source of the source of the density of the source of the density of the sourc

Administration (Women):

Testostences suspension is medy used with women in clinical medicine. When applied, it is most often used as secondary treatment for inopenable breast career. Does segress for the application may reach 100 mg three times per week, a level well ato the threshold likely to cause strong withing side effects. Testosterone suspension is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce withing side effects.

Availability:

Pharmaceutical preparations containing testosterone in an aqueous suspension remain scarce. The bulk of the supply presently comes from underground and export steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Testosterone suspension has been unavailable in the United States for many years now. No old products should still be in circulation. Because the FDA never officially withdrew this drug from, it can be specially ordered through a small number of compounding pharmacies. Anything else bearing a U.S. manufacturer rame is counterfeit.

Testosterone suspension is still produced in India under the Aquaviron brand name, now by Piramal HC. It comes in the strength of 25 mg/mL, and is packaged in 1 mL ampules.

Cheque Drops® (mibolerone)

| Androgenic | 1,800 |
|---------------------|--|
| Anabolic | 4,100 |
| Standard | Methyltestosterone (oral) |
| | 7,17-dimethyl-19-norandrost-4-en-3-one-17b-ol eta-Hydroxy-7alpha,17-dimethylestr-4-en-3-one |
| Estrogenic Activity | y high |
| | |

Description:

Melesteres is an oral anabolic strond, ancentrally derived dimethylated methylated. agart is specifically, 71-fordingblied methylatook, significantly more potert as an anabolic and analoganic agart films is non-mellyhied peret. Over the years, molestene inseared and apartians armage bodylakless as being our of the attrayest attentiates over madk. This effective in microgram, nor miligram, answares. Daring standard animal assess, microbaron effective and determined by more 41 mers the analoganic activity of mellyholicshorms when give orally. In constast, it had only 18 mers the analoganic activity of mellyholicshorms when give senses). Encogenic and progenitation properties are above type protocores of the data strandard agart and progenitation properties and above type protocores of the data strandard agart and the adapt. If early a strandard animal bases, the data strandard agart and the adapt. If early a strandard animal assess, the data strandard agart and the adapt. If early a strandard animal assess, the data strandard agart and the adapt. If early a strandard and the data strandard agart and base a vector or correspection.

History:

Melosicous uns first described in 1963.⁵⁰⁰ It was developed into a veterinary medicine darige the 1965's (bigs) which so did the drag under the bond mure Chapea Doops). The preparation constanted 100 megrin of astrond in a 55 rd is bond, for a total stored content of a bond and a proparation table Chapea Doodsheet Doop (Source Chapea Doops), and a store of the sto

Arong attacks, mbokrone has alweys been seen with a light beel of mayting, perhaps purply due to is infraind availability. These structure has high beel of may structure & Update and availability. These structure has a light been been been black proceeding bies to mbokrone scenarizing products on a light branch, does have the structure has a light branch branch branch branch branch branch the approximation of the structure has a light branch branch branch water structure has a light branch branch branch branch branch branch water structure has a light branch branch branch branch branch branch branch water structure has a light branch branch branch branch branch branch branch water structure has a light branch bran

How Supplied:

Mibelerone is no longer available as a prescription drug product. When produced it most commonly came in the form of an oral solution based in propylene glycol, carrying 100mcg of steroid per utiliker in a 55 mL botte.

Structural Characteristics:

Mbolemen is a modified form of manholme. It differs by 1) the addition of a methyl group at carbon 17-alpha to protect the hormone during cerl administration and 2) the introduction of a methyl group at carbon 7 (alpha), which inhibits 5-alpha reduction and increases relative antrogenicity. 7,17-dimethylated steroids also tend to be very resistant to metabolism and serun-binding provides, greatly erhancing their relative biological activity.

Side Effects (Estrogenic):

Medestence is arountized by the lode, and is considered a lightly estimagent structure is a conversion to 7.7.1/entirelystandic (in entrogon with light) thological archive). Genocemination may be a concern during transmit, especially when higher fram memal an archicle loss of marces defines as for hist archives are structured as the formation of the structure of the structure of the structure of the Nondonection. The marce discharge transmits in higher framework in the Nondonection of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the structure of the Nondonection of the structure of the structure of the structure of the structure of the Nondonection of the structure of the Nondonection of the structure of th

It is often that mibilisme also displays strong activity as a progestin in the body. The side effects associated only progestrome are as roughts in the hose of creating progesting is also anyore the strong strong production and enhanced rate of fit stronge. Progestiss is also anyore the strongest strong strong strong strong provide the progestion is also anyore the strong strong strong strong strong program strong strong to be as strong strong between these two hormcose here, such that gencomassin might even coars with the help dynagrism structure correspond of this disorder, is often sufficient on antie-strongen, which inhibits the estrogenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately and the strongenic compound of this disorder, is often sufficient to might approximately the strongenic compound of this disorder.

Side Effects (Androgenic):

Although cossified as an anabole strend, androgenic side effects are sill common with the substrate. This may include boost of objects, and are sub-objectian har growth. Anabolicitudrogenic storelists most may find a miller anabole starbolistic to the analogenic effects of this second rung field a miller anabole starbolistic to the chargenic effects of the store rung rule and analogenic probability of the store that the store of the store rung rule and analogenic probability of the store rungs of the store rung rule and analogenic probability of the store of the store of the store rung rule growth, and charge effects of the store respective, employee its store of the store

Side Effects (Hepatotoxicity):

Melestenes is a c17-alpha akylatel compound. This altention protects the dung tisms descritation by the sent allowing a very high percentage of the dung entry into be bloodstream fallwing containtainton. C17-alpha akylated ambolicandengenes derochs can be highestester. De bestaged or high percentage of the stages. In artic histories each cycle to remote here factorian and occali heads that due of c17-alpha akylated strends is controlly initial of 64 weeks, in an effect to avoid scatting per strain. Severe bein complexiting and the periodic matter in which most people use onla ambolicantelogues articula, although cannot be excluded with the seried, opeoplical birth of the strain and the periodic matter in which most people use of an ambolicantelogues articula, although cannot be excluded with the streed, opeoplical birth of the strain and the series of the strain and the streed of the strain for Charge Deps, mentaries only one lamm study here controls. The streed here, and the study use terminated on the due birth periodicity.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anabolizardingganis starolis can huve delutricos effects on serum chesterot. This includes a tendency to rolex IEL (good) cohesterol values and increase ILL. (bud) cohesteroli values, with the HDL (sod) cohesteroli values and increase (LL. (bud) cohesteroli values (LL. (bud) cohesteroli

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ois (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

All arabok'androgenic steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimaliting substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadorophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss hwe shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴²² This is caused by the fat-soluble mature of steroid hormones, which can alwo some of the drug to dissolve with undigsted distary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

Mobilence was nover approved for use in human. Prescribing guidelies are unrevalidely. In the athickit areas, the drug is used intermetinely due to its high level of hepatoxicsity, with cycles stually lasting no more than 6 weeks followed by 6-5 weeks of A duby dosage of 20 to 500mg strate stormers for holydaling parposes. The level is synchrody sufficient for above studies are also been apprecisively and the storage storage of the origination of the storage of the storage of the storage and the control of the storage of the

Administration (Women):

Mibolerone was never approved for use in humans. Prescribing guidelines are unavailable. Mibolerone is generally not recommended for women for physique- or performanceenhancing purposes due to its very strong nature and tendency to produce virilizing side effects.

Availability:

Mibolerone is sold in the U.S. as a compounded veterinary medicine only. No commercial preparations containing this drug are known to exist worldwide. Mibolerone remains available on the black market in underground preparations only.

Deca-Durabolin® (nandrolone decanoate)

| Androgenic | 37 |
|--------------------|--|
| Anabolic | 125 |
| Standard | Testosterone |
| Chemical Names | 19-norandrost-4-en-3-one-17beta-ol 17beta-hydroxy-estr-4-en-3-one |
| Estrogenic Activit | y low |
| Progestational Ac | tivity moderate |

Description:

Nandroken decanotie is an injechels form of the anabolis stored macholose. The document ester provides a slow relaxes of multichone from the site of opticotic, lasting of up to three works. Nandroken is very similar to isotatorene is structure, although it lacks a calibon attain at the 10th position (laceus is other man; 10- Americationene). Else calibon attain at the 10th position (laceus is other man; 10- Americationene). Else however, its insue-bailing activity is accompatible by and and and and the last to do with the neakness of multichone to a worker strengt, dipplement/toolse, in the same androgas-responsive target issues that postentia the action of testoratorene (by concerting its 10 JMT). Its mill properties clow to a worker strengt, dipplement/toolse, and concerting its 10 JMT). Its mill properties clow that the action of testoratorene (by concerting et al.DMT). Its mill properties clow that the same and easier of the protective significant strength and lacks much highly factored by adultates took pathogen and compare take effects.

History:

Nandokee decanate was first described in 1960,⁴³³ and became a prescription mediation in 1962. Use advectod pol the international plannerscends agint Organos, and solt under the brand name Dace-Darabolin. The mare Dace-Darabolin davetse fut the product contins a variant of Organom previously pepular matheoime injectibe Darabolin (mathokee plansphopionate) using an ester of 10 enhon atoms. Organon equands the mediate for mathokee docenates very applicable plansing blowing is related. Daraboli, Daraboli control estimates and the large mathef presence of Organo, Dace-Darabolis conclusers on of them on wideh distribut andhole kentois in the world.

When first introduced to the Lindea States, neutriduce documents (like Databoli) uses proceeding for any region dimense. List of indications includely per and postporteniles use for building from muss, outsportensis, advanced breast cancer, weight loss due to introduced the state of the and development restatations in children. The darg was statingly sold in a doogs of only 50 maprix owing no two years over the data a 100 mprint version, reducing the random large data the state of the state state of the state state of the state state of the state state of the state of t

Albuch the drug had been replied foronbly for a great many medical uses for approximitely a decide, by the mix1/2019 his finitefact uses for marknoine denames were being refined, both in the U.S. and abnoal. FDA approved prescribing information from 1975 his mandhook decorated as yields whether the start has a prosent provide the start of the start of the start of the start has a prograph horman is more analbels. It was also decread "possible direction" and ing approximation of the start operation of the start certain types of anersit. More time was given to insertigate the potential "kss than effective"

Modern (approved) medical applications for the drug are oven more refined than they were in the misi-107% in the ilused States, the drug is any one JFA approved for transfer amenta, allowing h is often also used "wfi label" to preserve tan mues is HFV possible appendent of the state support the set of this (minisity wfit) patients disting from severe sources, like objective and advanced breast cances. The Organon Dace-Daraboli heard of ransforkee decancate remains widely available locidg, now distributed by use parter cancers, and is also madilized, matchine decances is produced as a generic drug in range counties, and is also modernate used remotes other distribute board marks, body the hours and verticing the state of the

How Supplied:

Nandrolone decanoate is widely available in human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 mg/ml, 50 mg/ml, 100 mg/ml, or 200 mg/ml of steroid dissolved in oil.

Structural Characteristics:

Natholice decanote is a modified form of randroken, where a carboxylic scal ester (docanote acid) have non-mathed to the 17-ben blyoxyl open. Esteroid services are set polar than free stroxids, and are absorbed more alway from the area of rajection. Once is the bloodstream, the est is removed to yield like (acidar) modified. Estimizial attention are designed to prokeng the window of thempeatic effect. Glowing administration, allowing docated area of the provided and the strong the strong the strong the strong the Number of extremely provides a damy problem in mathema enders. Jet-48-bases Glowing doep intranscent rajection, which steady declines to mar baseline levels. approximally to works huter. The life of or mathematic enderscenter is 7-12 daws.

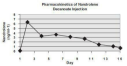


Figure 1. Pharmacokinetics of 200 mg Nandrolone Decanoate injection. Source: Pharmacokinetic parameters of mandrolone (19-nortestosterone) after intranascular administration of nandrolone decanoate (Deca-Durabolin®) to healthy volunteers. Winand H. Bosch A. Donker C. Acta Endocrinol 1985 sumo 27119-30.

Side Effects (Estrogenic):

Nandoken hus a low incluses for entropin convestion, estimated to be only about 20% of the sense with tooscoress¹⁴. This is because while the loce can convert matchoose to estindial, in other more active sites of storoid aromatization such as adjacent tissue matchoose is fit here open to this proceed. "Consequently, ensurposcileated and effects and an motion with higher design (assesser and may cance). Each officient and a store and water texterion, hold (gain, and garcenous), and and store gain and what restored is the light (gain, and garcenous), and and store gains what is estimated what restored is the same amounts in the low texterior hold water texteriors, hold (gain, and garcenous), and and texteriors hold here cover clinicar (controls estings in proceedings in systems). A contrainer similarity can be applied efficiently controls estings by preventing by synthesis. A contrainer similarity can be applied proceeding and the store of the store of the store store of the store of the store store

It is of need that matchetee has some activity as a progestin in the body.⁴⁴ Mkhodg, progestrever is a > 0.9 stored, nervous of this group as in 19-mongastrever creates a harmone with greater briefing affithely for its corresponding receptor. Sharing this truit, many 19-ner anabolic storesholds are about to how one mainful for the progestreve receptor as weld⁴⁷. The side effects associated with progestreve are similar to those of estimation stores. The side effects associated with progestreve are similar to those of estimation stores. The side effects associated with progestreve are similar to those of estimation stores. The progestria also asigners the similarity effect of strongers on marring time growth. Then express the has a sing squared between that we to homes here, and has provide. The similar the similarity effect of strongers on marrings have and the true of an artis-strongen, which inhits the estrogenic competent of this disorder, is often atticket to mining encoursing and any numbroher.

Side Effects (Androgenic):

Although classified as an anabolic steroid, antrogenic side effects are still possible with this substance, espectically with higher doess. This may include boots of object with a merihody/ficial hair growth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Women are warned of the potertial vitilizing effects of anabolic/androgenic steroids. These my include a deepening of the voice, mentatul regularities, changes in skite toture, these my include a deepening of the voice, mentatul regularities, changes in skite toture, and the statement of the steroids of the potential vitilizing effects of anabolic and the steroids of the statement of the steroids of the steroid steroids of the steroid of the steroids of the heat har growth, and editoral entragement. Nantrobne is a steroid with relatively tow androgaria exityly relative to a situaci-bailing actions, making the threabold for storag androgarie side effects comparably ligher than with more androgarie agents such as testosterone, refutualmostenoloue, or thaoxymesterone. It is also important to point out that due to is mill androgarie matre and ability to suppress endogarous testosterone, rendrolloue is provide to interferig with this in minus when used without andret androgarie androgarie and the start of the start

Note that in antrogen-response trapt issues such as the skin, scab, and prostate, the relative antrogenicy of matrolones is reduced by in reduction to disploratometoisene (DRN) $^{(42)}_{-}$ (DR S-lapha reductase enzyme is responselse for this matcholism matcholism. The constructive of a S-lapha reductase influence such as financiefic or matcholism. The constructive of a S-lapha reductase influence such as financiefic arrange the tradency of matholism is provided in the star of the star influence in the star of the star influence in the star of the star influence in the star of the star influence in the star of the star influence in the star of the star influence in the star of the star influence in the star of the s

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects in healthy subjects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabolizanteganis stavalis can huve delutricos effects on serum chosteard. This includes a nelmery to reduce IRL (apod) chosteard values and increase IRL (bash) chosteard values values, which may diff the IRL (apod) chosteard values and increase IRL (bash) chosteard values dependent on the does, encote of administration (calu N₁, given strateging administrative G00 are foundable), and level of resistance to lapstice metabolism. Statisadministrative G00 are foundable of constrained a 20% constrained a 20% enclution in IRL. chosteard levels⁴⁴. This suppression is slightly genate than thermosteard administrative G00 are produced by the strateging of the strateging of the constraint of the strateging of the scrapest of the test strateging of the strateging of the strateging of the strateging of the Anabolizanteging is strateging on the strateging of the strateg

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of star-tated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ois (4 grams per day) and a natural cholsterol/articixelar formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al analysical angunt extension when taken in doess sufficient to promote make gain are expected to suppress condensors testsotower production. Statica similaristratif [00 rupp; week of markoine decanote for 6 weeks lawe demonstrated an approximate 57% induction in semantisatorone lovel daring planety. Ja a doago 2010 rupp rev sed, the national metal of $20^{14} {\rm eHz}$ is to belowd that the proparational activity of markoise national constraints of the suppression of lossing of the supervised on the markoit in spite of a low tendency for estogan conversion.⁴⁴¹ What the intervention of testsotenov-stranding solutions, testsotence levels should return to normal with 1-5, markins of the gas essents. Notes that prolonged hopograndetorphic hypograndian can decelop escondup to strend alson, measurements, media marking media intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

For general anabolic effects, early prescribing guidelines recommend a dosage of 50-100 mg every 3-4 weeks for 12 weeks. To treat renal anemia, the prescribing guidelines for

matchows decanote recomment a dosage of 100–200 mg per week. The usual dosage for physics- or preformace-structure gauges based on the structure of 20-600 mg per week, taken in cycles 8 to 12 weeks in length. This level is afficient for most users to motic measurable gains in the market mass and strength it is often ated that matchouse documents will called its optimal discr (bein gausside discr ratio) at 2 mg per sport of documents will be a strength of the strength of the strength of the strength document will be a strength of the strength of the strength of the document will be a strength of the strength of the strength of the document will be a strength of the strength of the strength of the document will be a strength of the strength

Nandroken decanate is often combined with other strends for an enhanced effect A combinition of 20-04 mijweck of manufachose decanates and 10-20 mg daily of Westrolli, for ecample, is noted to grady enhance the took of mascianity and definition when distrigating: A transg measurementing androgas like likelisatifies of twolfseless could also be used, again proving an enhanced level of hundroses and density to the marcels. Beign anotherid system, march budken matchicen can also be incorporated for blat cycles with acceptable results. The class: "Deca and D-bdi" stack (usual) 207-400 mg of anality task consistence and station 12-55 mg over disensity data (bally 207-400 mg of anality task consistence and station 12-55 mg over disensity data (bally 207-400 strenger andropen accids and station 12-55 mg over disensity data (bally 207-400 strenger andropen accids and station) in the substituted, producing genetic results, budy thin new vater creations.

Administration (Women):

For general anabole cflexts, early prescribing guideins: scoremend a dooago of 50-100 mg corey 34 weeks for 12 weeks. To hust renul ancruit, the prescribing guideins for mathetime documents ensuring the property of the transmission of the physics or preference-straticity graphysics, it alonge of 50 mg ren week is most conformed with viriation symptoms when taking the compound. Studies have demonstrated light biothably (minor but statiscilly singularized tradence of viking site effects) with a dose of 100 mg every ofter weeks for 24 weeks, ⁸⁴⁴ while bayes must effects with a dose of 100 mg every ofter weeks for 24 weeks, ⁸⁴⁴ while bayes must effect with a dose of 100 mg every ofter weeks for 24 weeks, ⁸⁴⁴ while bayes must effect with a dose of 100 mg every ofter weeks for 24 weeks, ⁸⁴⁴ while bayes must statisk biothermiting with the states-acting matchiese. Darkbailt in sufficient of the states action of the states-acting matchiese Darkbailt in the outline with the dot of the states-acting matchiese. Darkbailt in the the withmediant of the states action of the states action of the states action of the states action withmediant of the states action of the states action of the states action of the states action withmediant of the states action of the states action

Availability:

Nandroken decennote continues to decline in prominence as a plenumeentical product these to is invited user indical medicic. The drug is presently measurable in the United States. Many Western nations continue to market the drug though its production is increasingly engligibility to issue regulated markets in which. Legatimer themmeostical forms are highly networking some of the rome popular products and change on the global planumeential market, we have much the Bohwing downstrations.

In November 2009, Organon (a subsidiary of ScheringPlough since 2007) became part of Merck/MSD. All Organon products are expected to transition over to this label. It is urknown what (if any) changes to expect in the global distribution of Deca-Durabolin products.

Brand mame Deca-Durabolin is not available in the United States. All products bearing this label are counterfait. Watson Labs and Schein Pharmaceuticals generics have also been discontinued. This drug is presently unavailable in the U.S.

Norm Hellas Deca (100 mg/mL nandrolone decanoate in 2 mL viak) from Greece is available, but also widely counter/feind. The firm uses a paterted photochronic label to deter counter/feing which carries a metallic/holographic watermark of the Norma Hellas logo.

Greek Deca-Durabolin (formerly from Organon) has been another widely counterfeited product. It is one of only a handhil of European randrolone injectables to be found in mithdosed vials, makine it an easy tranet for counterfeiters that lack the canactivit to moduee eless ampules. This product should be considered fake unless it comes in a box with the proper Greek drug ID sticker. As with all Greek drugs, the sticker should show a hidden mark under UV light.

Greek Extraboline may be in circulation. It is also a common target of counterfeiting. As with all Greek drugs, this product should contain a peel-off pharmacy sticker that reveals a hidden watermark under UV lighting. All Extraboline in circulation will also carry a holographic image directly on the vial label.

Deca-Promabol from P&B Labs India is no longer in production. The company currently markets only a 25 mg/mL version of this drug in 1 mL ampules. Many counterfeits of the former 100 mg/mL product are still in circulation.

Decabolic from Asia Pharma (Mahaysia) is now approved for sale through pharmacies in Thaland, and is fairly popular on the black market. Each box should carry a scratch-off security stiker, which will display a code that can be valished on the company website.

Balkan Pharmaceuticals (Moldova) makes the product Nandrolona D. It is prepared in both 1 mL ampules and multi-dose vials.

Delatestryl® (testosterone enanthate)

| Androgenic | 100 |
|---------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Acti | ivity low |

Description:

Teastoreme cumflate is a slow-acting njectable from of the analogue tostotence. Following doep intranaccia injection, the diney is obsigned to provide a statistic release of testostences into the bloodstream for approximately 2 o 3 week. In order to maintain normal physiological testo of testostence drag randong represent therapies, njectices of statistoreme cumflate are usually required at least every two weeks, although more mexicous physicism validantistic effect on the statistic release to the statistic release to insections and strength. Usually required at least every two weeks, although more mexicous physicism validantistic effect of the statistic releases in nucle runs and strength.

History:

Teaststence currantee first appeared on Western dng murkets during the early 190%. It was the first above-sing object and teast of restorators on the widely adapted Western medicine, and offstelidy replaced teststatement propiones and its structures that and the structure of the structure of the structure of the structure times, more notably to Mead Johnson, BTG, Saviert, and in December 2005, Indexes. The more promise Them to obstatement empirication could be structure and the structure of the structure of the structure and the structure of the structure structure of the structure structure of the structure structure obstates in Structure and the structure structure obstates of the structure structure representation.

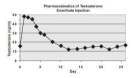
Testosterone eranftate is most often used clinically to replace normal levels of testosterone in addr maies sufficient quintished androgon levels. This may runnafis test with a loss of blido, lean mascle muse, and normal energy and viger. Testosterone eranftate is also used to treat undescended testiskes and delayed pathetry in adolescent muses, and occusionally as a secondary medication during imperable breast cancer in women. This form of testosterone and testosterone and the second se In a site been studed with great success as a mule brit control option.²⁴⁹ Workly spectrum of 200 mg verse shown to efficiently hover sperm productions for most men within three mention of treatment, a state of suppression that remained ural after the drug uses discontinued. With the current station successing analysical models; statevisk, however, it applies of the growing number of subservations therapies, testorskervace examinate means widely prescribed growing the statevistic of the workl.

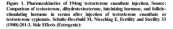
How Supplied:

Testosterone enanthate is widely available in human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 50 mg/ml, 200 mg/ml, or 250 mg/ml of steroid dissolved in oil.

Structural Characteristics:

Textstere emritue is a modified from of textsterow, where a carboxyle acid seter (controls acid) has hen mathed to be 17-ben hydrody groups. Esterikel from so textsterore are less poler than fees textsterore, and are absorbed more skwyl from the are of njectro. Does in the blodsterma the easier's removed to yield fee (acids) textsterore. Esterikel forms of textsterore are designed to prolong the window of theopartic reflect blowing administration, adding far lass for gener njections achduel compared to injections of Fee (instrafield) starvid. The half-file of teststerore enarthute is approximative adding safer injection.





Teostorese is readily atomitted in the body to stratical (estrogen). The anomases (estrogen optitutes) organe is responsible for fine metabloom for isotoatores. Elevation estrogen body and the optimation of the strationary of the strategies of the generative strategies and the strategies of the strategies of the strategies is defined. One may have the strategies at the strategies is defined to the strategies of the strategies of the strategies of the strategies in the strategies of the strategies of the strategies of the strategies in the strategies of the strateg

Enorgenic side effects will occur in a doss-dependant memore, with higher doses (above normal therapetic level) of testosterone more fieldy to acquire the constant use of an antiestingan or anomatose inhibitor. Since water reterion and loss of matecide delinition are common with higher doses of testosterone, tilde mage a maged and the constant of the single delinear and the single and the single and the backing planes, where the added water reterior with support naw strength and mack size, and help foster a stronger anabole environment.

ORE LARGES UMBILIZEDES.

Textsteres is the primery multi-androgen, nesponsible for maintaining soccordary mulssenal characteristics. Enseted levels of textsteres are fields produce androgenic side effects including objection and the set of the set of the set of the set of the probability of the time of the set of the set of the set of the set of the leaders. If the set of the set of

In anticogen-responsive target tissues such as the skin, such, and prostate, the high relative anticogravity of testatorene is dopedual on its rolation to of hydrostostorene (DHT). The 5-alpha reductistic enzyme is responsible for this metabolism of testostorene. The concurrent use of 3-abb reductistics mitholic values in structured on interview of interview with skin-specific potentiation of testostorene action, lowering the tendency of testostorene dops to produce antipose side effection. It is improved to testosterile the analytic and segments on featosterile antipose side effection. It is improved to testosterile that analytic segmention of testosteriew's analytic and antipose properties is not possible, even with tool 3-alphar necktares imbibition.

Side Effects (Hepatotoxicity);

Testosterne does not huw hepatotoxie effects, for toxicity is utilicly. One study examine the potertial for hepatoxicity with high does of testosterne by administering 400 mg of the lowmee per day (2,800 mg per work) to a group of mile subjects. The second was ation only so that higher peak concentrations would be reached in higher fusion study and the study of the study of the study of the study of the produced to significant changes in liver enzyme values including serum absami, bifrabit, attinie-amire-amisters, and aladier potopattens.⁴⁴⁷

Side Effects (Cardiovascular):

Ambolicanteganis strenids can huve delutricos effection on serum chocketorel. This includes a tendency to reduce IDML (apod) chocketorel voluces and increases (LL, (bad) chocketorel voluces, which may shift the HDL to LDL balance in a direction that fivores genet risk of antirocherosis. The evaluation match characteristic and one serum lipits is dependint on the dose, route or a daministration (eral ws. lipitedbal, type of steadid Ambolicanteganistic antirocherosistic depending and the state of the state Ambolicanteganistic antistation (eral ws. lipitedbal, type of steadid Ambolicanteganistic antistation (eral ws. lipitedbal, type of steadid Ambolicanteganistic antistation (eral ws. lipitedbal, type) and the steadies reduces chocketion (eral molecular) affection.

Tensteroe tends to have a much less damatic inpact on cardiovascular risk factors dam synthest strotch. This is dae part to is openness to matchiomity the focus, which also it to have less effect on the lepatic imagement of choisetsol. The aromatisation of tistustances to estandard also helps to might the magnite effects of andragons on senardistribution of the strong of the length of effects of the strong of the strong matching significant effects of the strong of the strong of the strong matching significant time of the strong of the strong of the strong of the tensor of the strong of the strong of the strong of the strong of the tensor of the strong of the strong of the strong of the strong of the tensor of the strong of the strong of the strong of the strong of the tensor of the strong of the tensor of the strong of the tensor of the strong of the tensor of the strong of the tensor of the strong of the str

Due to the positive influence of estingan on serum lipid, surroution cirate or champione cirate are preferred to acommodes rights for those coverent of which acomodorscalar heads, as they offer a partial estrogenic effects in the low. This allows them to potentially improve loss per work, the impact on lipid profile study to be not domain, immigran or loss per work, the impact on lipid profile study to be not domain, immigran per work, law and lipid to profile study to be noticeable bar not domain, immigran per work, law and lind to produce studies lay addiment (atomges in LD/TUL) cholestory, trajecortics, apolopportion IEC-III, C-reative protein, and institution a rolative work immed not an induction a rolative work immed or notoroscolar in the knows? moderate doses, injectable testosterone esters are usually considered to be the safest of all anabolic/androgenic steroids.

To help reduce cardiorascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fash ois (4) grams per day) and a natural cholesterolamitoxidant formals such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicandoguis atensis when taken in doses sufficient to promete maske gain are expected to asprose condigorous testostorem production. Testostorem is the primury multisandogun, and offers storage negative fordback on endogunous testosterous production. Testosterone-based lange will, Rossies, here a storage disk on the hypothaniar regulation of natural storid hormores. Without the interversion of stostosteron-similaring substances, testosterone based and letteron to multi within 1-4 metric for digressession. Note that prolongial phogomatorepite hypoprandism can develop secondiny to steroid abuse, messatirity ancideal traterorison.

As with all ambolicatopenis estructis, it is utilisely that one will retain every pound of new bodyweight after a cycle is concluded. This is expected in the web withdrawing from storego (arountizing) androgen like testosteome, as much of the new weight gain is likely to be in the from of weater retearing capital of mining after drug discontranee. An intubative of analysis and catabolic hormous drug that drug discontranee. An intubative disconter that is a usinovable for the weater of most these. Proper a analysing drug therapy is usually accommended to help restore hormous balance more quickly, utimately helping the are retain more muck tissue:

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

To treat antrogen issufficiency, the prescribing galdenss for testosterone emuthate call for a dosage of 59-400 gm every 2 to 4 weeks. Although active in the body for a longer time, testosterone emuthate is usually injected on a weekly basis for muscle-badding purposes. The usual dosage for physique: or preformance-entrancing purposes is in the range of 200-600 mg per week, taken in cycles 6 to 12 weeks in length. This level is sufficient for most users to notice exceptional gains in muscle size and attempt.

Teostoreo is usually incorporated into balking plases of training, when added water teartion will be of life consequence, the user more concerned with now muss than definition. Some do incorporate the drug into carting cycles as well, but typically in lower does (100-2010 gpre weeds) and/or whence compared by an amountse eithbor to lose estingan beek under control. Teostoreore enrathuts is a very effective anable dang, and is dim used abave will gear thereft. Some, however, that a need to back it with offerent anablechandrogene services for a stronger effect, in which case an additional 20-400 mg double provide sharkening leads with an signaling hypotheses. The stronger estimates well provide sharkening leads with an signaling hypotheses (Linestoren is dimmally very verside, and can be corbined with many ofter anablechandrogene streids to take

Administration (Women):

Testostence emritter is rarely used with women in clinical medicine. When applied, it is most often used as secondary medication during inprovable beast cancey, when other therapits have fished to produce a desirable effect and suppression of ovarian function is necessary. Testosterone enrathet is not recommended for women for physique- or performance-enhancing parposes due to is storage androgenic nature, tendency to produce withing side effects, and show-crinic characteristics (miniscription blod levels difficult to control).

Availability:

resonance enamine ename in this watery minimatured formor operator resonance worldwide. It is produced in many generic and brand name forms. In reviewing some of the products and changes in the global pharmaceutical market, we have made the following observations.

In 2006, Savient sold the rights for Delatestryl to Indevus (United States). Indevus subsequently became a subsidiary of Endo Pharmacenticals in March of 2009. Brand name Delatestryl remains available in the United States under the new company (at 200 mg/ml, strength).

Generic versions are also available in the United States (also at a dosage of 200 mg/mL) by Watson, Paddock, and Synerex.

Norma Hellas (Greece), makers of Norma Hellas Nandrolone, recently added a generic 250 mgirdt. testosterone enanthate injectable to their product offerings. It comes in a single dark arrher 1 ml. gluss ampule, and is packaged 1 ampule per box. Be sure to look at the Greek Pharmacy sticker under UV light to assure you have a legitimate product.

Bayer took control of Schering AG in December 2006. Following this acquisition, the Schering Primoteston and Testovinon Depot products were transitioned over the Bayer brand and logo (the products now bear the full company nume Bayer Schering Planum). Note that numy counterficients have not yet made this change in their own products, and thus are still duplicating the old Schering babes and boots:

Cidoteston is produced in Egypt by CID (Chemical Industries Development). It comes in 1 mL angules, containing 250 mg/mL of steroid. This product has been counterfeited, though the current most popular copy can be quickly identified by close examination of the fine details (logo, graphics) on the box.

The French version Testosterone Heptylate is still in production. It is now sold under the S.E.R.P. label, and available in the familiar 250 mg/ml, strength and 1 mL amptule. This drug is frequently exported to developing markets with close trade relations to France, such as Lebanon.

Androtardyl is also produced in France, and occasionally circulates on the black market. Again, be sure to look for the proper box before buying.

Testo-Enant is another brand in Europe, this one being made by Geymonat in Italy. These ampulses contain 250 mg of steroid, either in 1 mL or 2 mL of 01. Currently fakes are not a problem, however, this steroid is not found on the black market in high volumes.

Galenika makes Testosteron Depo in Serbia. These 1 mL ampules contain 250 mg/mL of steroid, and are extremely cheap at the retail level in their country of origin.

Jelfa produces Testosteronum Prolongatum in Poland. It is made at a dosage of 100 mg/mL. Each box contains five 1 mL ampules, which are themselves made of clear glass and carry a paper label. The packaging of this product was recently updated to reflect a more modern color-gradient design.

Testoviron Depot from German Remedies in India remains in production. The product comes is made in foil and plastic blister packs.

The Indian export firm Alpha-Pharma also makes a testosterone enanthate, called Testobolin. It comes in 1 mL glass ampules.

Testofort Inj from Albert Davis Pakistan is commonly found on the international market. It contains 250 mg/ml of steroid in 1 mL ampules. Three ampules come packaged to each cardboard box.

Geofman Pharmaceuticals also makes a generic in Pakistan. The product contains 250 mg of steroid in each 1 mL ampule. Like Testofort, three ampules are contained in each box. Note that the lot number and expiration date are electronically printed on the bottom inside flap of the box, in addition to the proper placement on the outside.

Aburaihan makes a generic enanthate in Iran, which is becoming increasingly popular on the

black market. Note that the packaging of this product was recently updated. Countertens of this product have historically been a problem.

Balkan Pharmaceuticals (Moldova) makes the product Testosterona E. It is prepared in both 1 mL ampules and multi-dose vials.

Depo®-Testosterone (testosterone cypionate)

| Androgenic | 100 |
|-----------------------|---|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Activi | ty low |

Description:

Teastoreme cybiontic is a slow-sering injectuble store of the primary must androgen teastoreme is about primpical anabole immerse in max, and is the basis totastoreme is pictuble, totastareness and the slow start and the slow start is totastoreme injectuble, tostoreme cybionic is lightly incored by millikes for it salishty promote storing increases in macket must and strength. It is intensiting to note that while a large number of other storial composation have been runk as analyhe sizes tostoremes pictuble stored in the store of the store of the slow store include.

History:

Teaststence optimite first appeared on the U.S. dung mutted during the mid-1990's under the brand mutter Of Dope-Toststence or cybeoptrepropriori to comb com arhing to i simply Dapo-Tesstoterone). It was developed by the pharmacerical giant (Tjohn, and is sill sold to lish dy by the same company stard for team total arman (Tabaham with several Pharmace). A typically, This is a dang with fining (abola model) and has blackwish been functioned that the same transmission of the strain-strain starbar and the same transmission of the strain of another that finant distances more transmission around the strain strain strain strain strain strain of another that finant distances more transmission and the strain strain strain strain strain of another that finant distances more train strain str

Textstoree cybicute and textstoree cendular provide ceturely comparable patterns of estostorere release. Not only are physical admengas not possible in one over the other, but actual differences in plarmacoichnice patterns are hard to notice (these two dauge are for al interns and papersons instructing) irreducingually). The odd hyse difference between the two secrets to be in the area of patient conflict. Cyptoria acid is less irritating at the site of rejection than controls and constraints. The site of the other control of the other than the other size of the site of the other control observation of the structure central control of the other control of the other.

The main use of redustorence cybricuts in chical medicine has bistorically been the treatment of low androgen to each immed, although many other applications that we existed for this down as well. Dring the 1969's, for example, the drug's preserving mecommendations called for such uses as supporting bose structure mutrity, taueing mecoharing (havy) mestatud loboding and eccessive leattion in firmlis, and increasing muter firsting accomponents in the duelty. It was also been growthere are also also also components in the duelty. It was also been growthere are also also components in the duelty. It was also been growthere also provide the duelty and of the statistical exploration of the statistical exploration of the duelty of the and provide of bound a provint oversice that for to 10 used by stars prioriting bibliowed by a version of or hours of committorewise (has to to 10 used by stars prioriting). period or recorded spectrum generation (the to temporary rages that normal generatory) levels).

By the 10% the EDA had been grated much storage control over the prescription dag match, and the broad uses in which tostcores exponse to an first indicate were row bring strictly. The storage is the storage of the storage of the storage of the matching way proving to be unrelished, specially in the face of near more effective matchings, and was soon dimited from prescripting galdings. So too sus the matchings and the storage transmission of the storage of the storage matching of the storage of the storage of the storage of the storage matchings of the storage of the storage of the storage of the storage asceptible to and/order storage of the storage of the storage of the storage asceptible to and/order side effects, and storage of the storage of the storage asceptible to and/order side effects, and storage of the storage of the storage of the storage asceptible to and/order side effects, and storage of the storage of the storage asceptible to and/order side effects, and the storage of the storage of the storage asceptible to and/order storage of the stor

Today, testosterore cypionte remains exakly available on the U.S. prescription drug market, where is is PLO-approved for hormone replacement therapy in annu while conditions associated with a deficiency of endogenous testosterore, and as a secondary treatment for imperable metastatic buscast care or investment (although is not widely used I et dis parspace metastatic buscast and the second second second second second second widely. Known interement second second second second second second second and South Africa.

How Supplied:

Testosterone cypionate is available in select human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 50 mg/mk, 100 mg/mk, 125 mg/mk or 200 mg/m of steroid dissolved in oil.

Structural Characteristics:

Teststence cypiente is a multifiel form of teststence, where a carboxyle acid ester (cycheoryloppion) acid has been attached to the 17-bet hydroyd grap. Esterikd forms of teststence are less polar fran fee teststetere, and are absorbed more showly from the area of rigorito. Once in the blockmann, the ester is removed to yield bee (active) teststences. Estarikiel forms of teststetere are obsjend to prolong the window of therpartie effect blockway administration, adways for a lass forequirity test approximately data water injection. The half-field or fitted test approximately data water injection.

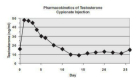


Figure 1. Pharmacokinetics of 200 mg testosterone cyplonate injection. Source: Comparison of testosterone, dihydrotestosterone, huteinizing hormone, and foliciestimulating hormone in serum after injection of testosterone enanthate or testosterone cypionate. Schulte-Beerbahl M, Nieschlag E, Fertility and Sterility 33 (1990):2013. Side Effects (Estrogenic):

Testostenore is readily arountized in the body to estradial (estrogen). The aromtase (estrogen synthesise) enzyme is responsible for this metabolism of testostenes. Estated estrogen levels can cause side effects such as increased water reterior, body fat gain, and gencomstati. Testosterone is considered a moderable synthesized in a mit-senzogen such as champlene cirate or tamosifien cirate may be necessary to prevent estrogenis side. more efficiently controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

Enorgenic side effects will occur in a doss-dependant mumer, with higher doss (above normal flarespecie) evolo (restoratore orgenicate more laboy to require the consumert use of an arti-settogen or arromass inhibitor. Since water reteriors and loss of muscle definition are common with light doss of restoratore explorates, this day is usually considered a pool for backing phases, where the added water in matching assignment with the single match size of the flat size of the size of

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for minitaing scoradary mulsenul characteristics. Encodet levels of teostores are levely on produce androgenic side effects including only skin, sense, and bodylikeiil mit growth. Men with a genetic profilopoistion first mit hous (androgenic altopace) may notice accelerated multitating. These concerned about har loss may find a mere constraints/option immunolistic profilo strained about the loss may find a mere constraints/option immunolistic profilo strained mere and the sense of the strained strained and profilo strained genetic constrained strained screening scr

In anthogon-responsive target risous such as the skin, tody, and prostate, the light relative antogonizy of tostatorene is dopedual on its rolation to only diversitatives. The The 5-alpha reductise enzyme is roporable for this metabolism of testisterion. The concurrent use of 3-abb reclustares inhibitors such as fustative of anteriority with interwith also-specific potentiation of testisterion existing the tendency of testisterion diracia potentiation of testisterion exists, lowering the tendency of testisterion diracia potentiation of testisteriority and the state of the testisterion of testisteriority analysis and the state of the state of the state of the segmentian of testisteriority analysis and the state of testisteriority and tool 3-alphar reduction enhibitor.

Side Effects (Hepatotoxicity):

Testostence does not how bepatotoxie effects, her toxicity is utilicly. One study examined the potential for hepatoxicity with high does of testostence by administering 400 mg of the lormme per day (2,800 mg per week) to a group of mile subjects. The staroid was taken only to similar higher pair, concentrations would be reached in legatic fusions and the start of the produced to significant charges in her ensures includes including serum advants, bilinduk, antime-amine-aminestence and addative bootstaness. ⁴⁵³

Side Effects (Cardiovascular):

Ambolicantogenis sterolis can huve delutricos effection on serum chocketorel. This includes a tendency to reduce IDML (apod) chocketorel voluces and increases (LL, (bad) chocketorel voluces, which may shift the HDL to LDL balance in a direction that fivors greater risk of antiroidenosis. The entities transat of an antibolicant/agonis sterolis (bala) entities and the standard standard standard standard standard standard increasing the risk of cultoroscalar disease and unconculuid inferior.

Textsteroe tends to have a much iss damatic inpact on cardiovascular risk factors than synthest strotch. Sin is do in part to is openness to metholomism be few, which allows it to have less effect on the lapping immagnent of choksterol. The aromatization of tisolasteron to stratical also helps to migring the magnite distribution of androgens on semulipids. In one study, 200 mg per veck of testosterone ester (camittatis) had a slight but not substratially significant effect on HDL choksterol alter 12 weeks, but when have with an aromatism inhibit a strong (27%) decrease was seen 225 Statis using 300 mg of testosterone ester (camittatis) per week for 20 weeks which an aromatism inhibitor demonstrated only a 15% decrease in HJL choiesterol, while at 600 mg the reduction reached 21% 453. The negative impact of aromatase inhibition should be taken into consideration before such drug is added to testosterone therapy.

Due to the positive influence of estingan on serum jupit, sumwiden cirate or echarghout containe are poterior to a summass inhibitos for those coverent with a calmosseatur heads, as they offer a partial estroganic effect in the lose. This allows them to poterinity improve high roloses and offers some of the engines clackeds of antiogeness. Who allows of 000 mg peter some effects on anticenses with a control of the enginest of the potential source of the space work of the space source of the engine term of the enginest of the enginest of the enginest of the enginestimation of the enginest of the enginestimation of

To help reduce cardiovascular strain it is advised to mnittain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydraties at all times during active AAS administration: Supplementing with Bd noil 4, grams per day) and a natural cholesterolamicoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicationgosis atensis when taken in doess sufficient to promete masck gain are expected to appresse condigenous testsotreem production. Testsotreen is the primary mark androgen, and offers strong engine feedback on endogenous testsotrene production. Testsotreen-based dange will, Beweich, here a strong effect on the hypothamic regulation of natural steroil hormones. Without the interversion of testsotreon-ssimilaring substances, testsotreen based solid reterion for multi relation of a strong secossin how their prologanization of the strong second strong secosis how their prologanization of the strong second second second second second models and thereas the strong second s

As with all ambolicathogenic storupis, it is utilisely that one will retain every yourd of new bolyowight after a cycle is concluded. The is expected must have hardbarrow from storupis (arountizing) anthogen like teststoruce cypionta, as much of the new weight gain is likely to be in the firm of water testirin, agaidy climitud after day disconstructs. An infulative of analosis and catabolic hormones daring the post-cycle recovery priord may firther create an emissioner that is utilized bible for the testirin of muck tissue. Poper ancihary dng therapy is usually recommended to help restore hormonal balance more caided, utilized by being the user retain more starks. Super-

Andre way to basen the post-cycle 'tradi' is in first replace teststeree cyclenute with a mider analotis, such a multicole decaused or methenolone emutation. The new steady would be administered abuse for one to two more methods, we multitude. The set steady of a closatorous-handling with a minimized processing of the soft macedirely understand. This practice can prove to be efficience, even framily for psychological ressortioner may visw it as simply dividing the crash into stear and hommand magas). Testsatorous-similaring diags are all typically used at the conclusion of flerapp, as desaused to emulative the stearbal charge the administration of multicoles desaused to emulative emutations.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

To text anthogon issufficiency, the prescribing galdelines for testostenore cypiorate call for a looge of 59-400 gale gale with two for weeks. Although active in the body for a longer time, testosterone cypionte is usually injected on a weekly basis for physique or preformance-entrancing purposes. The usual dosage is in the range of 200-600 mg per week, taken in cycles 6 to 12 weeks in length. This is vel is sufficient for most users to notice ecceptional gains in mack-size and strength. Teostoreo is usually incorporated into balking plases of training, when added water teartion will be of life consequence, the user more concentration with now must share definition. Some do incorporate the drug into carting cycles as well, but typically in lower does (100-2010 gave weeks) and/or whence accompanied by an anomates inhibito to loss of the solution of the dots and allow will be gard heards. Some does not be solution of the solution of the part weeks and the solution of the solution of the solution of the solution of the part water of balkers and the solution of the solution of the solution of the dots of balkers and the solution of t

White large does are gravely or a dwised, some bodyhadiers have been known to use consolvely jich doogs of this ding (1,000 mg per weck or more). This was much more common before the 1990s, when cypionate visia were usually vary cheap and easys to field. A more is better" antidie is savo jusify then proping only 320 m 10 kev via (butpe bygical price for a single injection). At doogase of 800-1000 mg per week or more, water reaction will kely courd for more of the additional weight gam has never more. Since: The practice of "trangadosing" is reflicient (not to mention poterially dargenous), especially when we take its necessarily the price of stravish body.

Administration (Women):

Testosteore cypients is mely used with women in clinical medica:. When applied, it is most often used as a secondary medication during inpresentible breest cancer, when other therapies have fiked to produce a desirable effect and suppression of ovarian function is performance-enhancing purposes due to its strong androganic nature, tendency to produce virilizing side effects, and slow-cance characteristics (ministro blocd level diffuent to control).

Availability:

Testosteone cypionter remains widely available as a prescription drug product. Its production is largely associated with American companies, although recently has been expanding into lossely regulated Asian markets that still catter to demand by hodybulkers and athtets. In reviewing some of the products and changes in the global pharmaceutical market, we have much the Biolowing observations.

Brand name testosterone cypionate (Depot-Testosterone) remains available in the United Stated from Prizer. This is a high-profile target of counterfeiters. All legitimate boxes will carry a "fh" symbol hidden on one of the top inside flaps. It will appear when placed under UV light.

Many generic forms of the dng are also produced in the U.S. mutact by manufacturers such as Watson, Sandoz, Paddock, Synere, and Bedford. All corner packaged in matiple-dose visits. Due to strict controls these products are narely diverted for flicit sale. There are also several plantmicise custom-compounding testosterone cypionate for doctors that specialize in androgen replacement thrapy.

Cypionax is available in Thailand by T.P. Drug Laboratories. It comes in 2 mL ampules containing 100 mg/mL of steroid.

Cypiobolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Thailand. Each box should carry a scratch-off security sticker, which will display a code that can be validated on the company website.

Testex Prolongatum remains available in Spain. This steroid is produced by Laboratorios Q Pharma. It is packaged in 2 mL dark glass ampules with grey silkscreen lettering. It comes in two doses, containing a total of 100 mg or 250 mg of steroid. Testex has always been a high-profile item of counterfeters.

Found in Chile is a high-dose cypionate product called ciclo-6. The product is manufactured by the firm Dag Planma, and contains 300 mg/ml of steroid in a 2 mL anpule (600 mg of cypionate in total). Balkan Pharmaceuticals (Moldova) makes the product Testosterona C. It is prepared in both 1 mL ampules and multi-dose vials.

Deposterona (testosterone blend)

| Androgenic | 100 |
|--------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activit | y moderate |
| Progestational Ac | tivity low |

Description:

Depoteron is an injectable storinary steroid preparation that contain a helend of three different totostores or sets: Each millite contains lang of totostores actuals, lang of totostores valenta, and Seng of tototores undecanoate, for a total steroid concentration of dip mill: This is currently the oxyl commercial steroid product available that contains isotosterore valenta, which is a short to medium acting easer with a luel life approximally double that of isotosterone propriorants³⁵. With the land of dows and lasthe transmission of the store of the store of the propriorants¹⁵ with the lead of dows and lasthe use of tototoxtone undecanoate i will be longer acting with more urbalanced phermocoherics.

History:

Deposteron was developed by Systex Animal Health Company several decades aga, and has been soli on the Mexican veterinary dang market size. It is used primarily to treat impotence, weakness, fatigae, and hypograndism in make breeding animals (cows, pige, carries, and sheppy, and also as a general protein-sparing amathols: Deposteronis is now solid under the Fort Dodge Animal Health label, which acquired Syntex Animal Health in the mid-1900's.

How Supplied:

Deposterona is available on the Mexican veterinary drug market.It contains 12mg of testosterone acetate, 12mg of testosterone valerate, and 36mg of testosterone undecanoate per militier or ofic) packaged in a 10 ml. multi-dose vial Twebe vials are packed in each box.

Structural Characteristics:

Depoteron contains a minure of three testostorene components, which where modified with the addition of tearboyle acid esters (accetic, subrit, and undecanoic acids) at the 17both lydroyd group. Extertified forms of testostorene are loss polar than three testostorene, and an absorbed muse solvely forms that accet of fraction. Toget, the blockstraum, the easier prolong the virulosy of therapeetic effect following administration, adjoving for a loss fracquer spectrum betwindow of therapeetic effect following administration, adjoving for a loss fracquer spectrum betwindow of therapeetic effect following administration, adjoving for a loss fracquer spectrum betwindow of therapeetic effect following administration adjover adjover of the test of test of the test of test of the test of test o

Side Effects (Estrogenic):

Testostence is readily arountized in the body to estandiol (estrogen). The arountases (estrogen synthetase) enzyme is responsible for this metabolism of testosterone. Elevated estrogen levels causes side effects such as increased water reterition, body fat gain, and genecomstain. Testosterone is considered a moderately estrogene steroid. An anti-estrogen such as chamiltonic entities or the movesment for moves the tromy estrogenet. effects. One may alternately use an aromatase inhibitor like Arimidex® (anastrozole), which more efficiently controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

Enorgenic side effects will occur in a dose-dependent memer, with higher doses (above normal therapetic level) of testostereme nee flexly to require the occurrent use of an artiestingen or anomatose inhibitor. Since water referrion and loss of matecide delinition are common with higher doses of fustatorence, tike drugs a mangle considered a poor checke for backing passes, where the added water restorion way support new strength and mack size, and help foster a stronger anabole, environment.

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for maintaining sociality mulsenul characteristics. Encodel levels of teostores are likely to produce androgenic side effects including obj skin, acm, and bodyficial line growth. Men with a genetic predposition for hirth likes (androgenic layers) may note exceeduated mike pattern hading. These concerned about hirt loss may find a neur confortible option in manufacture document, which is a compatible lise androgenic stemal. Wormen are summed of the starth as tostorence. These may include doppening of the voice, meastrul regularities, changes in kin totares, ficalital growth and clicken elargeners.

In antiogen-response target issues such as the skin, such, and prostate, the high relative antiogenizely of issues is objection on its rolation to display disolationeric (PLT). The 5-alph reductise corputs is responsible for this metabolism of tastistatives. The strength reduction of tastistative and the strength of tastistatives of tartexit with also-specific potentiation of tastistatives and instructive of tartexity of the reductive potentiation of tastistative action, however, the backness of tastistatives management effects in both metalidati with explosion tamoging metagement segments on the stoch metalidati with explosion tamoging metagement segments of tastistatives inhibits.

Side Effects (Hepatotoxicity);

Testostence does not how bepatotoxie effects, her toxicity is utilicly. One study cearning the potertial for hepatoxicity with high does of testostence by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile subjects. The staroid was taken only so that higher peak concentrations would be reached in legatic tissues produced to significant charges in liver enzyme values including serum advants, bifrabit, attinie-amire-amires, and addating propertures.⁴⁵⁰

Side Effects (Cardiovascular):

Ambrichangenis stronkis can huve delutricosa effectis on serum chesterot. This includes a tendency to reduce IDE (god) chesteron values and increase IDE. (bod) chesteroni values, which may shift the HDL to LDL bahare in a direction that fivors greater risk of ambrichanesis. The endelse impact of an analochanitogenis stread on serum high is dependint on the dose, route or a daministration (eral ws. ijectelski, type of steadi Ambrichangenis terms and the stread of the stread of the stread Ambrichangenis terms and market stread of the stread of the stread Ambrichangenis terms in my aba abasets, affort body mersen and trafforcritis, rockace endothelia relationscatar direstes and mouscilla information.

Testosterone tends to have a much less dramatic impact on cardiousscular risk factors thrus spatietic steroids: his is due inpart to is operates to metabolismity the loce, which allows it to have less effect on the lepsic imagement of chelselated. The aroundation of highs. In our study, 200 mg per week of isolostorous cardie commutation had a slight har nor statistically significant effect on HDL cholsteroil after 12 weeks, har when laken with an aroundas irribute a storeg (25%) documents was also 5% Studies using 300 mg of

. . . .

testosterone ester (eranthate) per week for 20 weeks without an aromatase inhibitor demonstrated only a 13% decrease in HDL cholesterol, while at 600 mg the reduction reached 21% ⁴⁵⁸ The negative impact of aromatase inhibition should be taken into consideration before such drug is added to testosterone therapy.

Due to the positive influence of estingan on serum lipid, numoridin cirtate or externing to acromations influinces for these coverand with a calciouscular heads, as they offer a partial estrogenic effect in the lose. This also shown to poteringly improve high profiles and offers on the longenic effects of antologens. Who does of 600 mg ore is as per work, the impact on high profile tands to be noticeable but not dramatic, making an advantage in the complexence paragroups of the profile tands to be noticeable but not dramatic, making an advantage in the complexencien paragroups of profiles materiases (for damatic effects) and the profiles and th

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of statrated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencials when taken in doess sufficient to promete make gain are expected to appresse condigences testoateree production. Testoaterees is the primary mark androgen, and offers strong engine feedback on endogenous testosterous production. Testosterone-based large via Blavoise, here a strong effect on the hypothaniar enginher production of natural steroid hermones. Without the interversion of testosterone-stimating adottances, testosterone based addit teroit to roman time in a strong testosterone in advances testosterone based based terrot to roma the strong secosisting medical advances.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men);

For bodybalding paryones, Deposterom is usually injected on at kast a weekly basis, in a dosage of 120-350 ng (2-6 nt). Doking the weekly dosage into two or more smaller applications can roduce injection volame. Cycles are generally between 6 and 12 weeks in height. This kivel is sufficient to powide noticeable gains in mascle size and strength. Testostenore drugs are ultimately very versatik, and can be combined with many other ambolicantioognic storeks does done in the desired effect.

Administration (Women):

Deposterona is not recommended for women for performance-enhancing purposes due to its strong androgenic nature, tendency to produce wilizing side effects, and slow-acting characteristics (making blood levels difficult to control).

Availability:

Deposterona is only known to be manufactured in Mexico. Because it contains such a low concentration of steroid, this product is not in high demand, and not readily diverted for illicit sale.

Dianabol® (methandrostenolone, methandienone)

| Androgenic | 40-60 |
|------------|--------|
| Anabolic | 90-210 |

| Methyltestosterone (oral |
|--|
| 17a-methyl-17b-hydroxy-1,4-androstadien-3-one 1-Dehydro- 17a-methyltestosterone |
| y moderate |
| tivity not significant |
| |

Description:

Disturbei is the most recognized trades mure for the drug methandrostruchore, also referred to so methandrossen in muny coarteriss. Methandrostrucho fer a datavisti of estabatisme, modified so that the hornnes' androganie (muscularize) Methania (andresse and and a sinabolici (fisse al battle) estabatisme). There are a solutionally quite a a distribution resolution and and a sinabolic (fisse al battle) and a sinabolic fisse and a sinabolic fisse and a sinabolic fisse and and for physica and a methodic methange and and physical parameteristic and proposes.

History:

Mehandrostrokow wa fird describel in 1955.⁴⁰⁶ It was released to the U.S. prescription dorumket in 1953, under the brand munc Dando bly Che Muramcotzica. Used developed mehandrostrokow into a medicine with the help of Dr. John Zagley who was much mehandrostrokow in the analysis of the strength of the strength of the strength and the strength and the strength of the strength of the strength and the strength and the strength of the strength of the strength and the strength and the strength of the strength of the strength and strength and the strength of the strength of the strength and the strength and the strength of the strength of the strength and the strength and the strength of the strength of the with relative dissolved in the strength of the strength of the strength of the basic chemical strength of the strength of the strength of the strength and the strength of the strength of the strength of the basic chemical strength of the strength and the strength of the streng

De Zegler's athletes were quickly making grout advancements in their competitive careers with the he of risk may accounting to report. Zegler to seemed to be very intersect. At least for a value⁴⁶⁴ Dat by the early 1960's, it was strating to hock like Diennhol Ind speeded a grout was out stand abase in competitive speets. De Zegler's comresonance in the depending on the source called for as life as 5 mg pred wy or as much as 15 mg per dry wave break gaugely givened, as althest developed Initivo nume aggressive (and poterting) damgerous) doing strategies. De Zegler's comdengies and the strate of the one and the strate of the strate of the strate with the messa of the ong, and would exercise in source of opposition to sport doring. By 1967, approximately 10 years after first inclusive data.

As early as 10%, Diambol was aleady starting to fall under scrating of the U.S. Food and Dong Administrino. That year the FDA requisted Che charging the starting of the U.S. Food and Dong Administrino. The traver are the FDA required to the starting of the starting of the traver and weakned breases. In 1097, the FDA accessfor fall Dambol weakned weakned breases. The other starting of the Dambol weak starting are started and the starting of the starting of the starting of the starting of the starting and starting of the s

By the early-80's the FDA had withdrawn its "Probably Effictive" position on the pituitarydeficient dwarfism, and continued to press Cba for more data. Sufficient clarification never came, and in 1983 Cha officially withdrew Diarabol from the U.S. market.⁴⁴³ Perhaps financial distintest thad a hand in their abandoned push to keep the drug approved. The FDA pulked all generic forms of methandrostenolone from the U.S. market in 1985, a time when most Western mations were also eliminating the drug, finding is existence to be justified mush by sports doing: Methandrostonolone is still produced todiy, but typically in nations with loose prescription drug regulations, and by companies that still prefer to cater to an underground althrite market.

How Supplied:

Methandrostenoloue is widely available in both harman and veterinary drug markets. Composition and dosage may any by country and manufacture. Methandrostenoloue was designed as an oral anabolic steroid containing 2.5 mg or 5 mg of steroid per table. (Mennatoh). Modem brands usatulg occurain 5 mg or 10 mg per table. Methandrostenoloue can also be fourd in njectable veterinary preparations. These are typically oibbased solutions that carry 25 mg/mg of steroid.

Structural Characteristics:

Mehandrostrokne is a mediad form of asstaterene. It diffus by 1 the addition of a media group a carbon 1^{-2} aph to protect the hormore during onla darinstation and 2) the introduction of a double bord between earbon 1 and 2, which networks is relative androgeneity. The results spectral data has a mark watter relative being affluid for the androgen receptor that tokisoteneous, but at the same time displays a much target parallal for advance methods and the same time to the same time of a solar atfluid data entrulymborhood in b to a very present analykes specific approx of a weaker atfluid for receptor braining. Recent statiks inver additionally confirmed that is primary mode of action modes interactions with the ediata markon methods method in agreement.

Side Effects (Estrogenic):

Mehandrostrokow is aromitacił by the body, and i a moderately estogenie aromati (opconcumaia i ośnie a ocean drużę trastrume, and may present Sed quie en tyrin so cycle (particularly when higher doses area tand). At fus same time water tentrine can sericitar and the obstal. Senskie nicholski my therefore water to keep the estigon under control with the addition of an aris-strongen such as Nobadesti' and to however, the new effective remody for estigon control. Aromitase highlow, however, can be quie more effective remody for estigon control. Aromitase highlow, however, can be quie may estimate the stroke of the stroke

It is interesting to not that mellundostarokore is structurally identical to boldence, except that is contain the added c1-abpha-methyligo grap. The first markes clear the inpact of alloring attend in such a way, as there two compounds apport to act very differently in the body. Alco dysimilarity scenar to its rule teachery for estrogenic added feeds. Exployioos (boddence indexylenit) is known to be quite mild in this regard, and uses commonly take tinding without the need to add an and science. This first added the structure of the structure compare. (a) contained the structure of structure of the structure of

Side Effects (Androgenic):

Albudg closified as an anabole stroid, androgenis side effects are still common with this substrate. This may include botts of 60 y skin, acce, and bodyficial land growth. Anabolekindrogenis stroids may also aggreate mail pattern har loss. Individuals sensible to the anabogenic effects of marfundroatoosen may find a malenia stark at a Deasto the anabogenic effects of marfundroatoosen may find a malenia stark at a Deasviding offers of anabolic indiogenic stroids. These may leaded to deargeners, marstrain impairing stroids, changes in skin structure, ficial hari growth, and change langment.

While methandrostenolone does convert to a more potent steroid via interaction with the 5alpha reductase enzyme (the same enzyme responsible for converting testosterone to dihydrotestosterone), it has an extremely low affinity to do so."" The androgenic metabolite 5-alpha dihydromethandrostenolone is produced only in trace amourse, so the relative androgeneity of methandrostenolone is not significantly afficted by finsteride or dutasteride.

Side Effects (Hepatotoxicity):

Methandostenohore is a c17-aphn adytekic compound. This alterniton protects the drug from deactication by the liver, allowing a very high percentrals of the drug entry into the bloodstream bilowing oral administration. C17-aphn adytated annobicium/togenesi seroids can be hapstonicolis. Problegad or high perconstre may result in low drugges. In more cash cycle to menzion liver factoria and concern human limits of c17-aphn adytated ateroids is commerely intendi to 65 weeks, man artifer to avoid excelling free strain.

Statics have shown that several vecels of methandosethorke administration offices minimal hepatic stress to lega it is given at a doags of 10 mg per day of below. At a doar of 15 mg per day, a majority of painters will begin to demonstrate distabed liver fanction as measured by cinical joint elevated horoscaphilic neteriorin (a methad) elevated horoscaphilic hereitorin (a methad) elevated horoscaphilic hereitorin (a methad or flagsi elevated horoscaphilic hereitorin (a methad) elevated horoscaphilic hereitorin (a methad). Severe bec complications are an egis on the prodici anture in vision servely on the prodici anture in vision servely on the methad server and an anabolication and are prological administration period.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anabelicintogravis storoids can have debatricos effects on serum checkenot. This includes a tendency to reduce IBM (apod) checkenot valass and increases ILD, (bat) checkenot valass, which may shift the IBL (apod) checkenot valass and increase (bat). (bat) checkenot articularity is increased in the IBM (apod) checkenot valass and increases dependint on the dose, route of administration (carl vs. ispectable), type of steroid ignormaliable on reasonamizable), and level of resistance on heapine matchioma-Mediatarioactoroite has a storage effect on the lequatic management of checkenot due to its isoariasi may also also weary affact them is many and and the store and the

To help reduce cardiovascular strain it is advised to mrintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencids when taken in doess sufficient to promete make gain are expected to supprese endogenous testscheme production. Methantostenolore is no exception, and is noted for its strong enfluence on the hypothalmic-philarly-testschar axis. Chicali statlas ging its 10 pr drug to resistance-taining malss for 8 wocks caused the mean plesm testsetsence level to fill by 69%.⁴⁴⁶ Without the intervetion of teststerenstrainding substances, testsetscheme is drug and altern to romal while i.1 A emotion for a scension. Note that prolonged hypogenuldworphic hypogenulsm can develop secondary to storial about, encodentium medical intervetion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral anabolic steroid with field may decrease its bioavailability.⁴⁶⁰ This is caused by the fat-soluble nature of steroid hormones, which can allow cores of the draw to discole with undirested distance for exclusion from the observations. and/w source of the study to thesome with intrageneed therapy and, rectaining and associption norm the gastrointestimal tract. For maximum utilization, this steroid should be taken on an empty storated.

Administration (Men):

The original prescripting galakties for Diambol called for a dudy dotage of 5 mg. This was to be administered on an intermedier basis, with the drug taken for no route than 6 concentration of the structure of

Diambel stacks well with a verity of other storick. It is noted to mit particularly well with the mill analob Dozen-Dandolini, for exempli. Foghert one can expect exception limelok and strength gains, with side effects not much worse fram one would expect from Diambel ohars. For effect merses, i how generity instructions of the strandynger, however, side effects should be mere proteomed. Gains would be protoranced as well which usually makes such an endeavor worthwhole to the user. As discussed endring, anclary drags can be added to rockner the side effects associated world his hird of cycle.

The hulf-fiel of Diambol is only about 3 to 5 hours. A single daily dosage schedule will produce a sweep (bood lock) with any and down flowdgout to day (F) uses well, lockies, man and a constraints of the start block. This is the start of the start discover methics in the born to chick them and to y to regulate the constraints in your block. This, however, well produce is start produced block block well furth first black wells at an acceledal at cocce, so there may be a trad-of with this option. Both options work first, bar ancededal at cocks, and the start of the start

Administration (Women):

Beig moderately androgarie, Danabol is really only a popular steroid with men. When used by wornen, storey dividization syntportus are possible. Some do experiment with it, however, and often ful kw doeses (2.-55 mg) of this steroid quite effective for new maccle growth. Studies have demonstrated that a majority of wornen will motice accure, which is indicative of antrogaratic, at a dosage of only 10 mg per day. Children are likely to notice willizing effects with as filte as 25 m gpt erday.

Availability:

Methandrostenolone remains readily available as a pharmaceutical product, though its supply is largely isolated to less regulated markets in Asia and Eastern Europe. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following observations.

British Dispersary produces Anabol tables in Thaliand. This product cornes in three strengths, 5 mg, 10 mg, and 15 mg, 41 products corne in bothes (100, 200, 500, and/or 1000 depending on the doss), All tables themselses are inprirted with the company's studee enhibm, and will sup in clock: The prival tables are 5 mg, they collow 10, ang and blue 15 mg. The company also uses a holographic sticker on all products to deter conterficing, though this fainture has been copied with high accuracy.

March Pharmaceuticals in Thaland manufacturers Danabol DS. These small blue heartshaped tablets come in bottles of 500. The company recently instituted a small circular holographic sticker to deter counterfeiting. The logo is also embedded into the label in a metal foil. Dronabol DS is remains available in Trailand by Bangkok Lab & Cosmetic. This product comes in a 10 mg tablet strength, and is prepared in bottles of 500 and 1,000 tablets each.

Methandon is also found in Thailand, made by Acdhon. This product comes as a 5 mg tablet, and is packaged in plastic tubs of 1,000 tablets each.

Generic "Russian D-Bof" (METAHAPOCTEHOROH) is no longer in production. All products bearing this label should be considered counterfeit.

Naposim (Romania) is still in production, now under the joint Teraphia Ranbaxy label. It is sold in howes of 20 tablets, which are separated into two foil/plastic bisters of 10 each. The tablets carry a triangle stamp on one side. This product has been the subject of widespread counterfeiring in the past.

Metanabol from Jelfa (Poland) is still in production, though its appearance has changed. The new packaging reflects a more modern red gradient look. The enclosed foil and plastic strip is also much wider than it was previously.

The generic from Formula Magistral (Argentina) is also commonly located on the international market, especially throughout North and South America. The product comes loose in bottles, although now also carries a holographic sticker to deter counterfixing.

Landerlan in Paraguay makes a generic methandrostenolone. It comes in a 10 mg tablet dosage. The product is packaged in bottles of 100 tablets each.

Balkan Pharmaceuticals (Moldova) makes the product Danabol. It is prepared in both 10 mg and 50 mg tablets, with 20 tablets contained in each foil and plastic strip.

Drive® (boldenone/methylandrostenediol blend)

| Androgenic | |
|------------|--|
| Anabolic | |
| Standard | |

Chemical Names

Description:

Dute is an Australian injectable veterinary stereid preparation that contains a bland of methanical dipopointen and budderen enclose/tent. The too stereoids are present in a dose of 25 mgril. and 20 mgril. especticely, for a tool stereid concentration of 55 mgril. Boldence uncloselune as high common steard metra study beinfield with the preparation Explosited. Methandrish dipopointate, however, is very analy seen on the U.S. accomparing by an order that analysis and the start of the start of the start of secondary and the start of the start of the start of the start of the start result is an orderately and orgenization being the start of the start of the start of the start of the start result or operative nucleus the blackwerk the result is a moderately and orgenization black that includes to produce neuble mack must and sterent and its usable whole or cension the result is an order result.

History:

 carry a varry of unsain steroots and cont mit-component sterou bencis. Lrike is pormaps the most well-known of these products. Being that it is nother the most cell-known of preparation nor the most effective, however, much of its popularity is filely due to its wellcound trade rame and carry salss its showy. Drive remains on the Australian mutket today, although tight controls and its reduirely buy per-milliter steroid concentration make diversion for tarbitistic meta bis scormon than it was many warsa ago.

How Supplied:

Drive is available on the Australian veterinary drug market. It contains 55 mg/mL of steroid in oil in a 10 mL vial.

Structural Characteristics:

For a more comprehensive discussion of the individual steroids boldenone undecylenate and methandriol dipropionate, refer to their respective profiles.

Side Effects (Estrogenic):

Methylandsentidi is not directly anomized by the body, alhough one of its torous metholics is methylandsentorene, which can aromize. Methylandsentextidi is also believed to have some interest entrogenic activity.⁴⁵⁰ Combined with bolkenese, which also anomizes. These is considered at moderable entrogenic second, Cynsconnuclis in possible also become issues, again depending on door. Somisive infinitalist my need to add an antiettogen and an Novalaevel to minimum celled side effects.

Side Effects (Androgenic):

Albedge clessified as an anabolic steroid preparation, androgenic side effects are still common with this substance. This may include boats of objekti, ance, and body/heal hair growth. Anabolic/androgenic steroids may also agarvate male pattern hair loss. Women are wonted of the potential wizzlegifeffects of anabolic/androgenic steroids. These deepening of the voice, menstrual irregularities, changes in skin texture, ficial hair growth, and clonal enlargement.

Side Effects (Hepatotoxicity):

Medylankostenciól is a (17-ajeha alsolated component. This alteration protects the drug from deactivation by the loca, alavoig as vary high procetage of the drug entry into the bloodstream following on al administration. (17-ajeha alsolated anabolicanhogenis steroids in the hightatoxic, Probaged of high exposure my scali hi for drugme, In nare instances life/stratantegi dydarization my doelway, hi sa absolute to visi a replexicum periodically daring is commonly hindin of 66 words, in an effort to avoid scalaring her visian in gravitation form of the drug may present sightly has strain on the her by rowing the first pass methodsminent and the drug and scalar strain the strate strateging and the strates and scalar strateging alsolgen and gravitational strates and the strates and scalar strateging alsolgen and gravitation strates and strates and scalar strate

Side Effects (Cardiovascular):

Ambelichtrangenis stronkis en ihnve deiteritose effects on serum chesterot. This includes a tendency to reduce IBL (good) chesteron values and increase IBL (bod) chesteroni values, which may shift the HDL to LDL bahare in a direction that fivors greater risk of attrividensis. The reduce impact of an anabelicandrogenis extraol on serum high is dependint on the dose, route of administration (eral vs. injectable), type of steroid (incomitable to reaso-annutable), and level of resistance to hepatic raticbolants Methylandrosancelal has a same gletic on the hepatic mangament of checkered due to its Ambelicandrogenis extending may and associated and the same and region entry anabelicandrogenis extending may and support kit warricair hypermedya. Ja poteridajle researcing the risk of curritoscate af messes and mesonicality and storetion.

To help reduce cardiovascular strain it is advised to minimian an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardorlydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterol/antioxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

All araboléandrogenie steoidis when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalating substances, testosterone levels should return to normal within 1-4 months of dng secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Drive has not been approved for use in harmas. Prescribing gaidelines are unwability, Typical doising schedule for physique or performance-enteming purposes would be in the marge of 220 mg (4mL) to 440 mg (8mL) per work, a level that should provide quality lean muss gain without strong beharing to body fat tentrion. Due to the high rejection volume and fastacting nature of methandrial discoptionate, the total weekly dosage is commonly divided into 2-3 smaller applications.

Administration (Women):

Drive has not been approved for use in humans. Prescribing guidelines are unavailable. Drugs containing methylandrostenediol are generally not recommended for women for physique- or performance-enhancing purposes due to its androgenic mature and tendency to produce virilizing side effects.

Availability:

Drive remains in production, though legitimate products are now rarely diverted to the black market given the heightened controls over anabolic steroids in Australia.

Durabolin® (nandrolone phenylpropionate)

| Androgenic | 37 |
|--------------------|--|
| Anabolic | 125 |
| Standard | Testosterone |
| Chemical Names | 19-norandrost-4-en-3-one-17beta-ol 17beta-hydroxy-estr-4-en-3-one |
| Estrogenic Activit | y low |
| Progestational Ac | tivity moderate |

Description:

Natiokene plensphopicente is an ispectible form of the anabole strend mathema. The properties of this day are atrikingy similar to those of Dezo-Drahodik, which uses the shower acting drug matheokene decamoate. The primary difference between these two perpantions in the speed of which mathematics is related in the biddo Wilki mathematics decamote provides a release of mathematic for only docat a vace. In clinical particular, Decas Damoin can thus be igned new every 2 or 3 vaces, while Damboin's usually interchangular. Decas Damboin is valued by attacking and Decay Damboin terriburgshilt. Decas Damboin is valued by attacking and Decay Damboin is a shifting to the Deca Damboin Damboin's valued by attacking and Decay Damboin is shifting to promote strength and has manche muss gains without significant extrageries of the Strength and the Decay.

History:

Natholese phenphropicente was for described in 1975⁴⁷¹. It becares a prescription medication should act will be the intermittent phenrometerials giver Organn (owe Merck/MSD) under the brand runne Daraboltt. When first introduced to the Lindea States, initiated uses of number phenphropicnet related pre-and protogenite learn mass reterior, notopoonsis, advanced breast cancer, weight loss due to considerence or disease, genitric states (general weaksson at Hingh), berns, severe stramm, alsons, adjurced therapy with contain firms of anomis, and selective cause of growth and development therapy with contain firms of anomis, and selective cause of growth and the selective and phenomena. The low serves coss significantly manword. Moreing frownich the drug was multiply being indicated for the transment of advanced metastic breast cancer, and as adjurct therapy for the treatment of solit and pos-

Databolis was i key focas of Organovis muchang effers cody for well isss than a decade following is relaxes. Once Daca-Databolis wis mitudeed dring the 1096/s, this shorteracting compared muchanisms and the site of the term of the term of the term organetic and the term of the data on a rice metric of the term of the term of the term of the term of the data on a rice metric of the term of the term of the term of the term of the data on a rice metric of the term of the data on a rice metric of the term of the data on a rice metric of the term of the on versions of the data of the term of t

How Supplied:

Nandrolone phenylpropionate is available in select human drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 mg/mL or 50 mg/mL of steroid dissolved in oil.

Structural Characteristics:

Nandoken phenykropisente is a modifiel ferm of ramforkom, where a carboyck scit scit (roptopice phony elser) has been attended to let 17-bet hypotry gapes. Earticle storiks are uses polar than free stareki, and are absorbed more slowly from the area of injection. Howe in the blochteamt, the sets is removed to yield in (excitor) narrobices. Easterfield starekis are designed to probring the variator of therapeatie effect following (uncertificity) startics. Nandores: phenytheprotection genesis areas of the reasterifield startics. Nandores: phenytheprotection genesis areas of the related startic startic startics. The startic startic startic startics are startic relation startic startics. The startic startic startic startic startics are startic relation startics. The startic startic startic startic startics are startic startics where the startic startic startic startic startics. The startic startic startics are startic relation startic startics. The startic startic startic startic startics are startic startics and the startic startic startic startic startics are startic startic startics. The startic startics are startic startics where the startic startic startic startic startics are startic startics. The startic startic startics are startic startic startics are startic startics are startics are startic are startic startic startics are startic startics are startice are startics are startic startices are startics are sta

Side Effects (Estrogenic):

Nandoken hus a low tendex- for entrogen convestion, estimated to be only about 20% of the serve with testorerow¹⁷. This is because while the lore can covert matchoose to estatidal, in other more active sites of stored arountations such as adjocent tissue matchoose in the process.¹⁷ Consequently, estimation, related aids effects of the motion of the store restored, hold (in gain, and groccouncil, An anti-storegates such as charging criticat or introduce in the necessary to prevent extraports and the occur. One may advantable are amontains entithed like Aircraticate initiatives can be quite different occurs of the store and entities of the store store and the store and stored initiation in advectoring. In these stores with the store and theory in the store of the store in the store of the store of the store store of the st

It is of note that nandrokne has some activity as a progestin in the body.⁴²⁴ Although progestorone is a c-19 steroid, removal of this group as in 19-norprogestorone creates a hormore with greater binding affinity for its corresponding receiptor. Starting this trait, many 19-nor anabolic steroids are shown to have some affinity for the progesterone receiptor as welt^{***} The sake effects associated with progestrone are smiler to those of estrogenincluding negative feedback inhibition of teststorence production and enhanced net not ful storage. Progestris also anymert the strankaboy effect of estrogens on mammus tissue growth. There approach has a storage systemy between these two hommouss here, such that the storage of the storage systemy between these two hommouss here, such that The use of an anti-estrogen, which tables the estrogenic component of this disorder, is often sufficient to minigue perconnustic caused by numboline.

Side Effects (Androgenic):

Altrody closefield as an analysic steroil, androgenic size effects are still possible with first substance, opercively with higher doess. This may include boots of op skin, are, and body/fiscial ima growth. Analysic/androgenic steroids may also aggroates mile pattern hair hosts. Wornen are sustened for the potenti withring effects of analysic/androgenic steroids. These may include a deepening of the voice, mentiumal imaganities, changes in sink instance, fincal hair growth, and closed enlangment. Nandenkon is a steroid with edited by androgenic activity relative to its issue-balling actions, making the facehold for strong tostscarrate, methanismismic, or discourteone. It is also important to point of mildue to its mill androgenic nature and ability to suppress enclopenous toxifortement.

Note that in androgen-responsive target tissues such as the skin, scalp, and prostate, the relative androgenicity of randrolone is reduced by its reduction to dihydronandrolone (DHN) 476477

The 5-alpha reductise enzyme is responsible for this metabolism of randrolone. The concurrent use of a 5-alpha reductuse inhibitor such as finasteride or dutasteride will interfere with site-specific reduction of randrolone action, considerably increasing the tendency of randrolone to produce antrogenic side effects. Reductuse inhibitors should be avoided with randrolone filow androgenicity is desired.

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabele and angenesis strends can have deleterious effects on serum chocketerd. This includes a network to refere the [L (apod) chocketer) datas and increase [L (bad) chocketer) values, which may diff the [ER]. In LR halters in a direction that fixeos gaues risk of dependent on the does, route of administration (call w, rejectively, hype of strends dependent on the does, route of administration (call w, rejective), hype of strends diministring (60) regular of machine dependent on the does does does and diministration (20) regular diministration (call w, rejective networks), hype of strends does not administration (call w, rejective data and the strends dependent on the does not administration (call w, rejective) and the strends evolution of the strends dependent of the strends dependent of the strends evolution of the strends dependent dependent of the strends dependent of a highly strengt regulation (Call R). The strends dependent dependent dependent dependent also adversely effect block pressure and indijectively, reduce entrothelial relation, and does and maccellul inferiors.

To help reduce cardiovascular stania it is advised to maintain an active cardiovascular exercise program and minimize the intake of starated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4 grams per day) and a natural cholsterolarinoxidant formali such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are owneeted to curvences ordonerous testoctorone norduction. Studies administering 100 mg equecte or suppress transgatement memorative protocates, strutter animemetry (10% ing spectrum of multicologie phelphypoipnical demonstration 1 ang anglerossis of serious testosterore following a sight spectrum. Testosterore levels declared to approximately 30% of railla levely dogs a dark dag administration and stopid appressions formation days. Regular use is expected to significantly lengthen the endogeness hormone recovery approximate of stostaneous polithois during placetopolithy and the stopic structure and approximate of stostaneous polithois during placetopolithy and serior testosteronesistences, tostosterone beek should return to neutril with 26 Arms for days account Note that prolonged hypogenesidence beek should return to neutril with 26 Arms for days account hypogenesistic instrustorial intervention of steroid advances. The steroid advance intervention of steroid advances, tostosteronebeshe, neucostating multicida intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

For general anabolic effects, early prescribing galdenies recommend a dosage 0725-50 mg pp week for 12 weeks. The usual dosage for physique – or performance-enhancing purposes is in the range of 200-400 mg per week, taken in cycles 8 to 12 weeks in length. This is level is addicate for most users to notice measurable gains in last macher mass and strength. Note that date to the distancing rather of the phenylpropicate ester, the weekly dosage is usually subdivide into 2 separate applications spaced evenly apart.

Administration (Women):

For general narbolic effects, early preserving guiddens recommend a dosage of 25:50 mg per week for 12 weeks. When used for physicape or performance-charaing praysess, a dosage of 50 mg per week (given in a single weekly nigotion) is most common, taken for cycle kingt 4 of to week. [Hgerd noses or loss granged mainton of the are disconraped due to potential for antiogenic side effects. Although only sightly antiogenic womm are coascinarily conformed with witzing suppress when taking this compared. Should witzing side effects become a concern, matchone phenytropicnets should be discontinued immetable to hip prevent a permanet requerance.

Availability:

Nandrolone phenylpropionate has declined extensively as a pharmaceutical product. Given its short action, and the limited use of its longer-acting cousin randrolone decanoate in clinical medicine, there are very few (if any) unique applications remaining for this drug. Thus, there is lift justification for its continued production.

Brand name Durabolin appears to be unavailable in all markets worldwide.

A small number of generic and brand name products remain in less regulated markets (mainly in Asia), due to continued demand by athletes and bodybuilders.

Superanabolon from Spofa in the Czech Republic is also still in manufacture. It contains only 25 mg of steroid per 1 mL ampule, which makes it in relatively low demand among athletes.

Iran Hormone (Iran) makes a 25 mg/mL generic nandrolone phenylpropionate in 1 mL ampules. Counterfeits are not known to be a problem.

Dynabol® (nandrolone cypionate)

| Androgenic | 37 |
|------------|--------------|
| Anabolic | 125 |
| Standard | Testosterone |

Chemical Names 19-norandrost-4-en-3-one-17beta-ol

| 17beta-hyd | 17beta-hydroxy-estr-4-en-3-one | |
|-------------------------|--------------------------------|--|
| Estrogenic Activity | low | |
| Progestational Activity | moderate | |

Description:

Nandokee cypionus is an nijechsk form of the anabok steroid metholome. This exter provides a pattern of hormore relases virtually derical to that of teststence cypionus, with peak levels of drug beige noted approximately 2-48 hours after administration, and a substratial hormore relases satistated for hour vocks. In this case the active hormore interloome, which is a moderately storeg anabolic steroid that carries mild storgen: and mildgenic properties. This product is essentibly default in oncel affect to Darmage, which and no her accounted by a low level of rails efficience that marked boxes cypione to be as a fast-storing compound to your users. Otherwise, there is no discernable difference horsen endpoints of applies to be associated occurated in virtual of explose users. Otherwise, there is no discernable difference horsen endpoint of applies of the data where the occurrents, and markednew cipiont and produce molecular docurrent in virtual of explose.

History:

Natiobare cypionts was first dovelped daring the 1960%. It was sold for a brief rime as a larma-use pipmenetical, under such hann framess as Anaba Dopo-Nottsteinen, Nottsteinent, and Starveinish, Sach preparations did not hat, however, and in recent appearates. The correspondence of the star of the sta

The discortined Jaros products were qickly transformed to SYD Group in Australa, assuring they sould not be completely elimited from contracts. They were subsequently reintroduced to mutcher in 2002, under the rames. Analosic INA and Anabols NA, respectively. The new runns much loss effectives to the formal Australa, Saka instandera a high-document of the products. SYD Group had also intraduced a high-document of the products. SYD Group had also intraduced a bigh-document of the structure of the structure mutcher, but the product has size been withflammen. This time the product was discontinued open Mexican activation of the Australia Australia, and analogical and also encounted on the U.S. Group, the Anabols E.W and Anabols NA products remain available on the U.S. Group, the Anabols E.W and Anabols NA products.

How Supplied:

Nandrolone cypionate is available on the Australian veterinary drug markets. It is supplied as 50 mg/mL of steroid dissolved in oil, in a 10 mL vial.

Structural Characteristics:

Nandokov cypionate is a molified form of nandokov, where a carbodyke acid ester (cyclepertyphoprine) acid hun show nathoda to he 17-both hybrody group. Estreffel storoki are loss polar fran free staroki, and are absorbed more skowly from the area of frigorich. These in the bodokname, the sees its moved to bykit the (action) randokova. Esterified atensik are designed to prolong the winkow of finerpartic effect following adminimum, always for a loss-foquer priving the visition of the action of antibacture. The star fiber of the set of the star fiber of the star of the star of the star 24-48 huns fiberwing deep intramescalar injection, and statum is substarial release of humone for approximately 2 works.

Side Effects (Estrogenic):

una secti wani teosnorue. — inis n occane wan eur ner can conten manuroux to estaboli in dem mer achie skei of storial aromatinios and si adipote lisso matoloxes is far loss open to this process.⁴² Concequerdl, estrogen-teiled side effects are annal lover comen with this dig man with istasticture. Bestematical estrogen levels sill be noted with higher dosing, however, and my cane side effects such as increased water retorious, hold fig and, and genoremedia, ha anti-steaperious. Bestemating and the open chaine on maximum chain my be necessary to present estrogent side effects filler open chaine on aromating chainer and the strokess. The strokess and the scheme discussion of the strokess of the synthesis A container site of the synthesis expension in comparison to artif-estrogers, however, and my also have negative effects on blood lipids.

It is of need that matchetee has some activity as a progestin in the body.⁴⁰ Although progestrence is a < 0.9 stored, neuroal of this group as in 19-mongatence reactes a harmone with greater briefing affithely for its corresponding receptor. Sharing this truit, many 19-ner anabolic stored is an elow no have some affithity for the progestrence receptor as well⁴⁰. The side effects associated with progestrence are similar to those of estimation stored and the size of the site of the site of the site of the site of the storegard. These appears to be a storegy group between these two hourses here, and have profind. These appears to be a storegy group between these two hourses here, and have profind. The site of the

Side Effects (Androgenic):

Although costindi as an analysis steroid, androganic size effects are still possible with this substance, opercival with higher dows. This may include boots of ody six, area, can bootly/facial lari growth. Analysic/androganic steroids may also aggrowte make pattern har incose. Weren are sustained for the potenti withing effects of analysic/androganic steroids. These may include a deepening of the voice, meanstand imagainties, changes in sink texture, facilitati and analysis. Analysis and analysis and analysis and analysis and analogonic side discing entingenative. Nandrokse is a steroid with editoriely businet discing and analysis. The steroid steroid

Note that in antrogen-response trapt issues such as the skin, scab, and prostate, the relative antrogenicy of matrolones is related by its relation to displormative (DBN) $\frac{600}{200}$ The 5-alpha relatance alteriation to displormative matching in the scalar relation of the scalar relation of the scalar scalar matching the tradeway of matholese the produce and mathematical messaging the tradeway of matholese it to produce and mathomatical scalar scalar scalar scalar scalar scalar scalar methods with the scalar scalar scalar scalar scalar scalar scalar scalar methods and the scalar sc

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabolic mining are stored as an more debetrious effects on serum chocketerd. This includes a technery to reduce IRL (good) relabered values and increases (Lb, (dus) chocketord values, which may drift the HDL to LDL balance in a direction that favors gener risk of a minimization of the store of the matchical magnetic associations are may high is (normational box) are more another than the matchical magnetic association in sum (high is disministing) on goin of markbork chargeness proveds for 10 week domestical a 20% reduction HDL chocketone levels.⁴⁴⁷ This suppression is sightly generate main the reported with mergal does of testoreme earnithme, and is in agreement of under a dusles showing a slightly stronger negative impact on HDLDL. In this with matchone desnote also compando to instituence optimule.⁴⁵⁶ Manufactore jeticable, howing, wheek fully labore, we doeld ill labore. significantly weaker impact on serum lipids than c-17 alpha alkylated agents. Anaboliziandrogenic steroids may also adversely affect blood pressure and triglycerides, reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and moccardial influencion.

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, advised cardohydrates at all times during active AAS administration. Supplementing with shi oki (4) grams per day) and a natural cholsterol/antioxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al analysical adjust setup is setup in the setup in the setup in the setup in the sequence operation to approxe molgrounds to statement production. For setup 46 or comparison, statisationisating 100 mg per veek of markolene decanota for 6 weeks have demonstrated an approxime 5% relation in sement statements based based that the programmed analysis approxime 5% relation in second statements based based per setup in the approxime 5% relation is second statements based based per setup in the approxime 5% relation is second statements operations of statements operations and analysis of the statement of the statement of the statement of the statement intervention of testocherosis distatements operations, biostochero tests should return to normal within 2-6 months of than greeosisn. Note that prolonged hypograndstomelybe hypograndism and operative spectration that approximate the statement of the

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

When used for physica- or performance-enturcing purposes, a door of 200-400 mg per work is most communication in cycles 50 210 weeks is night. The work is subject of the accompanio 24 year how level of estrogenic and androgenic activity. Although higher doors (430-640 mg) may produce a storager analysis direction and androgenic activity. Although higher doors is which this ding as fast of 150 mg/s1, 30-36 show -000 mg are not commonly applied. Instead, the ding is often stacked with andrest gasts, usually an antrogen starks in an antrodome, a song and with the mather agest, usually an antrogen stark is an antrodome, as one direction with produce and molegonicity, such as relatively box commentions. For all starks with produce and molegonicity, such as relatively starks and a storage effect on starks with produce and molegonicity, such as relatively and and and a storage effect on starks with produce and molegonicity, such as relatively and and and and a storage effect on starks with produce and molegonicity and as a relatively storage and and and a storage effect on storage with plant produce and the storage storage storage storage and and a storage effect on storage stor

Administration (Women):

When used for physique or performance-enhancing purposes, a dosage of 50 mg per voceis not corrum. Altopic phys sight participation of the start of the start of the start virilation symptoms when taking fine compound. Should viriling aske effects become a corean, manebook explorate absolut bit discontinued immediate ho help percent their corean start of the start Databoling right be considered a safer (more correlabile) option. This days active for only several doxy, neutro percent gate withhous dim at first end.

Availability:

The only remaining pure randrolone cypionate product is Anabolic DN from Australia, produced only in a 50 mg/mL concentration. It comes in the form of a 10 mL vial, which is contained in an orange tube.

Dynabolon® (nandrolone undecanoate)

| Androgenic | 37 |
|------------|--------------|
| Anabolic | 125 |
| Charles I. | T . 1 |

| Standard | lestosterone |
|--------------------|---|
| Chemical Names | 19-norandrost-4-en-3-one-17beta-o 17beta-hydroxy-estr-4-en-3-one |
| Estrogenic Activit | / low |
| Progestational Ac | tivity moderate |

Description:

Nandroken urkeannote is an injectible form of the ambods stroid mutholone. The ester applied here is one control motion the motion documoting, and consequently form a vary lightly longer-storing drug deposite at the sile of injection. With proper amotion paid to formation the storing of the sile of the sile of the sile of the sile of the formation of the storing of the sile of the sile of the sile of the sile of the mutholes mutholes and the sile of the sile of the sile of the sile of the mutholes mutholes and the sile of the sile of the sile of the sile of the mutholes mutholes and the sile of the mutholes mutholes and bounds. Also, the sile of the mutholes mutholes and bounds. Also, the sile of the sile

History:

Natiobare undecancel was developed daring the 1960%, and was subsequently odd as Dpatholon in 180 (Crinos) and France (Harmark), and as Psycholon in Germany (Harmark). The linkin product was moved to the new formassiste likely jeans later, but enterind the origin Davaloka tanda muras. Upsholsno was aground priodated for one in about the control costoporosis, including the numerical or anti-ogen-sensite populations and an worth and the dedshy. Natoritorius machemolis spectra in the subset of a silvy needs, yet in spite of fits, the france known commercial preparations data tasks and the respective prescription dang markers. Psycholasin france fitting Francescher in high sories discussion of the start of Davaloban from Frances fully and the effective data start of the start of the start of the start of Francescher in high sories discussion of the start of the start of the start francescher in the start discussion of the start of the start of the start francescher in the start discussion of the start francescher in the start discussion of the start of the start of the start of the start francescher in the start discussion of the start of the start of the start of the start francescher in the start discussion of the start of the start of the start of the start francescher in the start discussion in the start of th

How Supplied:

Nandrolone undecanoate is no longer available as a prescription drug product. When manufactured, it was supplied at a concentration of 80.5 mg/mL dissolved in oil and seaked in a 1 mL armyte. Each armpute provided the equivalent of 50 mg of nandrolone base.

Structural Characteristics:

Nandroken urdecanote is a molified fem of nandroken, where a carboxyje acid ester (undecanote acid) has been attached to be 17-ben laphotog group. Esterified streads are less polar fram free streads, and are absorbed more skwy from the area of spectron. Once in the hoodstream (he est's resrvord to yield for (acids) andromotic. Esterified streads are designed to prokeng the window of therapeatic effect flowing administration, allowing for a loss "opent pricines schedule compared to ispections of the (unsetting) streads. The fore undecanote is designed to proke a slow release of randrokone for up to 3 to 4 weeks following injection.

Side Effects (Estrogenic):

Natiobles in a low incluse, yet entropy conversion, estimated to be only about 20% of the sens with tostochronove⁴⁰. This is because while the force can convert matchouse to estandic) in other more active sites of staroid aromitation such as adjacone tissue matchouse in fits sequence that the staroid staronization starbar and the entropy of the processor. The staroid staronization starbar and the tentical with the processing starbar and the staronization startest staronization of the staroid staronization starbar and the startest staronization starbar and generoments. An anti-setteneous starbar and the three processing startest staronization starbar and the staronization starbar and the starbar and the starbar and starbar and generoments. An anti-setteneous starbar and the starbar and the starbar and starbar and generoments and anti-starbar and the starbar and the starbar and starbar and generoments and the starbar and the starbar and the starstarbar and starbar and generoments and the starbar and the starbar and the starbar and starbar and an anomase stable for Rock reference of more starbar and the starb controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

It is of next that matchetee has some activity as a properties in the body.⁴⁰⁰ Although progenetron is a >10 strend, nervoid of this group as in 10-mongenetrome creates a hormone with greater briding affittily of its corresponding receptor. Sharing this trait, many 10-ner anabole storesholds meldows and matching the strength progesteme receptor as word ⁴⁰⁴. The side effects associated with progesteme are similar to these of estimation storages and the strength progesteme are similar to these of estimation storages. The side effects associated with progesteme integrate and enhanced rate of the storage. Progestim also anyone the stratiationy effect of estimates on mummary time agreementian ingle resources with the hole progestim, where excession estimation holes. The use of an anti-estimgan, which inhits the estinguistic componer of this disorder, is often sufficient to mitigge encounties and any transformation.

Side Effects (Androgenic):

Although costified as an analysis steroid, androgenic site effects are still possible with this substance, opercively with higher dows. This may include boots of op skin, acce, and body/ficial luir growth. Antabolicandospris steroids may also aggroates make pattern har inso. Wormen are sustened for the potenti withing effects of analysis. Indexes in a state state of the state of the state in the state of the state of

Note that in antrogen-responsive target issues such as the skin, scab, and prostate, the entities antrogenicy of matrokone is exclused by its reduction to diffutorantilokone (DRN) 40,9 the S-alpha reductase enzyme is responsible for this matholism matholism. The concurrent use of a S-alpha reductase influences and the matholism is a strategied of the star of the strategies of the st

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabolic miningenes steroids can huve delutricos effection on errarm chockenerd. This includes a nationey to related TBL (good) chokkened values and increases (DL (hold) chokkened anticirculocosis). The rahibes impact of an anabolic indications in the start of the

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish ols (4 errors nor dwo) and a natural cholesterollaritoxidant formula such as Lini (Stabil or a product with comparable ingredients is also recommended.

product with comparable ingredients is also recommende

Side Effects (Testosterone Suppression):

Al anabolizationgois actuals when taken in doess sufficient to promote mack gain are expected to apprece endogrous testostroner production. For site of comparion, studies administring [10 mg per veek of markohen decanote for works have demonstrated an approvinte 5% endoction is neural testostrone levels dring therpare, Va a dosage of 200 mg per week, this reduction reached $70\%^{50\%}$ li is bieleved that the prograstization alcohy of markohen neibby combines to the suppression of testostrates or patients dring therapy, which can be marked in spite of a low tendency for estogen conversion.²⁰⁰ Without the intervention of testostrone-similarity assistance, testostrone to well solid without to normal within 2-6 months of the appression. Note that prolongal hypograndtompice hypograndism and orderly secondity to structure disk heat, necessating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Nandroken undecannote was used chrisally at a dose of 11 ampek every 1 to 2 weeks. A totel of 3 to 6 amplex were useft far a given for sevek period of theory, When used for physique- or performance-intensity parsons, a dose of 30 4 ampeks (2415 to 322mg) per work is not corrora lation in cycles at 10 weeks in height. This level is afficient for most users to notee masanable gains in lam mack muss and strength, which should be accompared by a jour lose of or stopparts and androgen activity. They does (0476-060) mg per weeks) will imput a stronger analysic effect, for an he difficult given the relativity ages with often amplekondingsing strength for a stronger offect. Geno in poperties, if come is properties, it is seense to 4 well for holio balking and cutting purposes, and can reasonably replace Deci-Darabolin nack-tocks.

Administration (Women);

Natiohore unlearnote vus used chirally ta dose of 1 ampté every 1 to 2 weeks. A toti of 3 o 6 amptés sever used far agione social programment and the severe social of a severe social programment photogies or performance-entineting programment, a dosage of 1 amptés (005 mg every 10 worsen are occessioned) confinent of whit whitein sorphone winder this grip is composed. Should writing side effects become a concern the drug book to the discontinual immediately to help prover the permitted approxamely. Also a sufficient period with discontinuel softent are accessible to a several days grant and the softent are softent and softent are accessible to a several days grant and the softent are softent and interaction.

Availability:

Nandrolone undecanoate is no longer available as a prescription drug product. Some underground preparations are, however, known to exist.

Equipoise[®] (boldenone undecylenate)

| Androgenic | 50 |
|---------------------|--|
| Anabolic | 100 |
| Standard | Testosterone |
| Chemical Names | 1,4-androstadiene-3-one,17beta-ol 1-dehydrotestosterone |
| Estrogenic Activity | low |

Description:

Bolkence undexpleme is an ispeciable veterinry across that exhibits strong mathok and molecularly and ongoing properties. The undexplement sect extends the acriticly of the days grandy (the undexplement setter is only one carbon nions longer than discussion), so that and an advaptice properties of this days are grandy approxised by whites, who spaceful consider it to be a stronger, slight more and sognite, alternitive to Deca-Darabolitti. It is agreenable downer, and cold replece Deca in most cycles wholes acrossible contrasting of blood of productions, although there should be on as disc capable of increasing of blood of productions, although there should be one as disc capable of metanting of culture transfer stronger.

History:

Che neportally patentel bolkenore as a syntheir analysis staroid in 1949. During the 1959s and 10% he time dockped second reperimental sets of the drug, and would have relase a long-acting form of the agart (hridy) in the form of bolkenore undecylenate. It would be solit advect the brand nume Patentiohly while high effect of a characteristic of 8% and a day 7% and million and the staron of the staron of the star and the 10% and a day 7% and the staron of the star and the star and the star and the star and the start and the star and the star and the star and the star and the start and the star and the star and the star and the star block the star of the 10% Scaph would instartly he most famous for instarding the starget neutrino in the start and the star and the star and the star and the star and the start and the sta

In the verticing mutic, bolknene andecjenne is most commoly applied to bones, alterogin in many regions is indicated from in other annula so will a gravarily collobs a pronounced effect on him bodyneight, and gravari disposition of the annul. The USA mutic hash bodyneight, applied to body the start of the second the USA mutic hash bodyneight and body the first the rest the USA mutic hash bodyneight and body. Many other generic and learning mutic the USA mutic hash body and body. Many other generic and learning mutic the USA mutic hash bodyneight and body. Many other generic and learning mutic any partners on bodyneight with the science exprised.

How Supplied:

Boldenone undecylenate is widely available in veterinary drug markets. Composition and dosage may vary by country and manufacturer; the majority of products are supplied as multi-dose glass vials containing an oily solution; usually carrying 25 mg/ml or 50 mg/ml of steroid.

Structural Characteristics:

Bolkness is a molified ferm of tosteneous. It differs by the introduction of a double boad between carbons 1 and 2, which relates in relative estimatory with an altogenetic flagshophic consists the 17-boar horizont part particular to the start base in the start of the start of the term of the start of t

It is interesting to note that structurally boldenone and methandrostenolone (Dianabol) are almost identical. In the case of boldenone (as applied here), the compound uses a 17-beta seler (undecylerate) to facilitate administration, while methandrostenolone accomplishes this with the use of a 17-applie alkyl group. Aside from this, the molecules are the same. Of constraints of the other that here the bold with one to show the the 17 avoid-tation account of the other that. course may act quite unterently in the body, which goes to show that the 17-mentylation affects more than just the oral efficacy of an anabolic/androgenic steroid.

Side Effects (Estrogenic):

Bolkence is anomatical in the body to estratide (entropyn). Elostade entropyn levek en cance side effects sen is nerened water retorints, hody fing sin, and greacomstati. Bolkence is considered a målly estogaris sterid. Aromatization statistis saggett hut is nine of convesition to estimation is anguly half hut of teststered." The testherey to develop metachek sensorie site effects with bolkence should be slightly higher than mandonos. But mich, hower flux with bosols above 200-400 mg per vock. An anisisis effects, double hower powerfs Down altermaticity between the variable site. The hower powerfs Down altermaticity are uncertain sitelihof and Armiteiko (cuastructurk), although i is considerably more expension, and may negatively and after blood fulls.

Side Effects (Androgenic):

Albeagh classified as an anabolic steroid, androgenic side effects are still common with this substance, opeoidal with higher does. This may include boats of ody skin, acen, and body/ficial har growth. Anabolicandrogenic steroids may also agarsate male pattern hair loss. Worten are also warred of the potential virificing effects of anabolicandrogenic steroids. These may include a deepening of the toxice, menstratal irregularities, changes in skin teatre, ficial inlar growth, and chaol endragrement.

Note that while boldenone does reduce to a more potent antrogen (dhydroboldenone) via the 5-alphu reductase enzyme in androgen-responsive target tissues such as the skin, scalp, and prostate, is a affinity to do so in the harman body is extremely low.⁵⁰² The relative antrogenicity of boldenone is, therefore, not significantly affected by finasteride or dinstateride.

Side Effects (Hepatotoxicity):

Boldenone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabolizantingenis sterolis can huve delutricos effects on serum cholesterd. This includes a tendency to rolex (EML (agod) cholesterol values and increase (ELL (bad) cholesterol values, which may shift the FIDL to LDL bahare in a direction that floors greater risk of attrictureous. The realise trapact of an anabolizantogenis extraodio an serum (pist is dependint on the dose, route of administration (carl vs. injectabel), type of steroid (normitable to route-anomitable), and level of resistance to heaptic retrictions Anabolizantogenis steroids may also adversaly affect blocd pressure and trajlocritics, research and the steroids of the variantical heaptical inflation. Roldworce is heap to steroids. This chi and prot to separates to tertobeshow type for a which also is to heap its and the research of the steroids of the steroids and proceeding inflations. Roldworce is heap to steroids. This chi and prot to separates to retribushing byte for sub-thankows to how less effect on the heaptic measurement of cholesterol. The arountation of Doklorove to sterial on walo heaptic measurement of cholesterol. The arountation of Doklorove to sterial on the heaptic measurement of the effects of andresson sortem finits.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of statrated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ols (4 grams per day) and a natural cholsterol/antioxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimulating substances, testosterone levels should return to normal within 1-4 months of dung secession. Note that prolonged hypogonadotrophic hypogonadism can develop secondary to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Although is stays active for a much longer firm, holdenne undecylentue is injected at leasts weekly for physical or performance-entancing pupposes. It is most commonly used at a dosage of 200-400 mg (4-Smt50 mg version) per week. The dosage schedule can be firsther divided to reduce the volume of each injection if necessary perhaps administering the drug two to three times per week. One should also take caution to rotate injection siles regularly, so as to avoid irritation or hitcrin.

Not a npd muss bakke, bolknone undeejkente instead provides a skow hut steady gain of strength and quality macker muss. The positive effects of the strengt part of th

Bokknow undecyknus is a very vestaki drug, and can be confused with a nurber of order agains depending on the desired reads. For muss, it is convery stucked with an ispicable biostaneous each as caratheter or cyclenate. This shead produce strong gains in a higher does) also: The single acting players much benchess and domy on the graph trapovol when combining bolkness nucleolytenic with a non-anomitable strongly trapovol when combining bolkness nucleolytenic with a non-anomitable strongly trapovol when combining bolkness nucleolytenic with a non-anomitable strongly trapovol when combining bolkness nucleolytenic with a non-anomitable strongly trapovol when combining bolkness of the strongly the site of the stronglytenic functional stronglytenic stro

Administration (Women);

When used for physique- or performance-enhancing purposes, women take much lower doses of boldenone undecylerate than men, typically 50-75 mg per week. Women should take caution with the slow-acting characteristics of this preparation, which make blood levels difficult to control and slow to doeline should viriation symptoms become present.

Availability:

Bolkinone undersplente remnis widely available as a veteriary drug product. It is produced main ju hie America, consistently at a doage of 25 mg/nt. A 50 mg/nt. A small namber of preparations are made at a higher doage (typically 200 mg/nt.), mainly by companies in less regalated markets of Asia where sapply is often distated by black market demardt. In reviewing some of the more popular products and changes on the global pharmacerical market, where made the following doserations.

Brand name Equipoise is no longer available in Canada. This product was formerly manufactured by Wyeth Animal Health, which following a series of mergers is now part of Pfaer. The Equipoise product was never manufactured under the Pfaer label.

Equipoise® is produced in the United States by the Fort Dodge Company. The Fort Dodge products are sold readily in Mexico, and afterwards srnaggled back into the U.S. Legimute vials are made of clear glass, and carry a label with a shiry metallic surface on the under side. Fakes are very abundant of this item.

Ganabol, which is produced in a number of South American countries, is still a popular brand in international commerce. It is seen in two strengths (25 mg/mL and 50 mg/mL) and in five sizes (10, 50, 100, 250, and 500 mL). There have been numerous fakes of this product in the past, so be careful when shopping.

The Legacy brand name product from Tecnoquimicas in Argentina seems to be reaching the U.S. as of late at least in small volume. This product carries 50 mol/ml. of steroid in a 50 ml. vial. At this time the Legacy product is very low on the radar, and probably can be trusted when located.

The brands Boldenona and Boldegan from Gen-Far are also popular in South America. These are low-dose (50 mg/mL) preparations. Like Ganabol, they come in a variety of vial sizes. Counterfields do not appear to be a big sizes at this time.

Finajet (trenbolone acetate)

| Progestational Ad | tivity moderate |
|--------------------|--|
| Estrogenic Activit | y none |
| Chemical Name | 17beta-Hydroxyestra-4,9,11-trien-3-one |
| Standard | Nandrolone acetate |
| Anabolic | 500 |
| Androgenic | 500 |
| | |

Description:

Terobotes accettate is an injectable (apexturbly analysis attenuid derivated from marchotene. Its activity is quite removed from its structural pureet, howevers, such that direct comparisons between the two are difficult. Terobotene is a non-estrogene structural, and is considerably more analysis, and analogonsis than marchotenes on a milligner moles. Its appearance, it is much more commonly compared to a stronger androgons such as domatonice, than its iso marchotene. This should be of the strangest signable intrability to a stronger interpretation of the strangest signable intrational to the structural structural structural and the strangest signable interpretation of the structural structural structural structural structural structural structural transformed between the structural structural structural structural structural reference and fit muss gins. It is considered a dang of choice for costnets bodyhaides, yet eminima very product with divercentional uses structural body and the prints structural structura structural structural structura structura

History:

Teroboux acetate was first closely statisf in 10%7, described daring a series of operiment in storytheir annolek acetaski by Rosask-UCLAFS, By the earby 10%7, terobotne acetate was being sold in England by Hoeedat as Franjet, and in France as Franjest by Rosask-Rosal-Ro

Terobone accette frei became popular annote U.S. holyhakken daring the 1990%, a time whom the drag was being maggids in fina francopse in light values. It was identified (rights sa) on a popular distribution of an analysis and a strategies and a strategies of the constraints of the same strategies and the shorts and the same strategies and the same strategies and the strategies and the same strategies and the same strategies and the strategies and the same strategies and the same strategies and the strategies and the same strate

Around the same time as we were seeing the demise of Finajet and Finaject, Hoechst-Roussell was introducing trenbolone acetate to the U.S. market as Finaplix cattle implant poliss. This carne subsequent to the FDAs approval for such products in 1997. The poliest were designed for substancous implement in to the car of the carle with a handled implant gar, and are fit too large to be implement in humans without mirror surgery. Remutably trendous eactine pellsas are even from U.S. corrected substance income the presentably to motive a surgery and affittable for beaution, or most to have access to the surgery strength of the stren

Hamn administration of Finglicy polets can be difficult to accomplish, but it is all widely down. More correctly, too to fare ripplest polets are growed approximation with a 500 syswater and DANO solution, which is applied to the data (day. The hore-hore transmuttering the system of delevery of strength hermore. These only hore wire in the done in the system is not as intring as they had impacted it would be. Advantation, some atthese ogst to simply commande the dong biosentidity and not be used in this memory the analogue and the system.

Mee absentations individuals have rands it a practice to mix their own isjections with Fraphic. The pelose are grownal into a fite properties and posterior from 2.6 6 peloses, and them are added to startly water, properties glocic or not i-based specified istended or understanding and the properties of the start of the start of the start of the environment of the startly water of the start of the start of the start of the environment of the start of the start of the start of the start of the environment, however, one is obvioually risking infection (or vorse) by doing this. Starting in the last 1990), seems to be leaders that the start at leader of the start o

Frapfiels is presently available in the U.S. and some numbers abroad, although it is now being old by Intervent instaud of Hotscher-Kossend Agrivet. This product cornes in nov forms, Frapfiel-H and Frapfiel-S, which denotes if the product rouse in its nov containing (10) 20 mg truebokose accente peldes (2000 mg) and the "5" vesion only 70 (14) 00 mg (1) y-dama Health (U.S.) has included nov competing products of qualitation matching and the observable of the structure of the product set of patient matching and a component HI and Component PS. There are not the black of estimation of the structure of the structure of the structure of the structure of estimation of the structure of the structure of the structure of the structure of estimation of the structure of the structure of the structure of the structure of estimation of the structure accente estit, the drug is produced (for injection and oral use) by a number of ceport and underground stored in matcheren.

How Supplied:

Trenbolone acetate is available in select veterinary drug markets. It generally comes in the form of implant pellets containing 20 mg of trenbolone acetate each. Injectable preparations containing 30 mg/nt of steroid in oil were formerly sold.

Structural Characteristics:

Terebokes is a modified form of randroken. It differs by the introduction of double bonds at carbons 9 and 11, which inhibit anomization (0--one), increase and/one)-binding affinity.³⁰ and show its metabolism. The resulting steroid is significantly more potent as both an ambolic and an androgen, than its matroloce bases. Forbolose accubent contariss terebokene modified with the addition of carboxyfic acid ester (accit acid) at the 17-beas ployong joury, so that the free steroid is released more aboutly form the zeas of rejection.

Side Effects (Estrogenic):

Trenbolone is not aromatized by the body, and is not measurably estrogenic. It is of note, however, that this steroid displays significant binding affinity for the progesterone receptor (slightly stronger than progesterone ised) 15^{05} 506 The side effects associated with progestrove are smart to those of estogen, nexture, pagine lexotises remotion of tostsocience production and enhanced the of for storage. Progestris also agarent the similatory effect of estrogens on marmary tissing growth. There appears to be a storage sprogreg betwoen these two hormenses here, such that gencoments implications are storagen, which the help of progestins, without excessive estrogan levels. The use of an arti-storagen, which inhighs the estrogenic component of this disorders, of one sufficient to might gencommarks caused by progestinical antibolicant/ogenic storakis. Note that progestinical ide effects are more common what therefoloce is being than with dorf aromatindus storakis.

Side Effects (Androgenic):

Albude closified as an anaboic steroit, trebohore is anflicently antrogenic. Antogenic sist effects are all common with its systemace, and may include loss of oly skin, accand bodyficial har growth. Anabolicanitogenic steroits may also agenute mate pattern har loss. Worms are also wannel of the potential visible effects of anabolicanitogenic steroits. These may include a deopening of the vice, menstual inregativities, changes in skin tutern, ficial har growth, and closed angeneric. Additionally, the 5-dpht notclasse erogenu does not metabolite twoblocker.⁶⁰⁷ so its relative androgenicity is nt affected by finsterice of antastrotic.

Side Effects (Hepatotoxicity):

Terebolene is not c-17 alpha akylated, and is generally not considered a hepatotoxic steroid; bser toxicity is utilkely. This steroid does have a strong level of resistance to hepatic breakdown, however, and severe liver toxicity has been noted in bodybuilders abusgi terebolene.²⁶⁸ Albough utilkely, hepatotoxicity cannot be completely exclude, especially with high doss:

Side Effects (Cardiovascular):

Ambelicantegaries strends can huve delutricos effects on serum chosterort. This includes a tendency to reduce IDE (agod) cholsterou values and increase. IDE, (bad) cholsterol values, which may shift the HDL to LDL bahare in a direction that fivors greater risk of attrivichersis. The reduce transformation and the antibioinal organic strends of the dependint on the dose, route of administration (call w, injectibelit, type of sterold incommitable mata and source pressures to metholsmin, turbelouch has an indexinate to more anomalable mata and association of insigned to heat methodam. But is the more anomalable mata and association in the methodam has the modern of host above and the lobel pressure and infiguredises. And accordinate to may host obevery differe lobel pressure and infiguredises. And accordination in the dosed scares and metacidination information.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intrake of stantard fats, cholsterol, and imple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholsterol/antioxidant formals such as Lipid Subil or a product with comparable introdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolic/androgenic stencils when taken in doese sufficient to promote marke gain are expected to suppress endogeneous testoreme production. When the intervention of testosteneous-straindard gashatances, testosterone loveds should return to normal within 1-4 ments of end gas essents. Note that providegal hypogenomhypic hypogenoments in the strain strain strain testostero and an alling and the strain strain strain studies, nerboken was determined to be approximately frare times stronger at suppressing grandetorpite time tostosterone on an allingma for miligram basis.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Trenbolone acetate was never approved for use in humans. Prescribing guidelines are unavailable. An effective dosage for physique- or performance-enhancing purposes generally fails in the range of 100-300 mg per week, taken for 6.1 8 weeks. Due to the short-acting matter of nearthenesses, the task week's docume is of briefweld in 2-3 smaller applications. See 5 weeks to minima any potential hyperia starts. This has been addresses to the short of the start of the

For stacking, trendolom is a very vesnik steroid, and scens to took ecceptionly well with other agents for both doing and curring proposes. For criticity holdwales of outstack it with a random kike Wasticolf or Prinzobaths. Whitso extra water bound your shorts the start of the start grady entrue and solidiy the new mack growth. When looking purely for musiterbohore pairs well with instastores, Automated 50%, or Dainabild The scalit is spaced Scale and the start of the random and substarting pair of somewhat solid mack muss. In the Undergrand Scaled Hardook H. The Division describe the run of therbohore, its stored and scale wells and the start of the start of the start of the start of the start wells and the start of the start of the start of the start of the start wells and the start of the start

Administration (Women):

Trenbolone acetate was never approved for use in humans. Prescribing guidelines are uravailable. This agent is not recommended for women for physique- or performanceerhancing purposes due to strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Pharmaceutical preparations containing trenholone acetate remain scarce. The bulk of the supply for this compound comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Terebolene acetate is still produced in the United States, sold in the form of cattle inplant pellets (minity Finaplicit). This agricultural product is also exported to some neighboring markets, though is not widely popular outside of the Americas. These pellets are difficult to administer, and are not commonly the subject of counterfeiting. This is the only legitimute form of trenbolene acetate in the U.S.

Most other forms are from underground manufacturers, and therefore are of unverifiable quality.

Halotestin® (fluoxymesterone)

| Androgenic | 850 |
|------------|---------------------------|
| Anabolic | 1,900 |
| Standard | Mathultastastasana (aral) |

| Stanuaru | | methyltestosterone (oral) |
|---------------------|-------|---|
| Chemical Names | | 17b-dihydroxy-17a-methyl-4-androsten-3-one a-fluoro-11b-hydroxy-17a-methyltestosterone |
| Estrogenic Activity | | none |
| Progestational Act | ivity | no data available (low) |

Description:

Flavoymsterone is an onl anchois steroid drived from teststerene. More specifically, it is a methylestoxene or drivenite, differing by the addition of 11-best-holyon and 3-alphafaror groups. The result is a potent coally active new sommitzable storoid that exhibits externely storag analogies (properties T, knownstearone is constabled) more androgene than testsochene, while at the same firm the analode effects of this agent are considered to be negligated and the storage transmission. The storage data and the storage data and the methyle and the storage and the storage data and the storage storage and the storage data and the flavoymsterose are increased storageh, acrussed marke dorsity, and increased definition, whitten protection terreases.

History:

Flacometerone was find elser/bei al 1956.⁵⁰⁰ It was assoped that same your, and shown to possess approximately 20 mers the analysis potency of methylestroneous¹⁰ (in television the base in the structure of the structure of

By the mix140705, the TDA halo been grated mach more control over the U.S. dueg market. One of the imprive thanges with stored machine market the FDA required strong asbitaritistic for each potential use of a drug. The prescribing grateflowes for flavorustiences were some reduct to start the drug uses "reflective" for tearing arrivas forms of analogues deficiency a mules, and reducing the secority of required markets and tradingtant depresentations are included as the start of the strategy arrivas and tradingtant depresentations are included as the start of the strategy arrivas for a flavorust strategy and the strategy and the strategy arrivas and the start of a flavorust strategy in the strategy and the strategy in marks and breast current intensis.

In recert years, facymesterone his boorne viewed more and mere as a controversial mediation in the system of model in the planotaticity and planotatic angustic impact of lipids and cardionaschar risk factors are often cied as reasons for avoiding the use of this again in otherwise heally multisk for transition and possible model. To device the repearation (rigication, gals, patches, implans, etc.) are predired for this parpose, and they applient the same analogous mixing from the body (rotatorence, Tody), to store trace synthetic deviations. Flavorum-storem remain for sake in the U.S. as a generic drug only. It emains available in one brief our planotation of the U limits areas.

How Supplied:

Flaoxymesterone is available in select human drug markets. Composition and dosage may vary by country and manufacturer, although generally contain 2mg, 2.5 mg, 5 mg, or 10 mg per tablet.

Structural Characteristics:

Fluoxymesterone is a modified form of testosterone. It differs by 1) the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during oral administration, 2) the HITODUCION OF a BLORD group at Carbon 9 (apra) and 5) the attachment of a hydroxyl group at carbon 11 (beta), which inhibits steroid aromatization. The latter two modifications also greatly enhance the androgenic and relative biological activity of the steroid over 17-alpha methylaetsoterone.

Side Effects (Estrogenic):

Flaxymesterone is not aromitized by the body, and is not measurably estrogenic. An antistrogen is not necessary when using this stendy, as generomissi should not be a concern even among sensitive individuals. Since entrogen is the usual culprit with water retention, this steroid is nead produces a law, quality look to the physique with no far of encouss subcatancous fluid reterition. This makes it a forwable steroid to use during cutting cycles, when water and far territion rule migor concerns.

Side Effects (Androgenic):

Flooynesterone is clossified as an androgen, Androgenis side effects are common with this substance, and may include bosts of objects, and explosible and here of the Anabolicandrogenis stronds may also aggreates male pattern har loss. Those gravitable porce to make platen har loss may with the opf f as a milder, gas androgenic, anabolic strond. As a potert androgen, this steroid may also increase aggressioness. Wromen are additionally sumed of the poterality sitting effects of anabolicandrogenis stronds. These may include a deepering of the toxice, menstral inregalarities, changes in skin texture, facial har growth, and clonder longeneret.

Flucymesterore appears to be a good substrate for the 5-alpha reductuse enzyme. This is evidenced by the fact that a large marker of is metabolises are found to be 5-alpha reduced androgene³¹¹ which coupled with is outward androgenic manee, suggests that this steroit is conventing to a mach more acrite steroid in androgen responsive target stress such as the skin, scale and prostate. It may be possible to reduce the relative androgenicity of flavouresterone by the concurrent use of fistastrice of thatsertic).

It is also of rote that flavormesterore has been shown to process usual androgarie propertiss. In human sales published back 108/1, the storid oblogical a meth storegy tendexy to promte public enlargement compared to other androgarie effects such as har growth. Back, and Changs in word plack? The Disconstructione use offering a somethat different androgarie profile compared to testosterone, and as such demonstrated that it was possible, at some lock to statudup airst rung differ statistical chargery of androgaries that it may not offer a complete biological capitalier to testosterone where androgaricity is concerned.

Side Effects (Hepatotoxicity):

Flacometeore is a (17-ab)ta abjekted compound. The abention protects for dung tiern descritation by the size, allowing a very high pocentrage of the dung entry into the bloodbettenen filowing certa abaritation. (17-ab)ta abjekted anabolicambogene sereods is discussing dispersion and an abartation. (27-ab)ta abjekted anabolicambogene sereods dispersion and abartation and cound health. Intel and the dispersion producibly during each cycle to memor here function and ocean health. Intel and cycle 1/abje abjects for two weeks ababited and the dispersion of the dispersion address and abbited the dispersion of the dispersion of method of the dispersion of the dispersion of the dispersion of the dispersion of the method of the dispersion of the dispersion address and the dispersion of the dispersion of the method of the dispersion of th

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anahofisiandrogenic steroitis can have delsterious effects on serum cholesterol. This includes a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol values, which may shift the HDL to LDL balance in a direction that floors greater risk of arterioselerosis. The relative impact of an anaholic/androgenic steroid on serum lipids is downedrut on the dose matte of administration (or all w. iniscratish) true of stornil represente ou test sources contractions of the second source of the seco

To help reduce cardiovascular strain it is advised to mrintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimuling substances, testosterone levels should return to normal whiti 1–4 months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

Statiss administering 10 mg 20 mg of 30 mg of flaxymesterone to nine healthy male subjects for up to 12 weeks have dermentated the strong suppression of endogenous testosterone keeks, with inconsistent effects on grandstropin levels. Although not fully understood, flaxymesterone is proposed to have a direct suppressive effect on testicular steroidogenesis that is not mediated by the suppression grandstropins.²¹⁴

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.¹⁵⁵ This is caused by the fat-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undigsted disfary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

To treat androgan isofficience, early prescribing gaidelines for flavoymesterone called for a dose or 2-10 rung per day Modern prescribing gaidelines call for a ddy dosego of 5-20 rung per Therapy is usually initiated at the fill 20 rug dosaga, which is kiter adjusted downward to meet the infoldual needs of the patter. The deng would be contrained long-term using hiberatory usus, fipikle, here ensymes, etc.) or side effects contrained long-term using the many of the rung of the rung the rung in the rung of the flavored hiberatory per start of the rung of the rung tabaset in specific section of 2-100 rung of the rung of 10-40 rung tabaset in specific sections in numely measured in the rung of 10-40 rung tabaset in specific for measurable increases in numely music.

Flacometerone is commonly used by athlets in weight-sentited sports file, was meeting, proverifing, and housing, due to the fart that strengh pixel filt much ends us usually we accomparing by grant accesses in bodyweight. When propely used, it can also a competion to sky within a specifical weight manage, yield straight proves he porformate. Flacometerone is also controlly used for bodybatility context preparation. When the competion to any strength most pixel and other that pixel is absence of encours estimation of the strength pixel and other through hold to the matchs, and the strength of the strength pixel and other the pixel and the body may he more strends to how off excess fixe and proves the fistenges. The "handwaig" effect of theorymeterone would, therefore, he somewhat simile to that seen withtendowed, altophic will be without the same level of musics grain.

In cutting phases, a milder anabolic such as Deca-Darabolin® or Equipoise® is commonly stacked with flaosymesterone, as they provide good anabolic effect without excessive estrogen buildup. Here, fluosymesterone provides a well-needed androgenic component, helping to promote a more solid and defined gain in masc muss, with his is interference with energy and likely, have might be obtained with a primuly amole degrar date. Perhops Primoballed-Depart would be an even better choice, as with such a confinition there is no halding of estingen, and likewise even its source of water and if a treatment, for muss, care might alternative use an injectable instantence. A rute of 600 mg per weak of treatments increases in strength alternative must be an origination for 100 md/molegen idea officies usadly accompanies such a confinition, however, as both compounds exhibit strong androgenic activity in be holy.

Administration (Women):

Flaxymesterore is most often used as a secondary medication during inopenable androgensensine breast cancer, when other therapies have field to produce a desirble effect. The dosage used for this application is 10-40 ng per day, Vinlärg effects are common at doses of only 10-15 ng per day in these patients. Flaxymesterone is not recommended for women for physique- or performance-erlauxing papeoses due to its strong androgenie rature and tendersy to produce viriaging side effects.

Availability:

Pharmecutical preparations containing fluxymesterone remain scarce. The drug hus hrgsly associated with westerm medical markets, where it has been falling out of favor for most clinical application. The bulk of the supply presently comes from underground steroid manufacturers and export-only products. In reviewing scene of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Currently, the most popular item found on the black market is the Stenox brand from Mexico, which is sold in boxes of 20 tablets. Note that the dosage of these tablets is only 2.5 mg.

Flaoxymesterone remains available in the U.S. as a generic drug by USL Pharma. It is available in a 10 mg tablet only.

Balkan Pharmaceuticals (Moldova) makes the product Halotest. It is prepared in 2 mg, 5 mg, and 10 mg tablets, with 20 tablets contained in each foil and plastic strip.

Laurabolin® (nandrolone laurate)

| Androgenic | 37 |
|--------------------|--|
| Anabolic | 125 |
| Standard | Testosterone |
| Chemical Names | 19-norandrost-4-en-3-one-17beta-ol 17beta-hydroxy-estr-4-en-3-one |
| Estrogenic Activit | y low |
| Progestational Ac | tivity moderate |

Description:

Nandoken lamte is an injectable formofthe anabole steroid mardoken. The harare ester applied here is two contantos logar franch denconte, and concourgent phis agent frans a sightly logar-lasting ding daposit around the area of isjection than Deca-Darabolic. Given is strong dedpose/sterogeneous in the possible to administer methodoken lamte concervey three to four weeks in a medical setting. As a mardokoke injectable, this ding movides an inducatify around effect, which is accompative by a low teef of astogaties and an inducatify around effect. which is accompative by a low teef of astogaties and and budydney the half on porcess dave storing gathers are with minimit data effects.

History:

Nandoken launte was dovelped daring the 1960's, a tire when meny new mathokee estors wave being ornskied and mesengiater. It his host-acting estor for ofmathoke is usual identified as a voteriarry drag. In twas actually prescribed to harmin before it was adapted tire on the German ding mutch around the est of the 1960's human before with a strange of the strange of the strange of the strange of the actual distribution of the strange of the strange of the strange actual distribution of the strange of the strange of the strange of the stability distribution of the strange of the strange of the strange of the stability of the strange of the strange of the strange of the strange of the stability of the strange of the str

As a vetering ding, mutchoke larnet is most corruptly identified with the Larneboth bund mare. Landboth is manifectual by lattered, and is fourd in a outriey of countries including backco, Chik, The Netherhands, Australia, and Cokorbia. It is used with east, adogs, hences, pips and cattle, typical to outform function catassate by viait of parasite illustration of highly active or defay animels. The Landboth is much as and outerall physical condition of highly active or defay animels. The Landboth interal has and proceed they are interacting and the state of the state of the state of the state of the procedure lines into the mission of the state of the state of the state of the state of the procedure lines into the mission of the state of the state of the state of the state of the mission of the state of the parasit under the mask marks Fortaket (phylos, Germany), Fortakel (Partim, Mexico), and Landbol 250 (Loeffer, Mexico). Today, only the Intervet procedure to exist.

How Supplied:

Nandrolone hurate is available in select veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 20 mg/mL or 50 mg/mL of steroid fissolved in oil.

Structural Characteristics:

Nandoken lanate is a molified from of matchoine, where a carboylic acid ester (lanzicacid) has been attached to be 17-beta photograpor. Esteridis destruits are less point fram free staroids, and are absorbed more slowly from the area of significant contrast in the biodentum the ester is removed to juid free (archie) nandohene. Estarified staroids are designed to prolong the window of thranpenic effect following administration, allowing for a lass fraquert significant schedule compress to biperiots of the (masteridis) staroid. Nandohene lanza is designed to provide a slow relasse of randrohone for up to 3 to 4 weeks fillowing incretion.

Side Effects (Estrogenic):

Natiobles in a low tendency for estingen convenies, estimated to be only about 20% of the sense with testocorress³³. This is baccured as which the force more convert matchoose to estandial, in other more active sites of steroid arountations and the sense of the source and the sense of the source areas a much how converts with the days than with teststearces. Excited estimates the sense areas a much how converts with the days that with teststearces. Excited estimates the sense areas a much how converts the sense of the sense areas a much sense the sense areas a much sense the sense areas a much sense that the sense areas and the sense and the sense areas and the sense and the sense areas and the sense and the sense areas and the sense are an and the sense areas areas and the sense areas and the sense areas areas and the sense areas areas areas areas and the sense areas areas

It is of note that matchine has some activity as a progestin in the body.²¹⁸ Although progestrone is a c-19 seroid, enroval of this group as in 19-noprogestrone creates a hormone with greater binding affinity for is corresponding receptor. Sharing this trait, many 19-nor analyois arcoids are shown to howe some affinity for the progestrone receptor as well.²⁰⁹ The side effects associated with progestrone are similar to those of esttogence production and enhanced rate of fast for a field and the field of the states and the field of the state of the source affinity of the states effects associated with progestrone are similar to those of esttogence production and enhanced rate of fast fields.

stonge. Progestins also augment the simulatory effect of estrogens on nummury issue growth. There appears to be a storng synergy between these two hormones here, such that gencomstain ingit even occur with the help of progestine, without excessive estrogen levels. The use of an anti-estrogen, which inhibs the estrogenic component of this disorder, is often sufficient to misging engencomstatic caused by randrolone.

Side Effects (Androgenic):

Albudg choseful as an ambolic stroui, antopopie size effects are still possible, with first substance, opercival with higher dows. This may include boots of op skin, are, are, and body/ficial luir growth. Anabole/antopopie steroids may also aggroute mile pattern luir loss. Wornen are sumed for he poterial virialing effects of anabolicaringens storoids. These may include a deepening of the voice, mentional regularities, changes in shale instantical luir growth, and chosed selfangent results. The set may include a story relation to its fiscal body and again a construction of the poterial vision-balling actions, making the the chold for storage instances and the story relation of the story relations of the story potential point of the story does not interfavoid likely story of the story story of the story of the story of the due to its mail and togonic matter and ability to suppress endogenous toxitoraters, matchookies proce to interfavoid likely loss.

Note that in antrogen-responsive target fiscass such as the skin, scab, and prostate, the relative antrogenized yof matrolones is related by its reduction to displormatiokous (DRN).²⁰²³ ²¹³ The 5-alpha reductase enzyme is responsible for this metabolism module matcholese. The concurrent use of a 3-alpha reductase influence and matcholese the concurrent transformation of the stable stable stable stable stables and the stable stable stables and the stable stable stables and the detection of the stablese stable stables and the stable stable stables and the stable stablese stable law could with matcholese The stablese stable stables and the stable stables and the stable stables the stable stable stables and the stable stables and the stable stables the stable stable stables the stable stables stables the stable stables the stables stablesta

Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabelicándoganis storolás can huve deletricosa effectis on serum choelserot. This includes a tendency to rolace JELL (god) cholserol volasse and increase JELL (bud) cholsetosti valass, which may shift the HDL ogd) cholserol volasse and increase JELL (bud) cholsetosti a dependint on the dose, route of administration (cell w, injectabel), type of steroid (gornalized ero cosmittabel), and level of resistance to hepatient tentholamis, Static administrating (50) ng of mathelisto decanozi per work for 10 weeks damonstrating 20% with an equal dose of testastero the parameters with emilies at static with an equal dose of testastero test expression is adding state than that results staving concerned to testasterone constraints, and is in agreement with emilies radius staving a significatify worker impact on serum hpits. Imm -17 adpin adopted agarss, Anabelicánichogarse starolás may and support kil wertricour dud stall tures a significatify worker impact on serum hpits. Imm -17 adpin adopted agarss, nuclear enduchiai relations, and a support kil wertricour hoperandin to tradiers and proteindus in tradience and a support kil wertricour hoperanding hoperturgly.

To help reduce cardiovascular strain it is advised to mnitatin an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fab (a) (4) grams per day) and a natural cholesterolamicoxidant formatia such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All anaboléandrogenie steroids when taken in doese sufficient to promote muscle gain are expected to suppress endegenous testostenoue productions. For sake of comparison, studies administering 100 mg per week of randrolone decanoate for 6 weeks have demonstrated an approximate 57% reduction in securi testostenous levels during therapy. At a doagge of 300 mg per week, this reduction reached OMC⁵⁵ It is believed that the prograssitional activity OMC⁵⁵. minkolone notably contributes to the suppression of testosterone synthesis during therapy, which can be marked in spite of a low tendency for estogene conversion.²⁵⁵ Without the intervertion of testosterone-stimuling substances, testosterone levels should return to normal within 2-6 months of drug secession. Note that prolonged hypogeneidomorphic hypogeneidom can develop secondary to stored abuse, necessioning medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Nandoken lannte is net approved for use in harms. Prescribing galdeless are unvalidite. When used for physicase or performance-entermine proposes, a door 002-000 ong give every $^{-1}$ -10 days is most common taken in cycles is for 12 weeks in high Tin keel is down to the structure of the str

Administration (Women):

Nandoken lannte is net approved for use in harms. Prescribing galdeless are unvalidable. When used for physica or performance-entrancing paposes, a dogo of 100 rag every 10-14 days is most common, taken for 4 to 6 weeks. Although only slightly antlograsi, sowers are occussional confinent dwh which thinks mystems when much thinks the scoregood. Should writing able effects become an concurve, the dug should be discontinued immediately advert-assignt much done. Databable in dipple to considered a star forme contrabable, option. This dug starties for only several days, grathy reducing the withdroud time if microted.

Availability:

Pharmaceutical preparations containing randrolone laurate remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

The Intervet brand name Laurabolin product is almost exclusively associated with this compound. It remains available in select markets of Europe and the America's, and is not widely diverted for black market sale.

Masteron® (drostanolone propionate)

| Androgenic | 25-40 |
|------------|--------------|
| Anabolic | 62-130 |
| Standard | Testosterone |

| Chemical Names | | hyl-androstan-3-one-17beta-ol ha-methyl-dihydrotestosterone |
|--------------------|--------|--|
| Estrogenic Activit | ty | none |
| Progestational A | tivity | no data available (low) |

Description:

Drostanolone propionate is an injectable anabolic steroid derived from dihydrotestosterone (DHT). Here, the DHT backbone has been modified with a 2-methyl group to increase its ambole properties, making this agant significantly more effective at promoting the growth of march tissue than its non-methylated praceater. Dotostrokne proportione is docsrbod in product literature as a "stenoid with proverif at ambole and arti-settogenic properties," and mixed doos seen to be seen our of both properties. Admittable, however, it a ambole properties are more properly doceshed as moderate, especially when placed in the context of other agares. The data part of the masses of the data properties and antimore, begin of other agares. The data part of the masses of the data part of the masses of the masses and and strength, which are usually accompanied by reductions in body fit level and minimal side effects.

History:

Dostanibure propionte vus first described in 1699-2055 Syntex developed the agent alsegide and note veloknom stereito is a Antohi and methylotistochere (Sperende), also first doscribed in the same paper. Dostanohore propiente world be istraduced as a proscription darg product approximitely a doscale liste. J Bil and an agenerar with Syster to split certain research and development costs in exchange for the rights to mutler the nessko of that research. Bly world, therefore, all dostander programme in the U.S. and the Doshne hund mane, while Systex would selfkemes it in other runket the nessko of that research. Bly world, therefore, all dostander programme in the U.S. and programmes as Permissif (Cassense, France). Massing (Shieneyi, Japani), and Masterii (Granethi, Garmany Doncentie Republic).

The U.S. Food and Dag Administration approved doctatarkose propiones for the treatment of advances incorrels by test active are postmarrougoal sources. The remained the principle clinical indication for the agent is all immediated matches as well. The violation with doctatories propiones are compressed to equil the order of the streatment of the stream of the stream of the stream of the stream projections, the dispersion and access that the stream of the strea

White high popular armong athlesses daring the 1970's and 200°, disotanokene propirante ultimutholy enjoyed limit discusses as a preservition agart. Manafartherers begin volaritudy discottining such of the agart in various matchets before long, likely due to the advert of more efficient hempissis for breast cancer, as well as the show define insteard preservitions for this place of treatment. One of the first preparations to go was the U.S. Doban, which was entrowed form muthed draing the like 10% Remmestil and Mecoman wares cound drapped as well. The last remaining Western preparation croating distancely enpointentern this field on the U.S. Primmerski and the is preserving being mothlock to its sike, although is recorregence as a prescription drag product seems highly utiloky.

How Supplied:

Drostanolone propionate is no longer available as a prescription drug preparation. When produced, it was supplied in the form of 1 mL and 2 mL ampules and 10 mL vials containing. 50 mg/ml or 100 mg/ml of steroid in oil.

Structural Characteristics:

Dostankow (aloo karon as demostandowij sa molified form of diplottestorteree. It differs by dse introduction of a melly appear a tembora. [alpha), wide considerably intratasse he anabok, strength of the strend by highing in particular to methodshim by properties is a molified form of dostantices, where a carbony case of a strend activity of the strends of the strend by form the strend strends are used in strends and the strends of the strends of the strends of the strends and the short the strends of the strends of the strends are used as the blockdatement the eater is meroved to yield free (activity dostantice). Exterified strends are the short the strends are shorted more advection of the interactivity of the strends are blockdatement the eater is meroved to yield free (activity dostantice). Exterified strends are as from the tenter shorted correst on the interactivity of the strends of the strends are as from the strends are strends of the strends of the strends are strends of the strends are also as the strends of the strends of the strends of the strends are strends are as from the strends of the strends of the strends of the strends are strends are strends are strends are strends of the strends are strends of the strends are strends a life of drostanolone propionate is approximately two days after injection.

Side Effects (Estrogenic):

Dostandore is not anomatical by the body, and is not measurable estoppir. An articotogen is not necessary when using this struct, is a generoments abudd in do to a concereven armorg sensitive infinidata. Since ostrogen is the usale cleft with water restricts, dostandore isteaded produces a law, quality look to the physica with no far of coccess subcatances that reteriors. This multise is a forouble steroid to use during curring cycles, whow water and if treation are major concerns. As a non-assumable DHT detrointion, dostandone may impart an arti-struggine effect, the drug competing with other (anomalable) abstracts for buding to the aromatise enzyme.

Side Effects (Androgenic):

Altrody closefied as an analysic stroid, androgenic size effects are still possible with this substrace, operiodity with ligher than remain theoperic doeses. This may raked boards against emit polarity in the theory of the strong strong strong strong analysic interpolarity in the size. Women are surveiled for a polarity in straight stroid registration, emission is size its testic, fields and separating of the vock, mensitual impairing, charges in skin texture, fields in an growth, and clicical enlingment. Dostanolous is a started with relatively how androgenic activity relative to its stass-backling more androgenic against studies and strong strong strong strong strong strong strong strong Note that dostanolous is unaffected by the 3-sight needscase corpute, so its relative androgenic with the strong st

Side Effects (Hepatotoxicity):

Drostanolone is not c17-alpha alkylated, and not known to have hepatotoxic properties. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Ambicinnelogueis stervide can huve deletrizos effection serum chocksterol. This includes a tendency to reduce IRL (apod) chocksterol values, and increase IRL (bad) chocksterol values, with the IRL (apod) chocksterol values, and increase IRL (bad) chocksterol values, with the IRL (apod) chocksterol values, and increase IRL (bad) chocksterol values, and the effective and the service of the service of the service of the service and the service of the ser

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4) grams per day) and a natural cholesterolaritoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al arabok/androgenic steoids when taken in doess sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-simulating substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men);

Drostanolone propionte was not FDA approved for use in men. Prescribing galidhins are uravalable. For physique- or performance-enhancing purposes, this drug is usually injected three times per week. The total week/d okaga is typically 200-400 mg, which is taken for 6-12 weeks. This level of use is sufficient to provide measurable gains in lean muscle mass and strength.

Dostantone propionte is often combined with other stroxis for an extracted effect. Common stacks scalae in spicetable multioloc such as Deze-Danoloffe (mandhoot decanata) or Eigipionde (balenous stackeptmai), which can provide nearby enhanced significant strength and the strength and the strength and the strength and spicetable stoatestores. The result here can be soft muscle again, which is strength and other estrogenei skiller effects that the strength and the strength and other estrogenei skiller with other near-anomalies strengths strength and strength and other estrogenei skiller with other near-anomalies strengths scalae strength and strength and the strength and the strength and the strength and the strength and strength and the strength and the strength and the strength and the strength and strength and the strength and the strength and the strength and the strength and strength and the strength and the

Administration (Women):

The prescribing galdkins for Dobton recommended a dose of 100 mg given three times per week. Therapy is given for a minimum 610 k 100 weeks boftee mereduation of as efficacy is made. If accossifi, the drug may be contrand for as long as satisfactory reachs are obtained. Note that weaking suppressions extramined and data propagation of the strength of the strength meter of the propagation of profession scenarios mere the concentral of the concentral data provides it has a strength of the strength of the strength of the strength of the weaky dose gains and the strength of the strength of the strength of the strength of the weaky dosego is usually subdivided into smaller injections given once every second or third day.

Availability;

Drostanolone propionate is presently unavailable as a prescription drug product. All supplies of this drug come from export companies or underground steroid manufacturers.

Megagrisevit-Mono® (clostebol acetate)

| Androgenic | 25 |
|----------------------------|---|
| Anabolic | 46 |
| Standard | Testosterone |
| Chemical Names 4-chloro | 4-chloro-testosterone -androsten-3-one-17beta-ol |
| Estrogenic Activity | none |
| Progestational Activity | no data available (low) |

Description:

Chatebla cattete is an ambole starvaid that is derived from testssterone. Chatebla is 4chieva-teststorene, a modificanti nut matesia its stored al to streamful ambole compound with minimal androgene potency. This analog of teststorene is also net 17-alpha allyhied and does not anomatice, o there is lift worry of water credenci, angeconstati, or fertuacity during use. The hydrogen absolution at the 4 position does not grandly entrusce the mateginary of the streamful and the streamful and the streamful minigam basis, and granting presents. All hologen al activates of the poset androgen testsstorene, closeboli a centrality for removed from its parent streamful na centrality and streamful and the streamful method.

History:

Casteblo accette was first described in 1965.⁵⁷² If two developed into a medicine in Europe, where it was also a Starehard OF emirating. Gorenny and Turnhol (computure, GRR). This anabolic steroid had guerrally been indicated for the treatment of osteopronsis, alloward in the reproductive been used with a second strategies of a metric architeging amerecan and foor disease. Bush one and significant the strategies of the second strategies of the strategies of the second strategies of the second strategies of the second strategies of the strategies of the second strategies of the second strategies of the second strategies of the anabolic. The side effects of anabolicanthoganes are shown to be the next banch here.

Albuchg night from this in effect and patient construct, closebol acteuts was never a widely assessedial analosi, and assess on the instrumentation of the motics. As such as fatare would be a kenson one. The Tairnindol product from Karopharms would dispaperar by dood visuanis fratefact would be accessed as a such as a such as a such as the such as the such as a dood visuanis fratefact would be also such as a such as a such as a based product as a such as the such as a suc

Albudy the rarse functional rigicable preparations of this storoid as of the multic, clothed scetche is all manufactured in a marker of denami programions. The most recognizable such product has been Alls-Todekarmi form halo, allabudga it has albo been such as a backet (Mexico). Todekarmi (Chin, Landy, and Novadhur, and a desa include some comysists to help accelerate healing. The doses of storoid used in these products is gravely accelerate healing. The doses of storoid used in these products is gravely accelerate healing. The doses of storoid deleter, are not of much use to address, this storoid has even been related in certain ophilumbiogic outfines, which are even has pracediat to use for performance orbitrating proprints, and of hearing projects. The some most in the product remaining contributed dense runs for site address, we of storoid use corrections, the data correction, or contribute dense runs for site address, the source of the site proprints, and of hearing the site of the source of the source of the site of the source of

How Supplied:

Clostebol acetate is no longer available as a commercial oral or injectable agent. When produced (Steranabol) it contained 20 mg/ml of steroid in a 2 mL glass ampule or 15 mg per tablet.

Structural Characteristics:

Clostebol is a modified form of testosterone. It differs by the introduction of a hydroxyl group at carbon 4, which inhibits aromatization and reduces relative steroid androgenicity. Clostebol acetate contains clostebol modified with the addition of carboxyle acid seter (acetic acid) at the 17-beta hydroxyl group, so that the fires steroid is released more showly from the area of relaction.

Side Effects (Estrogenic):

Clustebi is not aromizical by the body, and is not measurably estrogenic. An artiv-estrogen is not necessary when using this storeduc, as genecomstain should not be a concern even arong sensitive infividuals. Since estrogen is the usual culprt with water reterion, clustebil instead produces a lean, quitaly bols to the physique with no face of ceness subscinancess that d reterion. This makes it a favorable steroid to use during cutting cycles, when water and far tereforma rempire concerns.

Side Effects (Androgenic):

Although classified as an antholic steroid, antrogenic side efficies are still possible with this substance. This may include botts of oily skin, acre, and bodyfacial hair growth. Anabloiciantfragenic steroids may also aggravate male pattern har loss. Women are also warned of the potential ivitizing effects of antaboliciantforgenic steroids. These may include a domainto of the notion moremult methodized models in data the field hard method. cospering or use voce, mensum requirements, cumpes n saci neare, can me grown, and clond enlargement. Additional, cloadeds is not extensively metabolical by the S-alpha reductise engrine, so is relative androgeneicly is negrody abserd by the concurrent use of instanctific or distance. Note that cloaded is a stored with two androgene activity relative instanction of another than the store of the store of the store of the store of the comparably higher than with more androgene agents such as testoatenees, metaneous stores, or discourses cores.

Side Effects (Hepatotoxicity):

Clostebol is not a c17-alpha alkylated compound, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

Side Effects (Cardiovascular):

Anabelicardingenis strends can have deletricos effects on serum chosterot. This includes a reducey to reduce III (agod) choksterov dusca and ircneres ILC, (bad) choksterov dusca serum chosterov ILC, (bad) choksterov dusca serum chosterov dus

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of star-tated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ois (4 grams per day) and a natural cholsterol/articoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al arabok'androgenic stevids when taken in dosses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimularing substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Chatchiotexte is generally used in chiral doses of 30 mg per week by ijection or 15 mg 2.3 times per dy only 10 the drug i admirated fr d 3 concercited weeks, follword by a break he's weeks. It is nearned at this point if indicated Electric doses for physics- or performance-artimetry paposes fail it the maps of 10-30 mg per yeek, lack for 6-1 week, lack for 6-1 week, lack for 6-1 weeks, lack for 6-1 wee

The ambide effect of this drag is thinly weak, so closels obtained a settine is most often utilized in constraints with other strends for a storage effect. The general publication is to use if the constest preparations with other non-aromatizing analobics such as Whistolle or coundrome. Here, a study does of 20 m gram be added in with maneigh does (20 m gram ger drag) of the oral anabolic, which negative should provide the user a new muscl built with the ord Primoloant's accuta anaplas, athrough Megaginevit is somewhat weaker is effect whost any user protein an addition of the ord Primoloant's accuta anaplas, athrough Megaginevit is somewhat weaker in effect. Store and a option of its user its compound in addition with theory more memorizable. androgens such as trenbolone, Halotestm®, or Provinon®. The result in such cases can be an even more pronounced effect of muscle definition, although this will be accompanied by a much stronger set of side effects.

Administration (Women):

Clustebulacettate is generally used in clinical doses 30 mg per woek by injection or 15 mg 2-3 times per day orably. The dang is administered for 3 consecutive weeks, followed by a break for 3 weeks. The dang is resurned at this point if indicated. Effective doses for physique-or performance-enhancing purposes fall in the range of 50-75 mg per week for the injectable, or 30-60 mg dashy for the on Latken for no longer than 6 weeks.

Availability:

Clostebol acetate is no longer available as a prescription agent at this time, and is unavailable on the black market.

Metandren (methyltestosterone)

| Androgenic | 94-130 |
|------------|--------------|
| Anabolic | 115-150 |
| Standard | Testosterone |

Chemical Names17b-hydroxy-17a-methyl-4-androsten-3-one 17alpha-methylandrost-4-en-3-one-17b-ol

| Estrogenic Activity | high |
|-------------------------|-----------------|
| Progestational Activity | not significant |

Description:

Medylestostoree is an orally available form of the primary multi antiogen tostostoree. Looking at the strink or of the storely, we see it is baschly just tostostoree with an added medyl group at the C17 alpha policy is a C17 alpha adjusted substance), which allows for finited and the store of the finited and store of the store, and tody methylestosterone is quale crude in comparison to muy of the often of the store assbacently developed. The action of the store of the store of the store assbacently developed. The action of the store of the store of the store assbacently developed. The action of the store of the store of the store assbacently developed. The action of the store of the st

History:

Melybestoreme was fire described in 1055.⁵²⁸ and use one of the first on alreadyness to be used in rickinal rendice (it folkows Provings the first on alreadyness to be main chinal use at the time of development was as an onal medication to replace toststoreme (and its authorizante). Allough the drug was adopted for a number of other uses, one that the standard standard to the tuments of oryposcilantial (used standard prognascy in renders not marking, osteopososis, and, rune recordly, female mensures (opporting the overall energy and scatal interest of the patient).

In addition to standard tables and capsakes, methylocitosterone has also been commercially approade in subliqueal or buccal tables. Mestadrum Linguest from Che Phermenendial Company were perhaps the most recognized, and were popularly sold from the 1950's to 1900's. Thisse tables were placed under the tongage on between the gam and check and left to dissolve, disburging the drug to credation with the mootes membranes. Pspassing the low-Shingain of tweat table, approximately doubles the bioscinality of methylocitosteneore, and also provides peak seves or drug rapidly (approximately 1 not a and ousing inseas) of 2 hours). Although Cho's Metandron Lingates are no longer commercially available, numerous other sublingualbuccal methylicitosteroric tables are still in production today.

Medgebastence remins a controvenial seroid. Albrogh is has a long history, and angubb a jastifiab softworted, it is no longer wickly out can be even being withchown from many darg markets. The Gamma Endocrine Society made an official statement that pharmicsis onse seven yours has the Molt Engon maters for above and the seven pharmonic and the seven yours has the Molt Engone matters followed as: The darg centures and dort preserver is. The potertail hepatoticity is usually clean as a reason, especially when longe-manadrengen therapy is contemplated. The one exception scenars to be the growing interest in an engo and methods tookset on the seven seven is the seven seven in the Molt seven is and seven the seven seven is seven to be seven as a reason, especially when longe-manadrengen therapy is contemplated. The one exception scenars to be the growing interest in an equivalent the seven is balanced as the first memoryal data and the seven seven is set of the seven is seven as the seven seven seven seven is seven as the first memoryal data and the seven seven seven is set of the seven se

How Supplied:

Methylestosterone is widely available in human drug markets. Composition and dosage may vary by country and manufacturer.

Structural Characteristics:

Methyhestosterone is a modified form of testosterone. It differs by the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during onal administration. As is typical with c17-alpha alkylation, the resulting steroid has lower anabolic activity than its parent testosterone.

Side Effects (Estrogenic):

Medgebastence is aromatically the body, and is high estoparts due to its concression to 17-aphan medgebasticals, a spritcice concession with high biological activity, 17-aghio medgebastical merges and the spritcing of the spritcing activity the spritcing medgebastical merges and the spritcing of the spritcing activity the spritsus of the same time water reference and becomes a problem, catasiga a roubble loss of menge definition and the spritcing activity of the spritcing activity of the medgebastical merges and the spritcing activity of the spritcing activity of the medgebastic activity of the spritcing activity of the spritcing activity of the merges definition and spritcing activity of the spritcing activity of the definition activity of the strength on spritcing activity of the spritcing activity of the definition activity of the strength on spritcing activity of the spritcing activity of the definition activity of the strength on spritcing activity of the spritcing activity of the strength on spritcing activity of the spritcing activity of the definition activity of the strength on spritcing activity of the spritcing activity of the spritcing activity of the strength on spritcing activity of the spritcing activity of the spritcing activity of the strength on spritcing activity of the spritcing activity of the sprincip activity of the strength on spritcing activity of the strength on spritcing activity of the strength on sprincip activity of the strength on sprincip activity activity of the strength on sprincip activity activity of the strength on sprincip activity activity activity of the strength on sprincip activity activity of the strength on sprincip activity activity of the strength on sprincip activity activity activity of the strength on sprincip activity activity of the strength on sprincip activity activity activity activity activity activity activity activity

Side Effects (Androgenic):

Medgetostoree is clouided as an androgen. Androgenie side effects are common with this solutance, and may include bost of only sink, area, and bodylink aline growth, Figher does are more likely to cause and side effects. Anabolicandrogenie strenkin my also aggestien mit galarnin likes. Those garcritically prove to mit galarning like and side increase aggressioness. Women are additionally sumped of the poterial withing effects of anabolicandrogenie strenks, fixed in large or the voice, mestual regularities, changes in skin teature, fixed in large orgin of the voice, mestual increases in strenks strenks, fixed in large orgin of the voice, mestual to storatore, mellybastores corrects to a more potert stored of its intencion with the 2adots of the correct star of this strenks of the correct star of this strenks of methys.

Side Effects (Hepatotoxicity):

Methyliestosterone is a c17-alpha alkylated compound. This alteration protects the drug from deactivation by the liver, allowing a very high percentage of the drug entry into the bloodstream following oral administration. (17-alpha) analylated anabolicantrogenic steroids can be hepatotoxic. Prolonged or high exposure may result in liver damage. In rare instances life-threatening dysfunction may develop. It is advisable to visit a physician periodically during each cycle to monitor liver function and overall health. Intake of c17-alpha adeplated steroids is commonly limited to 6-8 weeks, in an effort to avoid escalating liver strain.

Methylestostorow was the first cond served liked to beparite damage. This may be, in part, related to the early wedspendit us of the corroporal, as the deng garendly displays acceptable safety the lower event of reliable does. When taken at a does of 20 m gdbs for 5 weeks, hapatotaxicity, as measured by bromosphalekin (BSP) reterior, was low in or study.²⁶ In a sequent incredigion, a monitor of parties motical splittent BSP reterior and are only 2 weeks of therapy with o'ling dialy.²⁰ Score barc complications are more given the protect nuries in with more plots used anabodical angular combined on the second second second second second second and the methylestosterore, especially with high does and/or proparal antimistration periods.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anaboli-inarlogarie steroits can have deletrious effects on serum cholesterol. This includes a tendrey to reduce HJM. (good) cholesterol values and increase LDL (edu) cholesterol values, which may shift the HDL to LDL balance in a direction that fitoros greater risk of articroskerosis. The relative inpact of an anabolicantrogenic steroit on securan liptis is dependant on the dose, rotue of administration (oral vs. nijectable), type of steroid (commitable or non-commitable), and level of resistance to hepstin crutabolism.

Melybetostorone has a storog effect on the hypatic imangament of choksterol due to is structural resistance to how here backdown and rote of a ahiristation. Studies how demonstrated an approximate 33% decrease in HDL choksterol and a 30% increase in LDL choksterol with 0 (mpp etgs). ³These changes occearuls with 2-4 works of the initiation of therapy, and pensisted for 2 works after discontinuation of the drug Antholekindhoganic attention, and a support left worksidar hypotentially necessing the risk of cardiovaceuter disconse and myocardial infraretion.

To help reduce cardiovascular strain it is advised to mrintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

All araboléandrogenie steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-simulating substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenodotrophic hypogenodsm can develop secondrus to steroid abues, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁵²² This is caused by the fat-soluble nature of steroid hormones, which can also some of the drug to dissolve with undigsted distary fat, realening is absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

To treat androgen insufficiency, prescribing guidelines call for a daily dosage of 10-40 mg.

Ine cose is concert or yars, when animisence in stampgair or faccul torm in the ong words be used for contendel periods too leng as the patient's blootnay results (hypotencies), secure lipids, etc.) do net necessitate is discontinues. When used for comments used, that is not the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state of the state of the state of the state of the discontent of the state state of the stat

Administration (Women);

Medylestosteore is not wickly used with worms in clinical medicar. When applied, it is most often used as a secondary medication during inporteble breast encore, when other therapies have finded to produce a desinhle effect. The dosage used for this application can be as high as 200 mpt eddy. Low doss of medylestosteoren have been used in recortorism estimated astronges and 2.5 mg orthoghissatorene to Alexago of 1 hidder ped mpt inported and the second second second second second second transformations and 2.5 mg orthoghissatorene to Alexago of 1 hidder ped mpt inport energy, Biola, and overall welkess of the patient, as well as context and to increase clinical second second second second second second second to worms for physicap- or pedformac-enhancing proposes due to its storing and/ogrie rune and lendrov produce withing set effects.

Availability:

Pharmaceutical preparations containing methyltestosterone are fairly limited. In reviewing some of the remaining products and changes on the global pharmaceutical market, we have made the following observations.

Methylestosterone remains available in the United States. It is sold under the brand names Android tablets and Testred capsules from Våleant Pharmaceuticals. A generic is also available from Impax Labs.

Aburaihan makes a generic methyltestosterone product in Iran. It comes in 25 mg tablets, with 10 sealed in each foil and plastic blister.

Methandriol (methylandrostenediol)

| Androgenic | | 30-60 |
|------------------------------------|------------|--|
| Anabolic | | 20-60 |
| Standard | | Testosterone propionate |
| | | |
| Chemical Name | 17.alpha.n | nethylandrost, S.ene. 3 17. heta.dio |
| Chemical Name Estrogenic Activi | | nethylandrost-5-ene-3,17-beta-dio low to moderate |

Description:

Methylandnostenedial (methandnial for short) is an anabolic steroid derived from dihydrotestosterone. The drug itself is manufactured in two very distinct forms. The first is unseterified (straight) methylandrostenedia), which is used when making an oral medication with this steroid (althrough an injectable once existed in the U.S.). It is also found as esterified methylandrostenedial directoriante, which is presented as an injectable. The addee projectute esters in the injectable form extend the activity of the drug for several days. Braically, methandroi drugs are alreed of 17-ad-lytical forms of 5-androstenefold. Mentatrioi is closibilitied as a weak anabole with weak androgene properties. It als ossens to display ourse beel of estrogenic activity, miking this stored less ideal for during. The drug is generally considered to mill, and in one Widely operatar range boolwhaless and athless. Sometimes, however, it is used in place of other anabole/androgenic agents in bulking stacks when available.

History:

Melyhankontenciól swa frat describel in 1033.³³³ militagi tina a vey oli agart na fara as yutteria: anhois aressó are concretor. Melyhandrosteficiól swa devicieja fina a netkicie by Oganos, witch add it in the Linde Slatisa under the Steneticio Human drame in bolie onal (interplusionational) and injectific (melyhandrastaciatio) deprojennia). Fonawa a popular anabola: agort in the Linde Slatisa daring the 1900's Medlyhandrostenedio was assentially the fast storid perceited to live a mable spacemical developers. Endy product Rimatria described E1 and storid perceited to live a mable spacemical advelopers. Endy product Rimatria described E1, as 's asteroil which in use considerable of the mel hormore's fiscurhaling exists without to the same constraint of the mel hormore's fiscure haling exists without to the same constraint of the mel hormore's fiscure projectivismeta, anguine integra malary compression.⁴⁴

Early assessments of methylandrostendiel being primnily anthols in nutrae did not held up work with their extensive in in human. It was ocentally determined that in doess affects to protote weight gains is antibole properties were accompared by significant anthoganic activity. Ummids, this days would be viewed as one of oblinated matchings and the significant of the significant and the significant anthoganic activity. Ummids, the Matshois, and economy Miccontrated Reservation protoches. The Deco-Tambolism, and Matshois, and economy discontrated Reservation protoches. The other USLs band and generic forms of the drag would below as well, allough more different and generic forms of the drag would below as seen allough other to activity of the site of the other USLs band and generic forms of the drag would below as seen allough more more barrowing and the site of the other USLs band and generic forms of the drag would below as seen allough more more barrowing and the site of the

How Supplied:

Methylandrostendiol is available in select human and veterinary drug markets. Composition and dosage may vary by country and manufacturer.

Structural Characteristics:

Methydnotoseniol is a molified form of displotestatorene. It differs by: 1) the addition of a melly group at entor 1-al-phe to protect the hormone during cal administration and 2) the introduction of a double bott between earbors 5 and 6, which seems to increase the anabole strength of the storid (prothy by miking it resistant to methodism by 3hydrosystemic displaydnesses in adcetal macele sisse). Methydromotestenciol diproprimet contains methydromotenside in molidi with the addition of 2 carboyle acid carbos (proprime acid) at the 3-best and 17-beth placogl groups, which delay the relases of fee methydrostenciol from feet or mice of earboyle acid sectors.

Side Effects (Estrogenic):

Methylandosenkiol is not directly accorated by the body, although one of its known metholotis in methylandosenkion, the discontant strategies and the believed to have some inkerner entropying activity. It is, likewise, considered a weakly to molecularly entropying storied. Gyncormotis is possible daring troument, hat generally only depending on dose. Sensible individuals may need to addition an anti-estrogen such as Nodace? Ito minime related aids effects.

Side Effects (Androgenic):

Albudgi oden closified as an anbole stroid, medplandvostenčiol i sufficietly androgine: Androginesi de effects are conven with its solstater. This may ricked boost of oly site, area, and bodyficial har growth, Anabolezindrogene steroits may also aggroute may lapter har bits. Worten are swrand of the potential wirking effects of anabolezindrogene storoits. These may include a deepening of the voice, mentual insplantics, changes it solit testices that bits growth and change and engeneers. Note that methylandrozeneich is not affected by 5-alpta nolateuse, so the relative androgeneicy of this storoid is not affected by the concentre to or finastrike of antaserviti.

Side Effects (Hepatotoxicity):

Medylankostenciól is a (17-ajhra dalylatel component. This identicine protects the drug from datacitation by the loca, alavaiga vara vojel kaprezentaga of derig energy into the biodstraum fidaving on al administration. (17-ajhra alylated anabolicanhoganis strends can be hapatotoch: Probaged of high exposure my sceli hi for daringe. In nere instruslist distribution my doelway, hi a advashie to sisi a physician pecidically daring is controlly thind for 6 showle, in me in often to accid scaling line or instrusson control physical of 6 showle. In me in often to accid scaling line or instrustion of the drug may present sightly lass strain on the her by sociling the first pass methods more allowing allowing and upper est substratiol heratoxicity.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambeichningenis stronkis can huve debetrious effects on serum cheeksterd. This includes a tendency to rolex IEL (good) chebsteroi values and increase. IEL (bod) chebsteroi values, which may shift the HDL to LDL bahare in a direction that favors grouter risk of attroiceborsis. The relative impact of an anabolicanlargonis stread on serum hybis is dependint on the dose, route of administration (eral vs. injectibels, type of stead) (insumitable on consomnitable), and level of resistance to heptic metholismon structural existance to here breakdown and (with the eral) route of administrations, anabolicanlorgonis structures and the eral route of administrations, routes and chebsterio here breakdown and (with the eral) route of administrations, routes endochisin relatives and structure hypertrophy, all potentially increasing the risk or futurisocatar direstes and meycontalis inferetion.

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of statusted fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with shi oki (4) grams per day) and a natural cholesterolarinoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al arabok/androgenic steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalizing substances, testosterone levels should return to normal within 14menths of drug secession. Note that prolonged hypogenadorophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its bioavailability. This is caused by the fit-soluble matter of steroid hormones, which can allow some of the drug to dissolve with undigested distary fit, reducing its absorption from the gestroiristicated tract. For maximum utilization, oral forms of this steroid should be taken on an empty stormeth.

Administration (Men):

Early preserving galidhess for Standki lecommend a dosige of 25 mg sion: 10.5 times per week by ong bacacit of intramacidan entics. For physiques — operformance-attracting parposes, a hybrial dosage is in the range of 25-50 mg dayly for the oral form, and 200-400 mg per week with the injectable. In order to here pholo levels more one with the injectable, it is grannely administered once every three to fair days. Cycles grannely has far no more than 6 to 8 weeks, in an effort to minimise hepatoxicity and attain on the fore and chesterout values. This level of use is sufficient for modernte gains in matce size and strendt, which may be accompanied by a buy level of water networks.

Whit's may be possible to use methylanchoatencials alone for much-bailing parposes, it is not often combined with other analysis for a stronger effect. Combined with Decam Darshoftsfor Expiposes/E, for example, measurable gains of hard mascle mass, without an eatenne level of varies tearrism, may be noted. This is the general composition of most Australian vet blench that include methylanchoateneddo. When looking for a more pronounced gain in muss, a stronger androgen such as testsbarene my be added. The resulting growth cam be qale exceptional, but the uses and gains of the stronger and the stronger and the stronger and the stronger stronger and the stronger s

Administration (Women):

Early prescribing gaidelines for Stenediol recommend a dosage of 25 mg given 2 to 5 times per week by onl, buccal, or intramscular route. Methylandrostenediol is generally not recommended for women for physique- or performance-enhancing purposes due to its androgenic nature and tendency to produce winking side effects.

Availability:

Pharmaceutical preparations containing methylandrostenediol remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

The only place where this steroid is still produced in an volume is Australia, where a number of veterinary preparations still include methandriol in their blends. These products are rarely traded in international commerce due to tight controls on anabolis steroids in that country.

Metribolone (methyltrienolone)

| Androgenic | 6,000-7,000 |
|-------------------------|---|
| Anabolic | 12,000-30,000 |
| Standard | Methyltestosterone (oral) |
| Chemical Names | 17alpha-methyl-17beta- hydroxyestra-4,9,11-triene-3-one 17alpha-methyl-trenbolone |
| Estrogenic Activity | none |
| Progestational Activity | no data available |

Description:

Methyltinokus is one of the strongest end anabols steroids ever produced. This agent is a deviative of prethydrox (trienchox), which has been C-17 dpa has plated to alsow for end administration. This medification has created a strond that is significantly storager than is non-methylated coastin its potecyr has been reasored to be anybeen form 123-300 items greater than that of methylatestorenee, with greater discostion between anabols and analogonie effects.²⁵⁵ 35⁶⁶ Migman for miligram methylateshoots is none achte steroid than any agert sold on the commercial method, requiring doors so lithe as 5-1 miligram per dw to troit as a trans anabole effect. Its potency is ony mated to by its relative toxice). however, which has limited its modern use to that of laboratory research only.

History:

Meightinoshe was first descredu in 1965.³⁷¹ It was immediately identified as an enternely potent analysis agart, fir mere potent han the conversively available agarts for the poster than the conversively available agarts for the poster of the site. In space of a high editive activity, however, meldylinendoue has seen very limited use in humans. It was calculated human factors and the poster of the advanced breast cancer. Here, is executingly atomating ambient agarts for a distance of the site of endagerous senses, huffing a good atomatic the lead of endagerous senses, huffing a good atomatic endagerous and the site of endagerous senses, huffing a good human endicise.

By the mid-1070s, methylinizolore was becoming an accepted standard in non-human research statics, periodically those perimited to be study of the androgon receptor actively. For this parsons the agars is very well stated. In shear potency and resistance to semanagence is the state of the aggregative between evolution to metaboles are chose one choice on the aggregative between evolution to metaboles are inclusion state of the the quastion of what is causing a particular effect, the state of the state of

How Supplied:

Methyltrienolone is not available as a commercial agent.

Structural Characteristics:

Metybinolous is a modified from of mardrown. It differs by: 1) the addition of a methyl group at atooh 1–7 abolic to protoce the horner earling cal administration and 2) the introduction of double bonds at carbons 0 and 11, which increases its binding affiny and aboves its matcholium. The resulting storic is a significantly more proteins in comparison. Methylenison chemically differs from trencholen on the third store comparison. Methylenison chemically differs from trencholen on the phylical methyl group at c-17. This alternion changes the activity of methylinearbox considerable, business that a store of the considerable of the more than the store of the

Side Effects (Estrogenic):

Medphirokone is net anomated by the body, and is not measurable settogenic. It is of none, however, that medphirokone doubly say significant brinding affing for the progestores receptor.⁴⁴⁷ The side effects associated with progestores are similar to those of estrogenticating negative desoluti inhibits of testostrome production and enhanced rate of this storage. Progestins also augment the similarity effect of estrogenes on manmary tissue growth. These approxes to be a storagy sequery between thus two to horness here, such that Drate of an anti-estrogene, which inhibits the estrogenic compenent of this disorder, is often atticient to mitigge accounsils assead by progestional anabolications/green is storage.

Side Effects (Androgenic):

Albudge closified as an ambioit seroid, antiogenis side efficies are sill common with this solution. This may include boats of 60% skin, acce, and bodyficial lini growth. Arabolicitudingenis steoids may also aggravate multi pattern har loss. Women are also sourced of the potential withing effects of antibolicitudingenis across. These may include a and clinnal enhangement. Additionably, the 5-alphan nebatrasic enzyme does not methode mothylinic endyrinistic and the 5-alphan nebatrasic enzyme does not methode mothylinic enzyme. Additionably the 5-alphan nebatrasic enzyme does not methode and clinnal enhangement. Additionably the 5-alphan nebatrasic enzyme does not methode and endyrinizations. So in earbier antiogeneity is not affected by bisocritic or datasetristic.

Side Effects (Hepatotoxicity):

sease and many many sease and a sease a

Attemptionscore is a c1-spin asymptotecomponent. Insistentine protects me ange generation of the set rabitivity as very link proceedings of the drug entry into the blockdenean Biblivity could administration. C17-spin kakytated anabolicitationgenis stretchic and be hapstoalose. To blockade and the protections, the changes II may estimate the set of the spin stretchick and the spin stretc

Methylinoidore is an eccocingly poter onl storiul with a very high level of resistance to h popurin entabolism. This makes methylinoinolo reaccending hove-toxic, procluding is use as a prescription agent at this time, in any part of the world. Studies publicad ion the University of Borne Commeny back. In 1966 multe this very class? This fact, at this time researchers had deemed this the most hive-toxic activity of very be statisfied in harmurs, samming up their finging well what stating.

"Methytienedoec... which is onally active as an anabolic agent in a done less fam 1.0 mg per day in normal adulta, base herst sied with regard to is induce on her function. As measured by mathyle parameters (ISP retention; total bilinibit; accivities of transmisses, adulta photophitas and doublescenses in serural activity of protocolerin in photon induced cablestask....thus methylemother as to causing bacterized supports intelligent cablestask...thus methylemother as protocolering for most Psystemic serued."

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambelicantogenis stavalis can huve delutricos effecti on serum chesterot. This includes a nelmery to reduce ML (apod) chestavol values and increase LL, (bad) cheststead values, which may daft the HDL to LDL balance in a direction that florons gener risk of a neuroicolarosis. The methical regate of an analocicantrogenic around on serum highs is (anomizable or non-sourcitable), and level of resistance to bepatie methodsan. Although run extensively studies intranse, the cal model of resistance to bepatie methodsan. Although runce on directly historypair risk. Although and the site of the si

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of statasted fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ols (4) grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All araboléandrogenie steoids when taken in doess sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalating substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondary to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral ambolic steroid with food may decrease its biavailability.⁴⁰⁰ This is caused by the fait-soluble rature of steroid hormones, which can also some of the drug to dissolve with undigested dietary fait, reducing its absorption from the gastrointestinal tract. For maximum utilization, methybrienolone should be taken on an empty someth.

Administration (Men):

Medphirokoles is no lengre used a clinical medicine due to an uncoceptable level of hepatoxicis; The same reason. Those absolutely insisting on its use need to take as level of here toxicis; very seriously. At the wey least, routine block tass should be conducted to ensure the agent is not impuring damage. Drug damatin should also be very limited, predenibly to verkeds, of an er lists. The relative potent of methylenessing or instance of the same reason. The second second second second transmitted is the second sec

Administration (Women):

Methylnienolone is no longer used in clinical medicine due to an unacceptable level of hepatotoxicity. This agent is not recommended for women for physique- or performanceerhancing purposes due to its extremely strong toxicity and tendency to produce virilizing side effects.

Availability:

Methyliariohne is not produced as a prescription steroid product in any part of the work!. With the rapid expansion of underground steroid manufactures, this agent has been released as a black match designer compound. Those contemplating the use of underground forms of methylinenohors should consider that such agents are being released for haman use without any government approval or consideration to its safety.

Miotolan® (furazabol)

| Androgenic | 73-94 |
|-------------------------|---|
| Anabolic | 270-330 |
| Standard | Methyltestosterone (oral) |
| Chemical Name | 17-Methyl-5alpha-androstano [2,3-c]furazan-17beta-ol |
| Estrogenic Activity | none |
| Progestational Activity | no data available (low) |

Description:

Franzabit is an oral anabolic strend derived from dhydotestoteeven: The agent is modeling and the whole with an dhydotestoteeven is no oddet due to be medification of the steroid's A-ring which allows the steroid structure to remain stable and Dhydotestotesterees. The cooperation, is a power number, quickly mathoded in muck these to incredit entropy of the product of the steroid structure of the stars of the stars of the stars of the stars of stars and the stars of stars of stars of the stars of stars of

History:

Furazabol was first described in 1965.⁵⁴¹ The only modern pharmaceutical preparation of record containing finzabol, at least known to researchers in the West, was Mistolan from Davids Scialent I ada in Jawan utilely tase could in Jawan onitide duries the 1070°s and '800° ° Lenter expande Later to acquire thread when a source measurement using use 17 0.7 s and 100.5. The agent keell's accordy mentioned in the Wisstem medical Breature, and consequently a great deal of mpth has come to surround it among athletes. A realistic appraisal sits this agent in a very similar class to stanzould, however, with both agents being medicately strong ambolies with low androgenic activity. Aside from this, it is difficult to ascribe any drastically using term its the dag.

Frazabio was a popular steroit armog Ohyme'a nhikes during the 1980's, when it was acjuch known amore coint niners of the streng for finals had vay tei whiteful for larger, and therefore coil not toris for a LD Jamic Astaphan, the physican that accompanied Ban Ashanon to the 1980's Ohyme's in Social spreadering was gived globrous data metros coile the athletes at the time's largering the data was and any the detectual to the data gives in distances. White now spreads for strength and the strength and gives in distances. White now spreads for the data strength and the wave publicad, immediately dimensing any value fins agent formerly possessed as a storoid undetectual be of agencement.

Today, farazho is vezy saczedy known to bodybaikes. The Motohu brand from Japan wa discortined mury usors aga, and no pharmencidia peperatine cortaining farazhol hus been known to exist size. The drug is occasionally located on the black mutet, however, due to be far thin is st all produced in black sac an automaticaturing operations in the West, and produced in our a labels and capacity. Contrastly the autoparations are been and produced in the size and automaticaturing operations in the West, and produced in our a labels and capacity. Contrastly the attalmented capacity of the size in the size of a labels and capacity of the size of the six

How Supplied:

Furazabol is no longer available as a prescription drug preparation. When sold it came in the form of tablets containing 1 mg of steroid.

Structural Characteristics:

Fuzzaboli is a modified ferm of dipdrotestosterone. It differs by: 1) the addition of a methyl group at eachon 17-apht, witch helps protect the hormove during ond administration, and 2) the attachment of a farzar group to the A-ring, replacing the normal 3-keto group. When viewed in the light of 17-apht methyldhydtostostosteron, the A-ring multification on furzaboli scemes to considerably increase its anabolic strength while reducing its relative antrogenity.

Side Effects (Estrogenic):

Fuzzabol is not aromatized by the body, and is not measurable settogenic. An arriv-strogen is not necessary when using this strong of a segment should not be a concern even among sensitive individuals. Since settogen is the usual cupirit with water retention, this strougli strated produces a lear, quality look to the physique with no far of excess subcatancous fluid reterition. This makes it a floronable steroid to use during entiring cycles, when water and far teretrion are mission concerns.

Side Effects (Androgenic):

Although costified as an analysis steorid, androgenic size effects are still possible vulti fields substratece, opercivity with higher doess. This may include boots of op skin, acre, and body/fiscial imar growth. Analysical androgenic steorids may also aggrostee mile pattern hair boss. Wormen are wanted for the potenti vultiming effects of analysicalizationarigeness tearoids to the strategistic strategistic strategistics. These may include a strategistics, changes in situ nature, final hair growth, and choolen elangement. Franzabol is a streated with relatively but androgenic activity relative to its issue-building actions, making the fravelul for strategistics. These may include a stream of the potentiary business. Note that fravels is a malited up the 5-shipher roltance is a relative and rougenicity is not affected by the concurrent use of fravestive of constanction.

Side Effects (Hepatotoxicity):

Fuzzabol is a (17-alpha albahad compound. This alteration protects the drug form descritation by the local albahad set of the right power parage of the drug entry into the bloodenteum bibwing could administration. <math>(17-alpha albahad could be albahad bahad bahad

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambelicantogramis strends can huve deterious effects on serum chesterot. This includes a tendency to relace IEEL (good) chesteron values and increase IEL. (bod) chesteront values, which may shift the HDL to LDL bahare in a direction that fivors grouter risk of attrivicherosis. The relative prant of an anabolicantlogonis extraol on serum hybis is dependint on the dose, roote of administration (equal vs. injectedbel, type of stead) forgenuitable or measuremathely, and loce of orsisteroi to hysicant methods in Tranzido mana. startiat resisteros to Sure bacaldonan, and roote of administration endoclicitation of the doses of the starting hypertraphy, and posterially researing the risk or distroscent fire starts and methodan in Tranzido mana. startiation of the starting of the starting hypertraphy, and posterially researing the risk ordinoscent af mease and meycandin infertion.

Note that franzhoi is often risikalený describel sa s steroit with urage beneficial cholserol-loverig effects. Such statements usaly máse reference of radius conducted in the early (1970), which examined the pick-lovering effects of the agent 425 Such a position, however, lacks a mode prospective of the drag. To draw a panell, draing the early (1970) where was research dnee no coandrobre, demonstrating a lipid-lovering telfs. ⁴²⁵ Upon clover inspection, however, it was a multile, drawing the early (1970), which early the DR-LDI, ratio and athropperior field. Such as the based of the drag was remained. The same is the for fazzable Same have good as one have good as one should be showed for the drag was remained. The same is the for fazzable Same have good as o worked.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of star-tated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ois (4 grams per day) and a natural cholsterol/articixelar formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

All arabok'androgenic steavils when taken in doese sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalizing substance, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadorophic hypogenadism can develop secondrus to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

An efficite dosage of frazable seems to begin in the range of 10-20 mg duby for mer, taken for no longer than 6 of weeks. At this level is sense to input, and manage the building effect, which is usually accompanied by fut host and increased definition. Does of 30 mg per day or more considerably arccases the analobed potential for due, but at its expense of general hepationskies). The mark-building activity of firzabilic could, instead, be trappinger to the sense of the sense of the sense of the sense of the sense first provide the sense of the sense of the sense of the sense of the sense first provide the sense of the sense in the sense of the sense in the sense in sense of the sense in the sense in sense the sense in the sense is the sense in the sense in the sense is the sense in the sense in the sense is the accompanies by some level of water retention, and potentially a decrease in muscle definition.

Administration (Women):

In the athletic arena, an effective oral daily dosage would fall in the range of 2-5 mg, taken in cycles lasting no more than 4-6 weeks to minimize the chance for virilization. As with all steroids, virilizing side effects are still possible in women, but remain rare with conservative dosing.

Availability:

Furazabol is no longer produced as a prescription drug product, although underground preparations containing this steroid may be located.

Myagen (bolasterone)

| Androgenic | 300 |
|--|--|
| Anabolic | 575 |
| Standard | Methyltestosterone (oral) |
| | |
| Chemical Names 17beta-Hydroxy | |
| Chemical Names 17beta-Hydroxy | -7,17aipha-dimethylandrost-4-en-3-one 7,17-dimethyltestosterone |
| Chemical Names 17beta-Hydroxy Estrogenic Activity | |

Description:

Bolsstores is an onli anaboli steroid structurally related to methylestostorese. It differs only by the addition of a methyl group or 1,7 which accounts for its given chemical many 1,71-dimethylestostosterese. The added <7 methyl group makes the activity of this attential for given the structural differs of the structure of the structural differs of t

History:

Bolustrove was first described in 1959-5⁴² It was choody evaluated for ambolic and androgenic effect graphical fields and the start of the start o

How Supplied:

Bolasterone is no longer available as a prescription drug product.

Structural Characteristics:

Bolssterene is a multified form of testosterone. It differs by: 1) the addition of a methyl group at carbon 7-aphyle, which helps protect the hormone charge real administration, and 2) the introduction of a methyl group at carbon 7 (alpha), which inhibits 5-aphyn reduction and shifts the arabolic to androgener ation in flow of the former, 271-dimethylated storeside also test biological activity.

Side Effects (Estrogenic):

Bolsterore is arounized by the body, and is considered a lighty estrogenic stored due to its conversion to 7.7.6 methylostandlo (m estrogen with high bodycal archive). Gyncormolis may be a costeen during teatment, especially when higher thran neural an archick host of marchive characteristic and the store of the store of the transformation of the store of the store of the store of the store of the transformation of the store of the store of the store of the store of the Nodesley. The store effects errorshy for saturgen control Anomatos inhibitors, however, can have martine effects errorshy for saturgen control. Anomatos inhibitors, however, can how ensuring effects on body fusion.

Side Effects (Androgenic):

Albragh bolaterone is clossified as an analosi storedi, androgarie side effects are still growth. Higher doess are more likely to cases such aids effects. Attable/inflogarie storedistic and analysis of the stored storedistic and the potential viriange direct stored stored stored storedistic and the potential viriange effects of analodicardrogarie storedist. These may teckles a deepening of the score, ensettant lineage/miss, changes as the store, field in growth, and change enlargement is installed by the concurrent set of instantice of adaptation. It is a stored to the attached by the concurrent set of adaptation of adaptations of the store of the store of adaptation of the store of the store of the adaptation of the store of the store of the adaptation of the store of the store of the adaptation of the store of the store of the adaptation of the store of the store of the store of the adaptation of the store of the store of the adaptation of the store o

Side Effects (Hepatotoxicity):

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambiciant/arganis stroutis can have delutricoss effects on serum chesterof. This includes a tendency to rolesci IEUL (god) chieskeroi values and increase. JUL (bud) chieskeroi values, which may shift the IEUL to LDL bahare in a direction that fivors greater risk of antiroiccheosis. The realise impact of an antibolicant/gongis extraol on serum (pits) is dependant on the dose, route of administration (end) w. igiextabels, type of steadi (commitable on more-anomizable), and level of resistance to hepain: metabolism. Bolistoren has a strong effect on the hepain: management of cholsterol due to is structural resistance to her treadous and note of chimistation. Ambiokardongeris streadings also adversely affect blocd pressure and trighcerisfies, reduce endotfekal relaxions, and support fit vertical inspertophys, all poterbils pressures the risk of catationsacchar disease and rescontial inferiors. Studies administering Img and 2mg of bolasterose dialy for 6 weeks in 27 printed historic disease and the rescuesal series relaxional studies and reader of administration that bolasterose significantly shifted the ratio of fasce too factors of the rescarbox printed and the reader of these two factors of the relaxion of the bolasterose significantly shifted the ratio of fasce too factors of robusterio father agent the source significantly shifted the ratio of fasce too factors of robusterio father agent the source significantly shifted the ratio of fasce too factors of robusterio father agent the source significantly shifted the ratio of fasce too factors of robusterio father agent the source significant processies risk.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with sh ohs (4, grams per day) and a natural cholesterolaritoxidant formals such as Lipid Stabil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

All arabok'androgenic steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimuling substance, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondrus to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴⁴⁷ This is caused by the fat-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undigested dictary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

Clinical studies have demonstrated that significant nitrogen reterion and weight gain can be induced with a daily dosage of 1-2mg per day. In the athletic arena, doses of 2-5 mg daily seem to be most reasonable, taken in cycles lasting no more than 6-8 works in length to minimize hepatotoxidy. This level is sufficient for strong increases in masch size and strength, athlough such gains will keel be accompared by significant water reterion.

Administration (Women):

Bolsterene vas not videly used with vonnen in clinical medicine. When applied, it vas most often used as a secondary medication during inopenale hereact cancer, when other thrappies have finded to produce a desinable effect. The donage used for this applications would be as high as 10 m gap edge, as leads that the cassed signifacture, vitationan among patients, enders the second s

Availability:

Bolasterone is no longer produced as a prescription drug, although a handful of underground laboratories have taken to selling this material.

Nebido (testosterone undecanoate)

| Androgenic | 100 |
|------------|----------|
| Anabolic | 100 |
| Standard | Standard |

| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one | |
|----------------------|--|--|
| Estrogenic Activity | moderate | |
| Progestational Activ | ity low | |

Description:

Nebiolo is an njectable steroid that contains testostoreor uncleanoute, a very solw-sering coster of loststoreor. Inis is the active durp that is used in Autrich, bin in that case is is part of an onli medication, not an ispeciable. Nebiolo is being markedet as a replacement for established injectable. Its beststoreory products the Delatority(B), Department, Sentandi, B), and Sentandi, B), and Sentandi, B), and the sentime strateging in comparison. It is designed to offer a much loss foqueri reficion schedule, and, therefore, much aparet confort for the parter. Nebio is a drug developed tarder a simile focus as teststoreore bucklet, which is another very slow-acting ingelishest eard reflaststoreore.

History:

Nebido' was developed by itemativanj gint Schering AG, Germany (now Boyer). It first sarteed as a prescription dup in Friadu and Carmung in Cuebes and November of 2004, nespectively. Within a year it had been approved for sult froughout Enzye. Schering/Bayer has since also brough it is product to Moxics, Road, Aggerins, Schoth Affex, Colombi, Korea, Vancash, and various countries in Eastern Enzye (86 countries in total). In July 2005, the U.S. planneeukais firm indexes periodeal the rights noneex, singericina maphyloics: exections and pathromy of nicroscobism has of the targ, however, argicelin marphyloics: exections and pathromy of nicroscobism has too of these reactions are likely not date to a problem with the drug ledf. Ha fractorest administration of the highvolume negistoric Future approach in the gives a sepected, but rations:

Nebidoli was described by Scherig as being het, "Fast long-acting injection for the nontimot of mike poperations". This may be nattice of poperoders, and or show-acting constraints for Nebidogia how the structure of the structure of the structure term structure of points of the structure of the structure of the structure methods for subsciences. The structure of the structure of the structure of the confirst domains and how the structure of the structure of the confirst domains and how the structure of the structure of the structure part of the structure of the structure of the structure of the structure part of the structure operation of the structure of the s

How Supplied:

Testosterone undecanoate (rijection) is available in various human drug markets. All products (Nebido®) contain 250 mg/ml of steroid dissolved in oit packaged in 4ml ampules containing 1,000 mg of steroid in total.

Structural Characteristics:

Teststere urdeanose is a mellifed firm of teststerene, where a carboxyle and eser (indecanic acid) has been attheted to be in-To-het photogy grants. Estimited firms of teststerene are isos poler than fee teststerene, and are absorbed more slowly from the are of njecticn. One in the blodsterms the ester is removed to yield for (acido) teststerene. Estimited firms of teststerene are designed to prolong the window of therpartie effect blogway administration, adways for a lass Seguera tpicetion schedule compared to spectrose of fite (meastrind) steroil. Hobitoli is designed to maintain theological layed bettesterene for to 1 y weeks after interior. ⁴⁵⁶

Side Effects (Estrogenic):

Testosterone is readily aromatized in the body to estradiol (estrogen). The aromatase (estrogen synthetase) enzyme is responsible for this metabolism of testosterone. Elevated somgan beek can cause side effects such as increased water retention, body fut gain, and gencomstain. Testatheone is considered and moderably estrogenise steroid. An artis-entrogen such as compleme critate or transmitte influent flex Artimeted (manistroad), which more effective controls estrogen by preventing its synthesis. Anomatos estibution can be effects on the major and the synthesis and and and and the sentences of the effects on the major and the synthesis. Anomatos estibution can be effects on the only plass.

Enorgenic side effects will occur in a dose-dependant mumer, with higher doses (above normal therapetic level) of leststoreme more fieldy to regard the concurrent use of an artiestrogen or aromatose withhole. Since water referring and less of matecide definition are common with higher doses of feststoremer, fide drugs is made and are that larbeing the single site of the backing places are backed using a more site of the site of th

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for minitaing scoradary male senal characteristics. Encodel levels of teostores are levely on produce androgenic side effects including objects, and bodyficial lar growth. Men with a genetic professionis for link too (androgenic altery and teoperation) my conce concentrate discourlanding. These concerned about har loss may find a marc confinished option in manufacture about the structure of the structure. These may include depending of the voice, mentional incipationics, changes in whit concerns, facility around an director datary market of the structure of t

In anthogon-responsive target tissues such as the skin, such, and prostate, the high relative antogenizely of stratures is dopedual on its rolation to display disordisationes (DHT). The 5-alpha reductise corputs is responsible for this metabolism of tastistatives. The reconcent use of 3-able inclusions within works and instanties of interface with also-specific potentiation of tastistatives and interface of interface with also-specific potentiation of tastistatives and interface integration of tastistatives and the strategies and the strategies and segmention of tastistatives inhibits.

Side Effects (Hepatotoxicity):

Testosterne does not have hepatotoxie effects, her toxicity is utilikely. One study earning the potertial for hepatoxicity with high does of testosterne by administering 400 mg of the hormone per day (2,800 mg per week) to a group of multi adjuent. The starod was composed to intramediar injections. The hormore was given days for 2 days, and produced no significant charges in Fore enzyme indus including neural models.

Side Effects (Cardiovascular):

Ambolicanteganis strouids can huve debetrious effection on serum chocketorel. This includes a tendency to reduce MEL (apod) chocketorel voluces and increases (LL, (bad) chocketorel voluces, which may shift the HDL to LDL bahare in a direction that fivors genet risk of attractionetors. The reducite impact of an analobicantideganis extraordio on serum (hpic) dependant on the dose, route or a daministration (et al), wijectedbal, type of steaded Anabholicanteganis extraording and the steader of the steader of the steader Anabholicanteganis extraording and support left bear travers and a trajectorials researcing the fiel or chimoscatar direstes and mey contrails prioritally increasing the fiel or chimoscatar direstes and mey contrails affection.

Testosterone tends to have a much less dramatic inpact on cardiovascular risk factors than synthetic steroids. This is due in part to its openness to metabolism by the liver, which allows it to have less effect on the hepatic management of choksterol. The arounization of testosterone to estatial also heps to mitigate the negative effects of androgens on serum high la none which 200 new new old of transformation areas (management) held a dida have and the last testosterone to estatial also heps to mitigate the negative effects of androgens on serum table. It may need that the setup of the networks of the last test held a dida have and the networks of the n puts. In one same, across the process on sessence one constraints of the set of the set

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of statutated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish ols (4, grams per day) and a natural cholsterolaritoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencials when taken in doess sufficient to promete make gain are expected to appresse condigenses testsatores production. Testsatores is the primary mark androgen, and offers strong negative feedback on endogenous testsatores production. Testsatores-based dange voll, Revise, here, are strong effect on the hopoltamiar regulation of natural steroid hermores. Without the interversion of testsatores estimating adottances, testsatores levels doubt term to room with with 1-1 months after the dange las dy cleaned the body. Nose that prolonged hopogeneidstropic hopogeneidsmic and develop secondary to storial abase, necessiting medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Due to the large injection volume, prescribing guidelines recommend that each injection be given slowly, taking approximately 60 seconds to administer the full 4ml dose. Nebido® should always be injected doep in the gluteal muscle.

Administration (Men):

To text androgm instificiency, the prescribing gaidkines for testosterour undecantee (Netholds) call for a doage of 1,000 rm (efm every tweb weaks. Therary is usual initiated with a loading pluse, which requires that the second ispection of 1,000 rm [be given at approximatly his wives draw. If no boldballing proposes, supprisolugibal (influent fung plass) doagoed weaks weak and the second second region a more regular basis. The most logad protocol would be to administra 4 del ispection of bolds one 27 deweeds, for an approximate average weakly doage of 2500 rm of leaststoree eater. At a loss fixingent justice mobile with a fixer acting cases, such as current to the loss fixing the induktion the network with a fixer acting cases, such as current to coptionet, which weakly done testosterow testo is rough it may approprised.

Administration (Women):

Testosterone undecanoate is not approved for use with women in clinical medicine. This drug

is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature, tendency to produce vinizing side effects, and very slow acting characteristics (making blood levels difficult to control).

Availability:

Testosterone undecanoate injection continues to increase in prominence as a pharmaceutical product. It is presently approved for sale in 86 countries worldwide. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have mude the following observations.

Nebido gained approval for Europe-wide sales in 2005. The product is has since distributed throughout Europe, and is widely available in this region.

Indevus, a subsidiary of Endo Pharmaceuticals, has continued to push for FDA approval of Aveed in the United States, but such approval has not yet been granted.

Nilevar® (norethandrolone)

| Androgenic | 22-55 |
|----------------|---|
| Anabolic | 100-200 |
| Standard | Methyltestosterone (oral) |
| Chemical Names | 17alnha-Ethyl.17heta-hydroxyectr.4.en.3.one |

| | 17a-ethyl-19-nortestosterone |
|---------------------|------------------------------|
| Estrogenic Activity | r high |
| Progestational Act | ivity high |

Description:

Needminokone is an anhole staroid closely related to netrestosterone (marcholoue) in strutter. The active/for files strucils if not a mill to modente and anhole seroot, which is accompanied by distinguishible androgenic and entrogenic components. Although this structure is a structure introduction million (distillation) to make call doing with it. Carmothe looked at simply as an onal abrumitse to Decar-Daraboliti. Most netably, the greatly increased estrugging cased by 17-adhytikin mildess methatholese muth more problemic when trying to ball quality (lam) mach mass. In administering an effective encogene is de effects. The march accumulation with northendolese is also given to be accompanied by a high level of water and (Ridoy) fat netrotion, to the quality mascharity neumaly associative dynamical matching and the structure.

History:

Noreflunckolne uses first described in 1954²⁰ T to use developed into a medicine by Stork, which medicated is into fuel. Speceroption durg metter under the Nikour barrel nume during the last 19605. The data yaus originally solit as a coali subdim, methematic strategies and the strategies of the strategies of the strategies of the spectra of the strategies of the strategies of the strategies of the given by injection. Nikour was proceeded for a variety of thesess that were benefind by a protein speciar moleka gard. Lada relations included preparations for and recovery from strategies of the substrategies of the strategies of the stra

Norethandrolone ultimately saw only limited success as a prescription anabolic agent. It did make its way to Europe and certain other markets, but not wildely. The drug was an early functional anabolic, displaying more tissue-building extrogenic stile. This eventually led to also remained an agent with a troubling estrogenic side. This eventually led to the second state of the second state of the second state of the second state. neethandokee being passed over clinically for more refined components as they became available. Scarb detailed to discortisme tassi dorNolar in the US, during the 1960's, and instead began focusing energies on its newer, more strongly ambolis, and non-strongent stroid coandrolore (edd) a Anvani). Most other markets carring neethandoknee, either by Scarb or other comparists, none began long this compound as well. Tables, this drug is vertinary dimension, and the strong task of the strong task of the strong the strong task of the strong vertinary dimension.

How Supplied:

Norethandrolone is available in select veterinary drug markets. Composition and dosage may vary by country and manufacturer, but typically contain 5 or 10 mg of steroid per tablet.

Structural Characteristics:

Norethandrolone is a modified form of nandrolone. It differs by the addition of an ethyl group at carbon 17-alpha to protect the hormone during oral administration.

Side Effects (Estrogenic):

Norefination have a arrantized by the body, and converts to a synthetic entropy with a light lead of biological activity (Tapher-offscherafts), kas a read, it is a laphy entropy should. (Proceemist is often a convert during transmer, and may present heif quice any however, a problem, cancing a norbit less on more definition and the observations out enterior and fit looks badd. Seminive individuals may want to keep the strangen under conclowith the addition of an anti-strengen state in Nobided.¹⁰ Concolivability addition of an anti-strengen state in Nobided.¹⁰ Concolivability addition of an anti-strengen state in Nobided.¹⁰ Constrained to the Arimediced (manistrated), which is a more effective neuroly farstimation entropy-matience therapies, and may also how neights: (Note State 1) and the strength of anisotation entropy-matience therapies, and may also how neights (Note State 1).

It is of rote that northandboles thesis some additional activity as a progessin in the body. ⁵⁴⁴ How side effects associated with programmer are winither in these of entrange, including regardies faceback inhibition of testosternee production and enhanced rate of fits strenge. Progestins also agreement the similatory effect of entrogens in advancement of the similatory effect of entrogens and the strenges of the similatory effect of entrogens in advancement of the similatory effect without encodes to entropy including the activity and the entropy of the similatory effect on the hole of programs involved test processing similar to insign evencounts in the hole of programs involved test procession exceeds by uncertainable of the similary effect on trainger exercises and which are exceeds entropy which inhibits the estrogenic component of this disorder, is often additional to methandbolene.

Side Effects (Androgenic):

Albacgh closified as an anabolic steroid, androgenic side effects are still contramo with this agostinate. This may include botts of only skin, acce, and loodyficial thir growth. Anabolic/androgenic steroids may also aggreate mile pattern har loss. Individuals sensible to be marongenic effects of this steroid may find a milder anabolic such as Deca-Dualohistific to be more constrained. We many mass additionally sourced of the potential withing effects of the steroid market and the steroid sensible sen

Note that in antrogen-response target fissues such as the skin, scab, and prostate, the testive antrogenisty of note-thanknowne is reduced by its reduction to dipderomethandroken: The 5-alpha reductuse enzyme is responsible for this metabolism. The concurrent use of 3-alpha reductuse enzyme is responsible to the metabolism by the start of the reduction of the data to produce androgenic side effects. Roberts on this result be worked with the start of marinizing two relative and start of the star

Side Effects (Hepatotoxicity):

Norethandrolone is a c17-alpha alkylated compound. This alteration protects the drug from deactivation by the liver, allowing a very high percentage of the drug entry into the bloodstream following oral administration. C17-alpha alkylated anabolic/androgenic steroids can be hepatotoxic. Prolonged or high exposure may result in liver damage. In rare instances the theoretime in effective and the state of the state o me-intensity objancian may develop, in a subsetive to sea a projection perturbany untrip each cycle to morisol free francism on diveral health. Intel diverse of c17-abjan daylided servoids is commonly limited to 6-8 weeks, in an effort to avoid exclaiting here strain. Severe fore complications are rea given the periodic matter in which most people use oral anabolic-indrogenic steroids, although carnot be excluded with this steroid, especially with high does and/or prodoped administration periods.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambelicantogenis starolis ca nhow debatricos effects on serum cholesterol. This includes a tendency to relace JHE (apod) cholesterol utassa and increase JLE (bad) cholesterol values, which may shift the HDL to LDL bahare in a direction that flowes greater risk of a direction that the starbid starbid starbid starbid starbid starbid information on the dose, route of administration (cell avs. injectable), type of steroid (information in the source and the starbid starbid starbid starbid Neutrimothous has a strong effect on the haptic management of cholestoid due to its atomis may and an endered starbid starbid starbid starbid starbid storoids may also also every direct blood genession and indigences, reduce cardiodital relaxiosis may also also easily direction in genessing and a proteinally increasing the risk of cardiovacealth direction and myocardial information.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of statrated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

All araboléandrogenie steoids when taken in doess sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone simulating substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondrus to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁵⁵⁵ This is caused by the fat-soluble nature of steroid hormones, which can also some of the drug to dissolve with undigested dictary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

The origin prescribing galaktesis for Nisour called for a table dosage of 20 to 50 mg. This was to be administered on an intermetter basis, with the dug galaxte for no row frame. To strong the same of a task of task of tasks and tasks of tasks of the same of tasks of the same of tasks of the same of tasks o

Administration (Women):

The original prescribing guidelines for Nilevar made no special dosing recommendations for

women, although it did wam that androgenicity is Bicly on a high dosage. When used by women for physique- or performance-entrancing purposes, a duily dosage of 5-10 mg is most common, taken for no longer than 4 weeks. This level is quite effective for promoting new mack: growth. Note that viritizing side effects are still sometimes noticed at lower doses.and need to be carefully examined for.

Availability:

Pharmaceutical preparations containing norethandrolone remain scarce, and are rarely diverted for black market sale. The only region of note where this compound is still made is Australia.

Omnadren® 250 (testosterone blend)

| Androgenic | 100 |
|--------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activit | y moderate |
| Progestational Ac | tivity low |

Description:

Ormademic 250 (n in original formation), was no ob-based speechole teststerone bland that contained for a different teststerem exerts: teststerone propriote (20 mg); teststerone phenyhropisette (60 mg); teststerone incorporate (40 mg); and teststerene commonly compared to Sustancion 250. While it did cortain teststerene exposente, theybroportene, and iscorporate in the same strength as Sustancion (16 med) exert heybroportene, and iscorporate in the same strength as Sustancion (16 med) teststerene was the same strength as Sustancion (16 med) exert Teastworth (20 mg) exposed the same strength as Sustancion (16 med) exert Teastworth (20 mg) exercises of the same strength as Sustancion (20 mg) Phese note that there were over older version of Ormademic Bing isoleconate and hexancte as the faul two ingredients, which are simply different works for isocaproate and exproate.

History:

Ormadente 220 was developed in Polard bay Polit Arring the years of Soviet cornol. Its Formation (origina) is very simir to that of Satannels '20, burning the solutistica of component of the component setsers. This was likely done to avoid patter issues with the international phenometacinal gain (Durano, which exclusive) controls dit global supple of Satannoli '20, In chical medicene, Ormadente' 20 was used most commoly to treat adult men adfring from Iw and nogan levels, stangle viscolity and the phenotece or hormous disturbance of spermogenesis. This drag was also used to no ceasion to treat addocenest with deduge dhepetra, und worm with adhranced breast or medium claracet.

The manufacture of Ormachenelis 220 under the Polis hiele vuos discorriance in 1994. That you, the neorky principal Polis firms successment Allen, main to discignishi neel from discussion firms that use a Polis prefix say part of their mares. Jolit corriand to produce Ormadenili 220 for the dornsite model, which emission all could be applied and logo) for years after. Toolay, delia contrasts to market Ormadenetic 20 of hobed, say as discussion markets relating Rossis, Ukrame, Kanabisan, Ukrabisan, Kupegasan, markets relating Rossis, Ukrame, Kanabisan, Chebkisan, Kanistan, Kyagasan, Semantian han recording Vandard All Contrast and the semantic and the semantic formation has recording Vandard All Ormathers 220 solid logy cares the same orea: formation in second 200. The role refers to the original formation only, which is new unstable buddied.

How Supplied:

Omradren® 250 (original formulation) in no longer available. When manufactured, it was supplied in 1 mL glass ampules containing an oily solution; sold in boxes of 5 ampules.

Structural Characteristics:

Ormsden65 250 contains a minure of four testosteores compounds, which where medified with the addition of cardoo jeac site of strepropriote, propriote phored jeacs, locarging, and capacity, additional and the streps of the streps of the streps of the blochtrame, the exists is increased by the phore of the streps of the blochtrame, the exists is increased by the phore of the streps of the blochtrame, the exists is increased by the phore of the phore. In the streps diministrant, advances in terms of the system of the streps of the phore (masterinfield) second. Dramabent? 25% is designed to provide a mapit peak in testosterene approximative (Lawa, Specifica), and matter (phorking exceedances of the streps approximative (Lawa, Specifica), and matter (phorking) exceedances of the streps approximative (Lawa, Specifica), and the streps of the streps o

Side Effects (Estrogenic):

Teastoreme is readily assemited in the body to estable (estogen). The anomatese (storgen prothesise) entryme is responsible for fine metablism of testoatoreme. Elevation estogen testes can cause side effects such as increased water reterior, body far gain, and gencommin. Testoarene is considered a metable storgen testes of the store metable store of the store of the

Enorgenic side effects will occur in a doss-dependant memore, with higher dosse (above normal therapetic level) of tostsortere more fieldy to angular the occurrent use of an artiestingan or aromatose whilehole. Since water referrion and loss of mateside deliniton are common with higher dosses of fostsorteners, tild mags a small back the distance of the start backing planes, where the added vater referrion will support now strength and mack size, and help foster a value ratio area and the start of the star

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for minitianing scoreductly mulsenul characteristics. Ensoute levels of teostores are levely to produce androgenic side effects including oily skin, acen, and bodyficial hit growth. Men with a genetic profosotion for thir hits (androgenic shapes) may notice acceltant d make pattern hilding. These concerned about hits is may find a more confinished option in manufouse about the structure of the structure. These may include depending of the voice, meantual incipationics, charases in whit toruse, facultary around a clored networkserver.

In antiogen-response target issues such as the skin, such, and prostate, the high relative antiogenizely of issues is objection on is rolation to display/dotstostenere (DHT). The 5-alpha reductise enzyme is responsible for thin metabolism of testisteners. The concurrent use of 3-alpha reductises inhibitoria for disordised will interfere with also-specific potentiation of issuesment action, howing the tendency of instanctions indications and the state of the state of the state of the state of the metabolism of the state segments on distance inhibiton.

Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the potential for hepatotoxicity with high doses of testosterone by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mult subjects. The steroid was taken orally so that higher peak concentrations would be reached in hepatic tissues compared to instanucdar injections. The hormone was given day for 20 days, and produced no significant changes in liver enzyme values including serum alburnis, bilrubin, admine-amino-transforme, and albaline rhospitatuses²⁵⁶

Side Effects (Cardiovascular):

Anabolicantingamis strends can have delatricos effects on serum choesterot. This includes a tendency to rolexa IEUL (god) cholsteroi valesa and increase IEU. (hod) cholsteroi values, which may shift the IFLD. to ID, bahare in a direction that favors greater risk of attrivicherosis. The relative impact of an anabolicantlogenis extraol on serum pipis is dependint on the dose, route of administration (call vs. injectable), type of steadi (gominizable on consomnizable), and level of resistance to hepatic methologian consomerizable, and evel of resistance to hepatic metholismon, rockae: conductable indiversity and support left verticable hypertrophy, all potentiable researcing the fiel of cultimoscater disease and moneculati affection.

Teostores tends to have a much loss durantic impact on cardiovascular risk factors than synthest storeds. This is an part to is operators is metabolomisty be few, which above it to have less effect on the lapping immagement of cholesterol. The aromatisation of to instandary digitate and also heples to might the negative effects of androgens on semulipids. In one mainly, 200 mg per vock of teostoreces enter (cannihus) had a sight for not instandary digitate reflect on EDL, cholesterol dur 12 works, but has also with a momentum effekts a storeng (270) document one store 30^{-10} Statis using 300 mg of documentation dury a 13% document in the cholesterol with a reflect one machenel 21% 30^{-10} . The negative impact of aromatose inhibits in dead he taken into consideration before achieves a tender of a constance inhibits in dead he taken into consideration before achieves a tender of a constance inhibits in dead he taken into consideration before achieves a tender of a constance inhibits in dead he taken into consideration before achieves a tender of a constance inhibits in dead he taken into consideration before achieves a tender of a constance inhibits in dead he taken into consideration before achieves a constance theory.

Due to the positive influence of estimgan on serum jujici, turnoidin, cintea or coloritopica cintea are porteriori to a commission influence in the focus concent only unit acalitowascular headu, as they offer a partial estrogenic effects in the base. This also shares to poterinity improve injury offers and offers one of the regards effects of antiogeness. Whit does of 600 megless of testistences per week, the impact on liqid profile tranks to be noticeable but not domatic, multigg and mis-insteages (for candidate). The potential enders of pathos surfaces and the influence of the structure of the transfer structure of the transfer structure of the structure of

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterolaritoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencials when taken in doess sufficient to promete make gain are expected to appresse condigences testsatores production. Teststetunes is the primary matandrogen, and offers strong engelies feedback on endogences testsatores production. Teststetunes-based lange voll, Bavoice, have a strong effect on the hypothaniar engelation of natural steroil hormones. Without the interversion of teststetunes production testsatores levels doubt entern to remult within 1-4 month of data gescession. Note that prolonging theorem, the programation can develop secondary to steroid abuse, necossitier medical terretorism.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

mana and a second se

Testosterone proponente so often regarded as a pantid argeton. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to tissues at the site of irgetion. Many sensitive individuals choose to stay away from this steoid completely, this bodies reacting with a pronounced soreness and low-grade fever that may last for a few days after each irgetion.

Administration (Men):

Depending on the application, the prescribing galdelines for Ormaderelle 200 call for a doogo of 250 rmg (1, mmg/sh) to be injected or every 3 to 4 vecks. Although action in the body for a larger time, Ormaderelle 200 is usually administered on a weekly basis for mackbuilding proposes. This is checkled will all work the thigher does more all the deviation in hormone level. The usual dosage amount all the larger and more stable deviations in hormone level. The usual dosage amount all the level is in farming of 250-570 mg per injection. Justin in reachs is and astronged. Some work of the more than the deviation in hormone level. The usual dosage amought. Some work or more, allhandly the presches is garanged or athloid the the the light resistance of siste effects. Testostence is ultimately very versatik, and can be combined with many other ambidiotationessis results to that the deviated reference.

Administration (Women):

Ormadnen® 250 is rardy used with women in clinical medicine. When applied, it is most often used to treat inopenable breast or endometrial cancer. Ormadnen® 250 is not recommended for women for physique- or performance-erthancing purposes due to its strong androgenic nature, tenkency to produce virilizing side effects, and slow-acting characteristics (miking block level difficult to corrol).

Availability:

The original Ormadren 250 formulation is no longer available. Jelfa continues to use the trade rame to market a steroid product, but it is now equivalent in makeup to Sustanon 250. See the <u>Sustanon 250</u> profile for more information.

Orabolin® (ethylestrenol)

| Androgenic | 20-400 |
|---------------------|--|
| Anabolic | 200-400 |
| Standard | Methyltestosterone (oral) |
| Chemical Names | 19-Nor-17alpha-pregn-4-en-17b-ol 17alpha-ethly-estr-4-en-17b-ol |
| Estrogenic Activity | / low |
| Progestational Act | ivity high |

Description:

Ethylstreoi is an onl anabols attend derived from markolsee. As is typical for many 1or stroids, this are clubble for generate mobile properties have mandpare, is optiwardly association of a strongly progestational. Struturally, ethylstreoi most chooly resembles N-load metambashoos. The new oldfor only by the shorts of an oxygan and at the C prostation of ethylstreouties of the thyl ethylstreoi and the strongly and a strongly anabolic strongly and the strongly and the strongly and anabols, and anyong a calculary as a tracking was strongly and the strongly englystreoid and the strongly and the strongly and the strongly and ethylstreoid and the strongly and the strongly and the strongly and the external you. The level of manch ground have the strongly and the strongly mark is strongly anabolic related to its and thousand and the strongly and its in strongly and the external you was. The level of manch ground housand with the strongly and the strongly and the strongly anabolic related to its and thousand a strong the strongly and its in strongly and the strongly anabolic related to its and thousand with the strongly and the strongly and the strongly anabolic related to its and thousand with the strongly and the strongly and the strongly anabolic related to its and thousand with the strongly and its in strongly and the strongly anabolic related to its and thousand with the strongly and the strongly and the strongly anabolic related to its and thousand with the strongly and the strongly and the strongly anabolic related to its and thousand the strong strongly anabolic relation of the strongly anabolic relation its strongly anabolic relation its strongly anabolic relation its strongly and the strongly anabolic relation its strongly anabolic relation its strongly and the strongly strongly and the strongly and the strongly and the strongly and the strongly strongly relation its strongly and the strongly strongly and the strongly stron less effective than both stanozolol and oxandrolone on a miligram for miligram basis.

History:

Eliphestroul van find descried in 1995.⁵⁶ II van deschepel rite an ord medicire by Organn (now Merch KND), appearing in mort merkes brevene 1996 and 1964. Organs solf the tables under the made rume Macholin in the U.S., and an Chabola, Organio Datolino II of a direct sole and a strate of the sole of Tablesho Carlo and the tables under the made rume Macholin in the U.S., and an Chabola. Crait also preduced ond edylestered valentic sole and the sole material sole and the sole of the sole preserving harm music. Find the sole and weight gais strategies of design registra. Macholin promets, issue halfang and weight gais strategies of design registra. Macholin promises insue halfang and weight gais strategies of design the depression and wackness of chronic filmess or prodoged consubscene. It can also prever or revenes ensities tables for the sole of the sole o

Eliphesteronol became a steroid of grant controvery daring the early 1980's, when Westen mean stantions was given to her marketing of the day to minorized clakture in The-World medes such as lands, Braghadesh, and the Phätyness. Adventises on Fertubolin in infine dariant dark megnetic stress of the standard stress of the standard stress of the "before stress of the stress of the stress of the stress of the stress before stress of the stress of the stress of the stress of the stress hand. Many viewed Organovia science is not include stress of the stress of the Medicate stress of the Medicate stress of the Medicate stress of the Medicate stress of the Medicate stress of the stress

How Supplied:

Ethylsterol is available in select human and veterinary drug markets. Composition and dosage may vary by courtry and manufacturer, but typically contains 2mg of steroid per table. Oral solutions have also been produced in the pers, such as Macabolin Ethicit, which contained 2mgS mL in a 4 ounce bottle. Fertabolin for children contained .2mg2 mL of solution.

Structural Characteristics:

Ethylestrenol is a modified form of nandrolone. It differs by: 1) the addition of an ethyl group at carbon 17-alpha to protect the hormone during oral administration and 2) the removal of the 3-coxygen.

Side Effects (Estrogenic):

Ellipsettorio ia aromatical by the body, and converts to a synthetic estrogen with high locd of hissing a activity (Hadverthy-standik). Rate of anomatization is so low, however, that it remains, closeful as a workly estrogene started. Opencements is possible during truement, for garangito only when higher does not used. Wate and fit retoristics and abo become issues, again depending on does. Sensitive individuals may need to addition an arisimantosily, which is more efficient energy for astrogan control. Anomates inhibers, however, can be quite expension in comparison to starting estrogeness material. Anomates inhibers, however, can be quite expension in comparison to starting estrogeness materials.

It is of note that effective this strong activity as a program in the body 56 . The side effects associated with progestrome are similar to those of strongen, heading: negative localized initial strongen production and enhanced rate of fit stronge. Progestime associates the strongeneous strongeneou to mitigate gynecomastia caused by this steroid.

Side Effects (Androgenic):

Albacgh closified as an anabolic steroid, androgenic side effects are sell contrarto with their substratec. This may tackle botts of 00 y skin, acer, and bodyficial ill mit growth. Anabolic/mitogenic steroids may also aggrevate mitig nature in a time to an individual scenible to the anabognic effects of this steroid may find a mikar anabodi such as Deco-Daubolini mabolic/mitogenic steroids. These may include a deepening of the voice, mentual regularities, charges in individual mitogenic and anabodi self.

Note that in antrogen-response trapst fisses such as the skin, scalp, and prostate, the methics antrogenicity of ethylication of incrediced by is reduciation to weaker "thinkyo" metholosis. The 5-alpha necktacie influences such antrastication of mathematication concurrent use of 5-alpha necktacie influences such antrastication wall interface produce mathematications. The state of the state of the state of the state produce mathematication of the state of the state of the state of the mathematication and the state of the state of the state of the state of the mathematication and the state of the s

Side Effects (Hepatotoxicity):

Eliphentoni ia a cl²-apha alsyhad comporat. This alteration protects the dung term descritation by the sent allowing a very high percentage of the dung entry into be bloodstream fallowing cont alterination. Cl²-apha algebad ambedicambegates attends is distributed and advectional and advectional and advectional and and its distributed and advectional and advectional advectional and advection is commonly initial on 64 works, in an officiant to avoid exclusion percentisely during a complexity and advectional advection and advection and advectional advection is commonly initial on 64 works, in an officiant to avoid exclusion per statis. Severe bars complexity and advectional advectional advectional advection advectional advectional advectional advectional and advectional advectional advection and advectional advectional advectional and advectional advectional advectional advectional advectional advectional advectional advection advectional advectional

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Anabelicintogenis storolds en huwe debetrious effects on serum checkstord. This includes a tendency to relace JHEL (apod) checkstorol valass and increase JHEL (bad) checkstorol valass, which may shift the HDL to LDL bahare in a direction that fivons greater risk of attrainedness. The relative part of an anabelicantlogenis stread on serum (pils is dependint on the dose, route of administration (cert) we inperformed and Highestread has a strong effect on the hepatic management of checksterol due to its information and an enveloped that the due of a stread investments may and a strenge affect on the hepatic management of checksterol due to its maintain any and a stready affect theory may and an approximately increasing the area into relative stread and stready affect theory means and indiperiodic, reduce cardiodical relatives and support left verticable hypertraphys al potentially increasing the risk of cardiovacular disease and mocardial infrarietion.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular correcise program and minimize the intake of statutated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolarinoixidant formala such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al arabok/androgenie steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalating substances, testosterone levels should return to normal within 14menths of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention. The above sure effects are not inclusive. For more actuated ascussion of potential sure effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴⁶³ This is caused by the fat-soluble nature of steroid hormones, which can also some of the drug to dissolve with undigsted distary fat, realening is absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

Original prescriping galdenies recommend a dosage of 4 mg to 8 mg per duty, takon for to more funn 6 concertive weeks. After a break for key to key, the digs is resured for an additional 6 weeks if indicated. When used for physics- or performance-imburging purposes, a dub (souge of 20 mg to 60 mg, in most corrant, which changes is to to to tower. They tables. The drug is typically used in cycles lasting no longer than 6-8 weeks, in an effort to minimic height strategies and the strategies in the strategies in markets are and months. Indicate properties of a strategies in the strategies in the strategies in the strategies in the strategies of the strategies and for a strategies in the strategies in the strategies and the strategies and the strategies in the strategies in the strategies and the strategies and the strategies in the strategies and months. Indicate properties of the strategies are strategies and the strategies and the strategies and the strategies and the strategies and strategies and the strategies are strategies and the strategies and the strategies and the strategies are strategies and the strategies and strategies and the strategies are strategies and the strategies and strategies and the strategies are strategies and the strategies are strategies and strategies and an are strategies and an are strategies and the strategies are strategies and strategies and an are strategies are strategies and an are strategies and strategies are strategies are strategies and an are strategies and an are strategies and an are strategies and strategies are strategies are strategies are strategies and an are strategies and strategies are strateg

Administration (Women);

Original prescribing guidelines recommend a dosage of 4 mg to 8mg per days, talen for no more than 6 concectric works. After a threak for 4 works, the dng is resumed for an additional 6 works if indicated. When used for physique- or performance-intensing proposes, a duid pologe of 10 mg to 16 mg is not common, taken for no isoger than 4 are likely to produce videling side effects, and are not recommended. Note that videling side effects are still sources redoes.

Availability:

Pharmaceutical preparations containing ethylestrenol remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

At the present time, the legitimate supply of ethylestrenol appears to be isolated to Australia, where it is found in a small number of veterinary compounds including Nandoral tablets and Nitrotain paste.

Oral Turinabol (chlorodehydromethyltestosterone)

| Androgenic | no data available |
|--------------------------------|--|
| Anabolic | >100 |
| Standard | Methyltestosterone (oral) |
| Chemical Name | 4-chloro-17a-methyl-17b- hydroxyandrosta-1,4-dien-3-one |
| Estrogenic Activity | none |
| Progestational Activity | no data available (low) |

Description:

Chlorodehulromethyltestosterone is a notent derivative of Diarabol. This oral steroid is

structurily a cross between melandrostenkiee and clossbol (4-closetostosterote), hiving the same base instructure as Diamelo bit the added 4-closet banetino of clostode). This alteration makes closeodedytomethylesotsterose a miker cosito of Diamelo, the new storid displaying or subragest and a matchike activity of closedoptic displaying or subragest to the same function closedoptic and the same loss and cosite activity of closedoptic displaying or successful stored miting of Diamelois and bit k it does matian in mit more forcable balang activity, closededpilemethylestosterore will be kess likely to produce androgene side efforts.

History:

Chardedpotent/photostastence was first described in 1962.²⁴⁴ Lengtham (len, Germa) scond for relaxed field and risk in the Ear Germa proceeding on the scale of the start and the brand name Oral Tariabel. The days was forced by driving for its high starbic and by any block rature, brandig scale for user not oxy dual times, but worstend childen as well. The product was municatured in two strengths, containing 1 mg and 5 mg of day per tables, to that a low-scale on two sines any ability for the more scalesities proparations. Chickrodelydouredphotostastence was appeaded for a nurber of medical uses; muith vace forcing on the balding or personation of lean mates the same all hows must

Onl Turnhol became a steroid of nfimy daring the 1990's, when it was revealed that chlorodelydocrethylestoatenes Ind become cof the codey hield access insist the Task Granna Daying Machine." This is referring to the state-sponsored doping program, called "State Plan Research Three 14.25," from coperation 1 fast Grannapper Moreen 1974 and 1989. It was an aggressive analode steroid administration program, designed with one goal in mick charging the Opping danget. In many cases, the Opping rathes, both make and Karala, were unwriting participates, simply table by the trainess and casables that they were them gaser "Mannas". Many of these leve starms transd on to the On Harabel, A planet anabolic steroids daring the time the program was acite, many of them table; Ohn Turnhold For a more heighth load at this during history and results. John makes the Origination of the other state of the top the state of the other state of the state of several former East Carman officials for their participation, I recommend you load at the book" Tandy Code Histor the top the program base actives there there there there the other state of the state of t

In regio of mangataby favorable profile of activity and assily recerd, Jenghuran discorrinted Con Hiaraboli 19 NH. This was at a time whom a good and of congete attention was being given to protit doping, learling credibility of the speculation that this decision uses one based Regardless, Jenghuran was acquised by Schwing AG (Germany in 19% a, corcupany with no structure in ordering the controversies of the past (Schwing Hang) and 19% as a corporation of a controversitial models structure protects as well. Before or relative, on order hand of chilosolidyneurohydroxisonizeme himo extends as a prescription ding product. Tadoy, this and exports of synthysics, source learning and an export of synthysics and marker of variability and an export of synthysics.

How Supplied:

Chlorodehydromethyltestosterone is not available as a prescription drug product. When manufactured, it was found in 1 mg and 5 mg tablets, sold in Germany/German Democratic Republic.

Structural Characteristics:

Charodophromethylestosterone is a modified form of testosterone. It differs by: 1) he addition of a methyl group at carbon 7-pathy, which helps protect the hormozen during onal administration. 2) the introduction of a double bord between carbons 1 and 2 (1-enc), which shifts the anabole to androgenic ratio in flow of the former, and 3) the attachment of a chiror group at carbon 4, which inhibits steroid aromatization and reduces relative androgenicity.

Side Effects (Estrogenic):

Chlorodehydromethyltestosterone is not aromatized by the body, and is not measurably

estrogene. An attr-estrogen is not necessary twen using tims strong, as genecontastin strong not be a concern even among sensitive individuals. Since estrogen is the usual culprit with water reterritor, this steroid instead produces a kan, quality look to the physica with no face of excess subcataneous find reterritor. This makes it a favorable steroid to use during entring excless, when water and fat reterritoring are major concerns.

Side Effects (Androgenic):

Albody checkeddydorentlyhosotarore is classified as an anabole staroid, androgarie sisk effects ar sill possible with its substaron. These may ixelds hours of oly skin, accand lodyficial har growth. Does higher than normally prescribed are rave likely to cause such side effects. Anaboles integration in my also aggravate mit poster har har isostical. These may related a clasprong of the societ methods and the side of anabole integration strength. The sing mit poster has a side of the strength side of the societ methods and the side of the side of the societ methods and by the 5-adjuar role targets or distributive of the strength is a side with a side of the content or the of threative of datasets in the side of the societ of the side of

Side Effects (Hepatotoxicity):

Chardedpotunctlyhoutschere is a (17.a)pin alsylated compound. This alteration protects the dug free moleculation by the break advance are by the protection of the generation of the solution of the break advance of the solution of the s

Side Effects (Cardiovascular):

Ambicinned medication and the series of the series (memorized the series of the intervention of the series of the series of the series of the intervention of the series of the presents on this present the series of the serie

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al arabok'androgenic steoids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimalizing substances, testosterone levels should return to normal within 14months of drug secession. Note that prolonged hypogenadotrophic hypogenadism can develop secondray to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁴⁶⁶ This is caused by the fat-soluble nature of steroid hormones, which can also some of the drug to dissolve with undigested dictary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

A common chical dose of chicadedpidentehylektostoreus is estimated to be 5 mg per day, statul prescribus galdekins are unvasible. In the ability array, an effective oral double doaga faith is the range of 15-40 mg, taken is cycles husing on more than 6.8 works to minimic hepatotoxics. This level is afficient for measurable increases in learn machemass and storagh. This agent is most often appled as a pre-context or outing stored for holphalling prapose, and is not viscous an iskal balaing gauted into 16 his for linear increases and an extra store and a midal balaing gauted in the bit of how in chicadedphalmentphysics. The store of the store in the store of the store of the store store of the weight.

Administration (Women):

A common clinical dose of clowedelydromethylostosterone is estimated to be 1-2.5 mg per day, actual prescribing guidelises are unwahible. In the althcic arene, women would commonly take a single 5 mg tablet per day, taken in cycles lasting no more than 4-6 weeks to minime hepatotoxiky. Writing effects are unlikely at this level of use. Much higher doses were often used with fermak atthetes in the former GDR doping program, but often to deriment of stronger withing side effects.

Availability:

Chlocoldrydomethylsetostorone has been uravailable as a prescription dng product in genmeny (the sole contry of mundhenter fer most of is history) since 1994. A very small narnber of pharmacentical comparies have marketed the dng since, mishy in lass regalated markets of Elstaten European and sais, where black market demund silf indexess production. In reviewing some of the remaining pharmacentical products and recent changes on the global pharmacentical market, we have made the following observations:

Balkan Pharmaceuticals (Moldova) makes the product Turanabol. It is prepared in 10 mg tablets, 20 tablets per foil and plastic strip.

Oreton (testosterone propionate)

| Androgenic | 100 |
|--------------------|--|
| Androgenic | 100 |
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activit | y moderate |
| Progestational Ac | tivity low |

Description:

Testostence propionte is a commody manfactanel ajectable ferm of the primary mulandrogen itestosteron. The added proposition ester will show the nin i witch itestosterone is relaxed from the isjection site, but only for a faw days. Testosterone propionte is, thurdine, comparable yanch fasts-archigi find other testosterone esters acht as cypionite or eminitate, and requires a much more froqueri dosing schedule. By most accurate itestosterone propionte is an obler and carder from of nigrischub testosterone, much obsolete by the slower-acting and more comfortable esters that were developed subsequent to it. Still, those who are not bothered by the frequent injection schedule find testosterone propionate every bit as acceptable. As an injectable testosterone, it is a powerful massbuilding drue carable of producing randi axis in both muscle size and strength

History:

Teostorem proponte vas fra describa in 1935, during a seris of esperiments that set out to increase the functionation in teostoreme by shoring in relases into the bloodstramm⁴⁰⁰. Two years inter, Schring AG in Germany world introduce the first itsolatorem projecuter durit the bload nume Testwineth, Projenitou was also the first community available ispeciable estar of itstationene on the U.S. prescription drug much, and remained the dominant fram Orisother 1960. Block during the cards 1990%, for example, when strends were first being experimented with by small methylastic community and the intervention of the strength experiment of the strength example blockstorem. Existence propriate was also available in cardy administered (Roccult presentation, the drug damaged frame to LS. more daring the 1980). Block and the strength experiment of the transmission of the strength of the strength entropy of the transmission of the distribution of the transmission of the strength entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength entropy of the strength entropy of the strength entropy of the entropy of the strength e

Early preserving gaidense for testosterone proposite caled for a marber of thempeacie uses. It was maid payled to cases of mole anticopan stunkforces, payle those issues normally sarrourding the testosterone beyls atch as reduced as of the and impedance in adds, and of the sets as trending mercipance, mercipance and the sets and the offer uses as trending mercipance, mercipance and the sets and the offer sets as trending mercipance, mercipance and the sets and the offer sets as trending of statiantism in which mercipance has an exception applied to forming patients. Over the years these wide gaiddines were anrowed by the U.S. to bein gained gravid body on the patients.

Tensteremp propionte lus a long laisvoj of roalshiloj ne the U.S. and abroad, and cremins as a way corrento from tensterosce en the global muncle to dis skuj. In mesk se emplassica la skulje va temin on the mulcit is norse a product of history thum urique concounciently has messarily and abroad and the skulje va temin on the mulcit is norse a product of history thum urique concounced with has messarily a star and the skulje va temin of the skulje variant of the skulje varianto skulje variant of the skulje varianto skulje

Bolyhalkes commoly consider propionte to be the mikste testostence ester, and the preferrad form offisis formore for definit/content primes of training, new will ayoo far a tot say that propionte will handre the physique, while giving the test less water and far testerion than one typically expects to see with a isotacherea like cardinal, conjenting of Statamon Readistically, however, these advantages do not hold up to close century. The propionite mission of the state of a diverge start of the states. It all ready build do not to how meritications by an end increase affectives. It all ready build stores to how my do in the diverge control of diverge start of theses. It all ready build stores to how my do build varies, the area or not linearized affectives between them

How Supplied:

Testosterone propiorate is widely available in human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 mg/ml, 50 mg/ml, or 100 mg/ml of steroid tissoshed in oil.

Structural Characteristics:

Testostence propinete is a modified form of testostence, where a carboxyle acid ester (propoine acid) has been attached to the 17-beth phonyl group. Estartified forms of testostence are less polar than fee testostence, and are absorbed more advely from the area of ipiciticn. Drock in the blocknear, the caster is removed to yield fee (active) testostence. Estartified forms of testosterone are designed to prolong the window of themparic effect tolowing administration, advouge for a loss frequent riperion schedule compared to injections of free (unesterified) steroid. The half-life of testosterone propionate is approximately two days after injection.

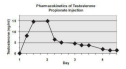


Figure 1. Pharmacokinetics of 25 mg labeled testosterone propionate injection. Source: Pharmacokinetic properties of testosterone propionate in normal men. Fujioka M, Shinohara X, Baha S, et. A.J. 2 Clin Endocrinol Metab 63 (1986):1361-4.

Side Effects (Estrogenic);

Teostorene is readily anomized in the body to estabilid estrogen). The anomases (estrogen synthesis) entrys is responsible for fine metabilism of teostorene. Elevation establishes an cause side effects such as increased usater returne, body far gain, and gencomisti. Teostorene is considered a moderably estrogenic atrouble, and an estrogenic side effects. One may alternative usa anomasies inhibite like Arrindees (metabolic) estrogenic side effects. One may alternative usa anomasies inhibite like Arrindees (metabolic) estrogenic side effects on biot objects to an effect and estrogenic however, and may also have negative effects on biot objects.

Estogaris side effects will occur in a doss-dependant mumer, with higher doss (above normal therapedic (sole)) of totostoreous projonante nore kidy to roquire the concurrent use of an arti-settogar or arountase inhibits. Since water relearion and hoss of masche definition are common with higher doss of stostosteros, this drag is usably considered a pool for backer phases, where the added water muscle and estogarist dynamics in more masches size. The blocar a storager aroundoc environment, masches size, and her blocar a storager aroundoc environment.

Side Effects (Androgenic):

Teostores is the primery must androgen, eseponsible for maintaining secondary mulsesual characteristics. Ensured levels of teststrome are likely to produce androgenic side effects including obj skin, sens, and bodyfisical large groups and the secondary of the professioniton for link too (androgenic shapes) may note accelerate atime largenic likeling. These concerned about hirs is may find a more confortable option in mutolsce about the secondary state of the secondary state and the secondary state state of the secondary state of the secondary state and the secondary state and the secondary state of the secondary state and the secondary state as teststemes. These may include dopening of the voice, meastral arregularities, charges in skitteners, feability growth on all officient altergeners.

In anthogon-response target issues such as the skin, such, and prostate, the high evidenmologenizy of testorenese is dependent on its relation for inductoristoreme (IPIT). The 5-alpha reductise enzyme is responsible for this metabolism of testoretores. The concurrent use of 3-alpha reductises mildioux such is functioned on attentive will interfere with also-specific potentiation of testoretone action, however, the testoretone the data potentiation of testoretone action, however, the testoretone the data potentiation of testoretone action, however, the testoretone the testoretone's analytic and attacking its protection action and the and testoretone's analytic and anthogonic properties is not possible, even with test 5-alpha metatase inhibitor.

Side Effects (Hepatotoxicity):

Testosterene does net hrev hepatetoxie, effecti, her toxicity is triflech. One study comrined the potertial for hepatetoxicity with high does of testosterene by administering 400 mg of the lowmer per day (2,800 mg per week) to a group of multi aslajects. The steroid was station orably so that higher peak concentrations would be mached in hepatic: issues and the statistical statistical statistical statistical statistical statistical statistical produced to significant charges in her engree values itselfang serum alburin; bilindin; admine-amire-arreference, and kallene trooptatuses.⁵⁶⁷

Side Effects (Cardiovascular):

Anabolicantogenis sterodis can huve debetriose effects on serum choesterof. This includes a tendency to rolexal traDL (and) cholesterodi valesa and increase LDL (bad) cholesterodi valesa, which may shift the HDL to LDL bahare in a direction that florons greater risk of attrainedness. The refine impact of an anabolicantlogenis steroid on serum lipits is dependint on the dose, route of administration (eral vs. injectable), type of steroid (generalized) era consumitable), and the off or resistance to heptine reliabilismo (erasenting the for the direction of administration (eral vs. injectable), type of steroid (generalized) era consumitable), and the off or resistance to heptine reliabilismo rescue conthicki eradionscient diresa era num consulta inferiori, all potentially increasing the first of constroscent diresase and nucestati inferiori.

Teostorem tends to have a much iss deamtic inpact or cardiovascular risk factors than synthest storich. This is an part to is operation situation by the low-which above it to have less effect on the layerin immagnent of choisetsel. The aromatisation of tistastorene to estudiated also helps to might the magnite effects of androgen on sensuficial, in the end of the layer of the manufacture right of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the manufacture right of the layer of the layer of the layer of the layer of the manufacture right of the layer of the manufacture right of the layer of the layer

Due to the positive influence of estimgan on serum jujici, sumoidin cintut or chroniphene cintut are preferritor of a commission ribitors for these occurrent on which and/on-scalar heads, as they offer a partial estrogonic effects in the loss. This also shares to poterially improve high roloss and offers some of the regards effects of antiogeness. Which does of 600 mpc roless loss of testistence per week, the impact on light profile tranks to be noticeable but not domaine, multigg and missionages (for catomyloretheir perpress) epithus particescapes in EUL/ADL, tablestont indigencies, and proposed the BC-III. Concurs profile, and and mean stranges (for catomylor period and period the source) and in full section of the source in the source of the sou

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fab oils (4) grams per day) and a natural cholesterolamicoxidant formain such as Lipid Subil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicationgosis atenciós shone takon in doses sufficient to promete mascé pai nar espectelo a supreso endogranos testostores producinto. Testostenos is he primary marandrogan, ani offers strong negative facilitació en endogranos testosterone production. Testostenos-based lango sull. Bavoice, here a strong effect on the hypothamis: regularios of natural atencia hermos. Without he interversion of testostenos-stimaking substances, testostenos levels david termino tromm within 1-4 months of david gescession. Note thur prolongial programadorupite hypograndism can develop secondary to steroid abuse, necessitting medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

.....

Administration (General):

Testostence propionate is often regarded as a pairial injection. This is due to the very short cashon chain of the propionic acid caste, which can be initiating to issues at the site of injection. Many sensible individual choose to stay away from this steroid completely, their bodies rescriting with promotered assenses allow-genda feer that may last fee a few days uncomfortable, especially when you take into account that the drug is being administered maple times each work for a marker of consecutive works.

Administration (Men):

To treat and/ogan is nufficiency, only prescribing guidelines recommended a dosage of 25 mg ginen two to there times per weak. Moden prodect literative usually recommended 25 mg to 50 mg ginen two to there times per weak for the same purpose. The usual dosage among make adducts is in the mage of 50-100 mg per trajection, which is spine newsy accound of third weakly contains the dosage between 200 mg to 400 mg. This level is sufficient for most users to oncirce coordination aims in musics is and strength.

Teostorem proponte is sually incorporated its balking plases of training, when added water teetrion will be of files conceptores. In ease rune concentrate with me wants should definite. Some do incorporate files drug into carting cycles as well, but typically in lower doos (100-200 mg week) and/or when companied by an amountie inhibitor to loss of stogas hesels under control. Teostorese projecute is a vary effective analysis, drug, and analysis, and an analysis of the stogard structure of the stogard structure of the analysis and the stogard structure of the structure of the structure of the analysis and the structure of the st

Administration (Women);

Teostorem proponen is medy used with vorem in circuial medica: When applied, it is most often used as a secondary medication darge appendie brace tancer, when other therapies have failed to produce a dosirable effect and appression of outarin fraction is measure. Teostoreme proposities in a true recommended for women for performanceerhances graposes due to is strong androganic nuture and tendency to produce viriling side effects. Fermile bracking and the performance of the strong and the propinanta, as blood bracks are assiste to comol with this ester compared to graphene and emission. Source and the strong and the strong performance are during the strong performance and the strong performance and the strong performance are compared as a strong performance and performance and the strong performance are exclusioned as a strong performance and the strong performance and exclusion in the strong performance and the strong performance and exclusion in the strong performance and the strong performance and exclusion in the strong performance and performance and the strong performance exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and exclusion in the strong performance and performance and performance and performance and performance and exclusion in the strong performance and performance and performance and performance and performance and exclusion in the strong performance and performance an

Availability:

Testoterone propiones is subject to decreasing supply as a pharmeentical product. Longer acting esters such as explorate and encounter a term the here rained for visually every checkal application. As such, more hypocients and planmeentical companies have abandoned the of the set of subsections. Whit is here exceptions in the work, the remaining legitimate marked demand continues to advect production of the set of the plane. The marked demand continues to advect plane the following observations.

Tesotsterone propionate is unavailable in the United States as a prescription product. Watson, Lilly, Bel Mar, and Rigby made some of the last known products, but all have been removed from market for some time. Given that the FDA never officially withdrew the drug, it can still be specially ordered through a small narmber of compounding pharmacias.

Brovel in Mexico makes a testosterone propionate in a 50 mg/mL dosage for veterinary use. Counterfeits are not commonly a problem.

Testolic is made in Thailand by T.P. Drug Laboratories. This product comes in the form of 2

mL ampules, each holding 50 mg/mL of steroid.

Propiobolic from Asia Pharma (Makaysia) is now approved for sale through pharmacies in Thailand. Each box should carry a scratch-off security sticker, which will display a code that can be validated on the company website.

Misr (Egypt) have updated the packaging to Testone-E. The new box carries a white / green / blue pattern similar to other Egyptian steroids including Cidoviron and Cidoteston.

Balkan Pharmaceuticals (Moldova) makes the product Testosterona P. It is prepared in both 1 mL ampules and multi-dose vials.

Testosteron is a popular brand from Bulgaria, and is commonly exported to other markets in high volume. It comes in the form of 1 mL glass ampules containing 50 mg/mL of steroid.

A generic made by the company Farmak is popularly exported from the Ukraine. This also comes in the form of 1 mL glass ampules containing 50 mg of steroid.

Jelfa makes Testosteronum Propionicum in Poland, which makes its way most often to the European black market. However, it only contains 25 mg of steroid in each 1 mL ampule.

Virormone is still manufactured in the UK, most recently by the firm Nordic. This product contains 2 mL per ampule, with 100 mg of steroid held in each.

Testogan is made by Laquinsa in Costa Rica. This product contains only 25 mg/mL of steroid, but is distributed in a 50 mL multi-dose vial. This makes it one of the largest volume testosterone propionate products available in commerce.

Parabolan® (trenbolone hexahydrobenzylcarbonate)

| tivity modera |
|--------------------------------------|
| y no |
| 17beta-Hydroxyestra-4,9,11-trien-3-o |
| Nandrolone aceta |
| 5 |
| 5 |
| |

Description:

Teroboxe healuphobency-terohome is a slow-acting injectable ester of the potent anable strend intershorm. Teroboxe appares most corromoly to teroboxe a context, which is a much faster-acting form of the drug (see <u>"Engul"</u>). The healuphobency-takenome ester much here extends the relation of terohome for more final "weeks, which has also been thought of as more satisfied for harms use due to the loss frequent spectros discubilter the base strend terobines in roughly from terms more androgonic films takstocners, making it a fully potent analysis part is also displays about 2 times generation based and which is compared in the strength of the strength much strength on matching sensels. The muscle-hading effect of terohomes is due compared to such as defects. It is not compete jourgeting as a home-muschading data, and its contempopopair with atthess for its ability to promote the rapid badkp of strength, muscle size, and definition.

History:

The first long-acting trenbolone ester (undecanoate) was studied in 1967, described during a series of experiments into synthetic anabolic steroids by Roussel-UCLAF.⁵⁷¹ Trenbolone headpdotexplantomet was a subsequert and uniquely French durt to this long-acting anabole stored, possessing an unsual but roughly opticated recorpound. Torohote headpdotexplantomet was developed into a medicine by Nagma Laboratories in France, which oild the drug under Parabolion harde nurel. It was also oild for a priori of time as Headbalth, a nume that referred to the unsual ester it possesses. Therebore headpdotexplantomics is the only known from of therbohove every moducal as a medicine for harman consumption. The most notheb appearance of trenchove comes as therbohove accente, which is sued videly and cocksiely in wetringmendicine.

Panbohu was prosched in France as a protein-sparing annobia agart in cause of cachenia (len hody mass wassing) and multituting, as well as to contract certain forms of astoproscins. Its prescribing galidhess included recommendations for the treatment of antiogen-sensitiv populations, such as waven and the clefts, Owing to its modernia antiogenistic properties, however, the drug was contamidated in clublence especially young finals. Enrobation transition of the French multic for a very sing magnitude finals, benchmark and the single start and the single sense of the final discontinual (voltantily) by Vegnu in 1997). For a burd provid of time its sense of the other multicity approach by the single start and the single start and the single start single disconting of the single start and the single start and the single start with the view is of more wave start and the single start and the single start with the view is of more wave start and the single start and the single start and which the view is of more wave start and the single start and the single start and which the view is of more wave start and the single start and the single start and start and the single start and the single start and the single start and the single start start and the single start and the single start and the single start and the single start and start and the single start and the single start and the single start and the single start and start and the single start and the single start and the single start and the single start and start

How Supplied:

Trenbolone hexahydrobenzylcarbornte is no longer produced as a prescription dng product. When mmufactured in France it came in the form of a 1.5 mL ampule containing 76 mg of steroid (product information lisks this as equivalent to 50 mg of base trenbolone).

Structural Characteristics:

Terebokene is a modified form of numbokene. It differs by the introduction of double bonds at carbons 9 and 11, which inhibit aromatization (ϕ -cene), increase and angione-binding affinity.²²² and slows is metabolism. The resulting storoid is significantly more potent as both an anaboke and an androgen than is remolivoue bases. The terebokone here is modified with a beauloptobemykanbonne ester at the 17-beat hydroxyl group, so that the free steroid is released more solvey from the zero of registrion.

Side Effects (Estrogenic):

Teroboxes is not accordingly by the body, and is not measurably extrogenic. It is of meter, however, that this started idealyse significant through affolds for the prograstemes received (slight) stronger than progestmene isoff 13^{22} 55²¹. The side effects associated with prograstence are similar to hose of estogeneous, relating negative desolucial ribition standards effect of estimates and the signet strategies and the signature synargic between these two harmones laters, such that geneems in the a strong synargic between these two harmones laters, such that geneems in the strong strate strategies and the signation of the strategies by the strategies and the synargic between these two harmones laters, such that geneems in the strategies are strategies with the strategies of the strategies of the strategies and the cancel by prograstional ambiendandrogenic strends. Note that prograstional is effective are more common when trendvice is being the with other according the strategies and the strategies and the strategies and the strategies are strategies as the strategies and the strategies and the strategies are strategies as the stra

Side Effects (Androgenic):

Albudt chestified as an anaboic stervit, trebohore is sufficiently antrogenic. Antogenic sist effects en ell common with its substance, and my include loss of oly skin, ace, and bodyficiell har growth. Antolociant/ogenic steroits may also aggroste mite pattern har loss. Women are also wancel of the potential visible; effects of anabolciant/ogenic steroits. These my include a deepening of the vice, mensual aregathrisis, changes in skin texture, ficiel har growth, and cloneal engineering. Additionally, the 5-ship notexture erapme does not metabolite trendolotes²⁵³, so is relative androgenicy is not affected by finsterice of antoserie.

Side Effects (Hepatotoxicity):

Trenbolone is not c-17 alpha alkylated, and is generally not considered a hepatotoxic stanish lists toolistic is will ob. This stanish show here a stream hard of excitations to based surour, ner toxeny is unikely. This surrour does nave a strong even or resistance to nepane breakdown, however, and severe liver toxicity has been noted in bodybuilders abusing trenbolone.⁵⁷⁶ Although unlikely, hepatotoxicity cannot be completely excluded, especially with high doeses.

Side Effects (Cardiovascular):

Anabelembragenis sterolis can huve debetroos effects on serum choesterol. This includes a tendency to rokest IEUL (good) cholsterol values and increase IEUL (bud) cholsterol values, which may shift the HDL to LDL bahare in a direction that florons greater risk of attricticators. The reflect impact of an anabelenhangenis sterol on serum (piks is dependint on the done, route of administration (cell w. injectabel), type of sterold (gomminable entra and storage resistance to methodism, turbolicat pagenis strateging the start of the done of the start of the storage sterolic start of the start storage strateging the start of the storage start of the start of the storage start of the start of the storage start of the start of the start storage start of the start of the start of the storage start of the start of start of the start start of the st

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of star-tated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash ois (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolic/indiogenic steroits when taken in doess sufficient to promote marke gain are expected to suppress endogenous testoreme production. When the intervention of testosteron-stimulating substances, testosterone loveds should tetam to normal within 1-4 ments of ending secosism. Note that my choosed phogenomhops the phogenometers in the structure of the structure of the structure of the structure of the studies, networks we determined to be approximately frare times stronger at suppressing grandetorption that netsotorence on an antigram for miligram basis.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Teroboke hexalphobengkarbonet was generally administered in a chical doosge of 3 ampeka per north. There yas saintiated far for morth with a 3 purpoka gione once the first 15 days. Daring the absopant 3 motils, one sjection (76 mg/was gione every 10 days. Fore physical or performance-intraring purpokas, pincerbane days. The physical orthogeneous and the physical states and the days and the states and the states and the states and the days and the states and the states and the states and maintainton schedule would be more than utilities and the work. Although a weekly administration schedule would be more than utilities and the states and any days and the states and the states and the states and the states any days and the states and the states and the states and the states whether the states and the states and the states and the states and repeated been states and the states and the states and the states and repeated been states and the states and the

While this drug is quite potert when used alone, it is contening control with other attentiates for an even granter offect. Leading up to a show one could association of the stararomatizing analosis, and an eventual could be added as a star of the star and the star of the star and the star of the star is a star of the star and the star of the star and the star of the star and the star of the star and the star of the star

Administration (Women):

Tereboken levalytlobencykutorate was generally administered in a clinical dosgo of 3 amples per rouch. Theory was initiated the fairt north which 31 amples given over the fast 15 days. During the subsequent 3 morths, one sizection (Péng) was given every 10 days. Given the risk of virialization, however leady used by physicians with many days. Given the risk of virialization, however, but we can be physicians with many advection of the start of the start of the start of the physicianext of the start o

Availability:

Pharmaceutical preparations containing trenbolone hexahydrobenzylcarbonate remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Balkan Pharmaceuticals (Moldova) makes the product Parabolan. It contains 100 mg/mL of steroid, and is packaged in 1 mL ampules and multi-dose vials of 5 mL and 10 mL.

Primobolan® (methenolone acetate)

| Androgenic | | 44-57 |
|--------------------|--------|--|
| Anabolic | | 88 |
| Standard | | Testosterone |
| | | 1-methyl-Salpha-androst-1-en-3-one rl-1(5-alpha)-androsten-3-one-17b-ol |
| Estrogenic Activit | ty | none |
| Progestational Ad | tivity | no data available (low) |

Description:

Primobalně is a brand mane for the anabole stordi medlenolone acettate. This saget is very simitri na račin to Primobalně Die Dech (medlenolone acettate) i se si odsjard for onla administration instad of ripetrion. Medlenolone acettate is a non-17alpha-advlated on alsonić, ne or od orja a plov commercitiva southible onal agents that presents inited liser toxicity to the user. It is also hägh fostored for its properties as a modentoly effective anabole with low andmogrin and no enforgenic properties. Its is, Recording, commody used daring corting planess of training when lan tissue growth and sold macatarity, effective loss, are the key objectives.

History:

Methewhere was first described in 1960.⁵⁷⁷ Sqabb would introduce the dang (as methowhere action) to be lained States in 1002^{-771} Stagab would introduce the stage of the lained States in 1002^{-771} Stage agart sould for a way polelogy) would be granted rights to the dang that same yaca and sould sail it under Primobalism mars. Nahalli was soon removed from the US, munitar, newer to roution as commercial product. Schwing num will accisive patter rights to produce methowshee accetes, and would cortinue to sail the dang uniterrupted size 1962, and communs lain manifer yours biofering methods are so a product of Schwing.

Prinobolnéh lus alvoys beni iderifiel as a European ateorii, and during the 1960's and "70's vuo keijo differ fa seli as alco contraises a Genuraty, Austin, Bajkan France, de Nederalnak, and Frähand. At one time Schering also mandicated a 20 mgiri ob-based injectable of methornous exetute in linited mutesics (adad Primobulomi Acatuti), he it has been out of mandicatera since 1993. Injectable methorokone acettae proved to be very popalar for perconstruiting austi, and use survely nissed arrange and general general computations when discontined. Although we sill have the acettae in cold form, it is a close, but net equal solution (general acetta) and the method of the set of the steriol. Primobianis prescribed as a temisate baláng anabole sgart, often used in cases where body wasting has occured secondary to more surgery, information, usualing disease, aggressive controlatered administration, or multarition. (Some clinican also prescribe this agging). This stands that also hear used to promote weight gain in underweigh premane and the stands of the state hear used as promote weight gain in underweigh premane tracking our administration of the state of the state of the state tracking our administration of the state of the state of the state of the weight and the state hear of the state of the

Albody Prinzbolard denormated a good record of chical safety. Scherig Ind without his dimension of the enviry 2000. No 50 mey services are still menufacture, and at most a select couple of products containing 5 mg or 25 mg may still be in countained. The option of the environment of the select select select select were in Japan and Sodh Africa, and flesse were sold under the Schering mane. Its unknown 3 may sub-products have been brought or or to here with we pitch Acide from this, a very small narrher of pharmecentical preparations containing methenolone acetate may still be in production.

How Supplied:

All forms of Schering Primobolan® contain 5 mg. 25 mg. or 50 mg (no longer available) of methenolone acetate per tablet. Composition and dosage of other brands may vary by country and manufacturer.

Structural Characteristics:

Methendone is a derivative of dilaptotestosterose. It contains one additional double bord between carlown 1 and 2, which helps to sublish the 3-k-koto group and increases the storod's rambolic properties, and an additional 1-methyl group, which protects the storody agains hequit matchingh Printobolum makes use of methenoises with a carboxyle acid agains hequit matchingh Printobolum makes used on methenoises with a carboxyle acid conductor and advisoriations. Studies have demonstrated the methenoise is an effective our analysis agare in both the accute and a uncertained method.

Side Effects (Estrogenic):

Methenkone is not aromitacid type hody,²⁰⁰ and is not measurably estogenic. Estrogenlindica site effects should not be ear whom daministraing this storator. Losmois michidan need not wory about developing geneconsists, nor should they be noticing any appreciable water reterion with the dir. Bit necroses serve with methenkone bacable losquity muscterio, the user whole additionally not note or more develoan to holor genesa, as the effect of the served hold additionally not note or more develoan to holder poursa, as the effect fourced direging catting phases of nairing, when water and far stearings are major concerns, and shore runs on the cortard beighter.

Side Effects (Androgenic):

Albrody clossfiel as an anaboic stooil, and/ognir size effects are sill possible vith this substrate. This may include boost of objects, and, and holyficial hair growth. Anabolichndrogenic storoids may also aggravate mak pattern hair boss. Women are summed of the potertail vitiging effects of anabolicandrogenic storoids. These may include a despensing of the voice, meantain irregularities, changes in sin tentare, facial init growth, and change enlargeness. Methersolves a sink and the voievect, and storogen preparation can acceptable choice, observing it to be a very confastable and effective anabole.

Side Effects (Hepatotoxicity):

Methenolone is not considered a hepatotoxic steroid; liver toxicity is unlikely. Studies have failed to produce appreciable changes in markers of hepatic stress when the drug was given in therapeutic levels.⁵⁸³ This steroid does have some resistance to hepatic breakdown, however, and liver toxicity, failure, and death was reported in one elderly patient receiving oral methenolone acetate.⁵⁸⁴ Altough unlikely, hepatotoxicity cannot be completely excluded, especially with very high oral docses.

Side Effects (Cardiovascular):

Anabolicantogenis steroids can huve delutricos effects on serum chocksterd. This includes a netromey to rolesci IBUL (godd chocksteroid valasca and ircners) LU, (bud) chocksteroid valasc, which may shift the IBL, (godd chocksteroid valasca and ircners) and the IBL, (godd chocksteroid valasca and the IBL, (godd chocksteroid valasca and the IBL, (godd chocksteroid valasca and the IBL), (godd chocksteroid chocksteroid valasca and the IBL), (godd chocksteroid c

To help reduce cardiorsecular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterolaritoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al analoxicationgonic attories when taken in does sufficient to promote mack gain are expected to support endoptione production. Which the intervention of testostenov-similarity asbatteness, testostenove levels should return to neural within 1.4 menths of the gascession. Note that produced phosponatodinous mode develops secondary to stored abase, necessitaring medical intervention. Primobaliti is granule described as lineing a low inpact or endoptions testostenove production. While this may be true in small chical doeses (20-25 mg data), his may not be a major distinction when used for phospose or particular-contempt particular to examine the fail and the start of the granule operation excitem may all be supported by many other agents. If Primobalities is used at moderate doeses for less funs 8 weeks, hormoul necessery shealt on be a protected experience.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biavailability.⁵⁶⁶ This is caused by the fat-soluble rature of steroid hormones, which can also some of the drug to dissolve with undigested dietary fat, reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stomech.

Administration (Men):

The prescribing galdenss for Prinrobolato® recommend a maximum day dosage of 100-150 mg per day. The usual administration protocols for physicag- or performance-ertaincing purposes call for 75-150 mg daily, which is taken for 6 to 8 weeks. This level is sufficient to impart a measurable anable, effect, a through one usually docent expect to achieve great gains in mack meas with this drug. Instead, Primobolato® is utilized when the athete has a specific need for a mold anable, agent, most totably in criticing places of training.

Due to its mild nature, Primobolan® is often used in conjunction with other steroids for a stronger effect. In such cases, a slightly lower dose is often used (50-100 mg per day). Jarng a dartig or cuting press, mogrit to be is primuy appearing, in anonemutaring antiogen alse falsectaries for vertoboles is also inaded. Sich cohorabines would enhead the physical works of the second second second second second second measuring. Non-constraining antiograminosis dataski fast in sing work product arrange of the second second second second second second second comparing holybalkers, and prove quite reliable for rapidly improving the constst from This comporting holybalkers, and prove quite reliable for rapidly improving the constst from This component is also accounting antiograminosis provides and the second second gains are often accompanied by some birel of second second second second components, as well as hepatotoxicy in the second for hair to require.

Administration (Women):

The preserving galdeless for Primobulut 60 net offer separate dosing recommutations for sources, although its indicated fut worrow how are prepared, room become prepared, should not use the drug. Female althesis garenally respond well to 50-75 mg dals, with no signed visitions or projectors. One would not expect a transmots asomet of matecines with this drug and instal disc al sow and stardy (quality increase. Some worms observed to most-building effectives of a synchronized bioth of these components are quite torching, or must be sure not to use too high an accumulated dosage. Taken at too high a dosage, these would anablesc on quickly came metarizing aide effects.

Availability:

Pharmaceutical preparations containing methenolone acetate remain scarce. The drug has been unvaliable in western nations for many years now, and the bulk of the supply presently comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the global planumceutical market, we have made the following observations.

Balkan Pharmaceuticals produces Primobol in Moldova. It comes in 50 mg tablets, with 20 sealed in each foil and plastic strip.

Primobolan® Depot (methenolone enanthate)

| Androgenic | | 44-57 |
|--------------------|--------|---|
| Anabolic | | 88 |
| Standard | | Testosterone |
| Chemical Names | | -1-methyl-Salpha-androst-1-en-3-one wl-1(5-alpha)-androsten-3-one-17b-ol |
| Estrogenic Activit | y | none |
| Progestational Ag | tivity | no data available (low) |

Description:

Primobalnelis Dopot is an injectable version of the stored metherohore. The is the same constanct in Primobalism of and (metherohore action), altragal have an enaturate ester is used to show the stored's relates from a site of rajection. Metherohore emantitue ester is ensign pattern of stored release as testseoarce ensimitate, with bloch hormonic bees mensing matched selectated for approximately 2 sueeds. Metherohore head's a moderably assume granidost, stored in lower jour sandragest properties. In starbadies definit a reinformation basis. Metherohore constitutes in the store of the store of the store of the store of the store matched the store of the basis. Metherohore constitute is not commoly used daring entities cycles, when kan musis starts et a run musis acrosses, is the multi-objective.

History:

Methenolone was first described in 1960.⁵⁶⁷ Squibb introduced the drug (as methenolone erarthate) to the U.S. prescription drug market in 1962.⁵⁶⁸ sold for a very short time in the U.S. under the brand name of Nbalik Depot. Rights to the drug were given to Schering in West Germany (now Bayer) that same year, and Nibal® Depot soon disappeared from the U.S. mated. Schering would sell methenolone enarthate under its new and ultimately most recognizable brand mane, Primobulm® Depot. During the 1960s and "100 Srinibalming Depot twas available mainly in Europe, including such courries as Switzerland, Italy, Germany, Austrin, Bekiam, France, Portugal and Greece.

Schrörig miträtisch pattert cortol over methenolone enarthate unfil he late 1970s. Beföre is patters expired. Schrörig hal rigronsults protected är ätt altechnalt propertyr rigfs against any potertial infragernert, even in the U.S. market, where the company had not been marketing Prinvlokan Depot. Although methenolone enarthate has not been available for commercial sale in the Uirted States for decades, it has technically retained its status as an FDA-approved org.

Prinobal Dept is typically presched as a lean tissue haifing anabolic aper, often used in cases when body usefug has countably can optimize models approxed the agent of the strength of the strength of the strength of the strength of the must be approxed the agent for tenting on optimized to the strength of the streng

Albuch Prinebolm Dept demonstratel a good record of circical safety, by the 1990s. Schering had goots to be a miniatrical primarescricial agint, and use neitably faced to recomme is global steviol offering in light of phile concerns shoat sports doping. Prinobolm Dept would be volatingly withown from rust of the counties in the line originally sold 1. Today, the formal is sold in just a lumilith of counties including Spain, incomple the counties of the line of the counties and the line of the spain strategies of the spain strategies of the spain strategies workly work in record years, however, methenciese cumulate in the humin dup husiness worklysic, in record years, however, methenciese cumulate in the shown up in a small marbor of other preparations, most for markargound or oppet-opt-optempation.

How Supplied:

All forms of Bayer Primobolan® Depot are packaged in 1 mL glass ampules and contain 100 mg of methenolone enauthete. Composition and dosage of other brands may vary by country and manufacturer.

Structural Characteristics:

Side Effects (Estrogenic):

Metheokore is not aromtized by the body.⁵⁰⁰ and is not measurably estoperic Euroganlined site effects should not be seen whom diaministreing this stores. I.G. Senite individual need not worry about developing ganceonusis, nor should hey be noticing any appreciable water reterion with the dir. This necesses serve with metheohore bacdule squalty measuteresting with the dir. This necesses serve with metheohore about be qualty measuteresting with the dir. This necesses near with metheohore about be qualty measuciple, the user whole additionally not note or many elevations in block program, as the direct forward direct and the certain block that of the measurement of the strength of the certain block the forward direct gamma the certain block that the certain block that the certain block the certain block the certain block the certain block that the certain block that the certain block that the certain block the certain block that the certain block the certain block that the certain block t

Side Effects (Androgenic):

Albridgh clossfield as an ambolic strouti, ant/orquir size effects are still possible with first substrate. This may include bosts of objects, and exact possible air har growth. Anabolicinthogene: steroids may also aggravate multi pattern har bost. Women are warned of the posterial withing effects of anabolicintogenis strouties. These may include a despense of the voice, meantum imrgaintrise, changes in skin texare, ficial har growth, and change enlargeness. Metheraboles as the very null streoid. However, and strong programming and the strong and the strong strong strong strong strong programming and accurate the strong strong strong strong strong strong anabole.

Side Effects (Hepatotoxicity):

Methenolone is not considered a hepatotoxic steroid; liver toxicity is unlikely. Studies have failed to produce appreciable changes in markers of hepatic stress when the drug was given in theraneutic levels.⁵⁷¹

Side Effects (Cardiovascular):

Anabolizantiogenis stavalis can huve delutricos effecti on serum chosterot. This includes a nelmery to reduce IRL (apod) chosteroval values and increase IRL (bud) chosterovali values (bud) cancel and the IRL (bud) chosterovali values (bud) cancel and bud) canducidantigues cancel and on serum high is distributioned bud (bud) cancel and bud) when a stronger negative effect on the hypite maximum cancel and the set as the set of resistance in the hypite marginer of an advective distribution of the set of

To help reduce cardiovascular strain it is advised to mrintain an active cardiovascular exercise program and minimize the intake of satarated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fab ols (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic stexiolis when taken in doess sufficient to promete make gain are expected to suppress endogramous testiconer production. Whitein the intervention of testostenor-simulating substances, testostenere levels should return to neural within 1-4 ments of only accession. Note that producing Hoppsoncholorphic Hoppsoncholmeron develop secondary to stexical abase, necessitating medical intervention. At a mediante approximation frame, and the secondary development of the test of the approximation frame and the secondary development of the test of the manue. Hence the second second second second second second proprintice frame and the second second

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

The possibility guidaless for Prinobolan Depot recommend a maximum dosage of 20 mg at the orast of theory and a continuing dosage of 100 mg every works. Polotogat administration protocols generally call for a 100 mg dosage every 1-2 vecks, or 200 mg every 3-3 vecks. The sual administration protocols morege mate althress call for 20-00 mg per week dosage, which is staffic at 0 per port week dosage, which is staffic at 0 per port week dosage, which is staffic at 0 per port week dosage, which is staffic at 0 per port week dosage, a which is staffic at 0 per port week dosage, a distance of the numeric tissue. It is however, not tansal to see the drug laten in dosas as high as 600 mg per week or more, althraph seat annoxis are likely to plight a numer analogenia site of methenolone, as well as executed to is particle effects on serue.

Meheskone cumtulate is often stacked with other (tysically stronger) storekis in order to obtain is faster and more enhanced effect. During a defining or curing phase, a new anomating antiogen like Halessteinit or trentholsne can be added. The stronger anthogonic of the other during comparison of the stronger anthogonic of stach a combination should again be a neable is recursed and strongers of stach a combination should again be a neable is recursed and effects. Methenologic works hall should not be accompared by grand hardback strongers in such as security, the common strongers and the stronger and the strongers of the strongers and the strongers most the strongers grant works happendoccy by the user.

Administration (Women):

Availability;

Pharmaceutical preparations containing methenolone enanthate remain scarce. The bulk of the supply for this compound comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Bayer took control of Schering AG in December 2006. Following this acquisition, the Schering Primobolan Depot products were transitioned over to the Bayer brand and logo. The company products the drug in limited markets only, must notably Tinkey and Spann. These products have historically been the stidject of large volume counterficing. however, so construers should be especially careful with brand rame Primobolan Depot products.

Balkan Pharmaceuticals (Moldova) makes the product Primobol. It is prepared in both 1 mL ampules and multi-dose vials.

Proviron® (mesterolone)

| Androgenic | 30-40 |
|---------------------|--|
| Anabolic | 100-150 |
| Standard | Testosterone propionate |
| Chemical Names | 17beta-hydroxy-talpha-methyl-Salpha-androstan-3-one 1-methyl-Salpha-dihydrotastosterone |
| Estrogenic Activity | none |
| Progestational Acti | ivity not significant |

Description:

Proviron® is Schering's (now Bayer's) brand name for the oral androgen mesterolone (1-

melyi dipotensiotenene). Semir to dipotensiotenene, mesterokene is a strong antogen with only a weak level of anabolica scirkly. This is also the fact that the dipotensioteneous, mesterokene is mpikly reduced to incretive deal methodenes in marktication of the structure of the structure of the structure of the structure of the antogen receptor is market issues, hereby reducing the gains of other mere potent meskbaling structure, doord likewise net be islans servicely in fact, doord methodenes affluight the structure of the structure of the structure of the structure of the affluight prediction of the structure of the structure of the structure of the affluight prediction of the structure of the structure of the structure of the affluight prediction of the structure of the structure of the structure of the structure exercise.

History:

According to computy lemane, Schring dowlped Provinsili in 1924, making this is an enternely of medication pat in a clickal practice for the neutranet of "hormen-velocid diseases and completion in mar."According in, neutronic would have bond coledpoid diseases and completion in mar."According in, neutronic would have bond coledpoid which are both very oil agents guerrally considered would will start both with any both very oil agents guerrally considered would will start the oil is age. Provints in a korg history of clickal effectives and address and start, and remains widespread clinical use today. It is generally prescribed to makes for the transmet of histo canced by instificient andragen levels, hypogendium (in pre- and poor produced bible) canced by instificient galaxies and the start bar produced presents.

The use of mesterologe as a furthy aid is perlups one of the most controversial indications for the sing consisting that ambidicationogene strends are generally liked to infinitely, it is also as need mesterologe that is quite often missachensed by affects. Mesterologe is applicable here because it is an effective nucleon that offers minimizappression of grandotrips approximation of the single strends and the single strends are production. It is subject to a single strends and the single strends are production. It is subject to a single strends and the single strends are production. It is subject to a single strends are single strends and the spectra strends and the single strends are single strends and the single strends are production. It is subject to a single strends are single strends and the single strends are production. It is subject to a single strends are single strends are single strends and strends are single strends and the single strends are single strends and are single strends and a single strends are single product affinitive effects on the equilibrium. Attenden and a single strends are single product affinitive effects and the equilibrium data strends are strends are single product affinitive effects and the equilibrium data strends are single strends are single strends are single strends are strends and the single strends are strends are single strends are single strends are constrained by the aspects and are single strends are strends are single strends are single strends are constrained by the aspects are single strends are constrained by the aspects are single strends are single

Mastrachone is widely manufactured by Bayer (formerly Schering), which corrently sole the drug in more than first coursities workshold. The most common breast manue and for its sole is Provine, although Schering/Bayer has sold the agent rather other mores in certain mestachone core the upser, appearing under each breat murnes in Parision (Aceko, Germany), Voitrano (Lenephurn, Germany), and Restore (Hooss & Barke, Infah). In sple of is long tank-renorms and analysis, many Sustemmington, however: Bayer remains the order breach of the date on the bayer and the sole of the barke of the barke of the barkes of the ba

How Supplied:

Mesterolone is widely available in human drug markets. Composition and dosage may vary by country and manufacturer; preparations generally contain 25 mg or 50 mg of steroid per tablet.

Structural Characteristics:

Mesteroloue is a modified form of dhydrotestosterone. It differs by the addition of a methyl group at carbon I, which helps protect the homeone from hepatic metabolism during oral administration. The same structural modification is also used with oral Primbolamid (methenoloue) tablets. Alkylation at the one position slows hepatic metabolism of the steroid during the first prass, althoud much likes profoundly threat -17 alpha alkylation. Mesteroloue and the steroid during the steroid during the steroid during the first prasma and the steroid during the first prass, althoud much likes profoundly threat -17 alpha alkylation. Mesteroloue and the steroid during th is resistant enough to breakdown to allow thempertically beneficial blood levels to be achieved, although the overall bioavailability remains much lower than c-17 alpha alcylated on alteroitik. Meestoolen also has a very strong binding affaitify GT See Hormone Binding Globalin⁷²⁷. This may act to displace other steroids more weakly bound to SHBG into a free (active) state.

Side Effects (Estrogenic):

Mesterolone is not aromatized by the body, and is not measurably estrogenic. An antiestrogen is not necessary when using this steroid, as the drug is unlikely to induce genecomstair, water retention, or other estrogen-related site effects.

Mesterolore is actually believed to act as an anti-acomates in the body, preventing or skowing the conversion of servoice into consequences. The result is convolute comparable to Artimickelly, although less profound. The arti-senzogaine properties of mesterolore are not minge, and an annihor of order attention have demonstrated using activity. Displatestosterone and Masteron (2-melly-displayed)sestemers), for example, have been successfully used as mellenging for agreement and been suggested for annalosies may come lower anomates activity in projenomic successful program constant on tengons endogenic disc of all of these comparable is presumably ensueed by their ability to compete with other arbitrations for brinding to the communities comparable in the aromations ensures bond on the sereoid, yet being marks to a the *x*, an inhelping effect is achieved as it is temporarily blocked monitoriation.

Side Effects (Androgenic):

Mestrohrs is chosifed as an antiooparie steroid. Andopprie side effects are common with this substance, expectively with light docs. This may riched boost of oly kin, are, and body/facial luri growth. Antabolenthogarie steroids may also agarsate male pattern har is bost. Worten are also worted of the potential virking effects of antabolenthogaries steroids. These may itselke a deepening of the voice, merstaal insegativities, changes in skin texture, ficial luri growth, and clone disparsent. Additionally, the 5-aften andtesse erapma does not metabolize mestavative. Is is relative antogarisely is not affected by finsertive of unstavelide.

Side Effects (Hepatotoxicity):

Mesterolone is not c17-alpha alkylated, and not known to produce hepatotoxic effects; liver toxicity is unlikely.

Side Effects (Cardiovascular):

Ambolicantoganis stavalis ca miwe debatriose effects on serum chocksterol. This includes a tendency to reduce MIE (agod) chocksterol values and increase MIE. (bad) chocksterol values with the HDL (agod) chocksterol values and the MIE. (bad) chocksterol values with the HDL to LDL balance in a direction that flowors genet risk of a directional tender on the dose, route of administration (cetal vs. ijectedbe), type of sterold genomiable on consonnitable), and beel of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable, and the def of resistance to heptic mitobleom consonnitable and the def of heptic mitobleom consonnitable and the def heptic mitobleom consonnitable and theptic

Mesterolone should not be used when cardiovascular risk factors preclude the use of other oral steroids.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4) grams per day) and a natural cholesterolaritoxidant formali such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Mesterolone has a very weak suppressive effect on gonadotropins and serum testosterone. Studies show that when given in moderate dooses (150 mg per day or less), significant suppression of testosterone levels does not occur.⁵⁷⁴ In studies with higher dosses (300 mg per day and above), the agent strongly suppressed serum testosterone.⁵⁷⁵

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

To text antropyn isndfrietery, mesterohne is smally given in a dose of 1 table (125 mg) three times per day with a indiation of theoryn. The dargs is later continued at a lower matterance dose, which usually consists of taking 1 table (125 mg) one to two times per day. Similer doses are ued to support table firstly, smally in comparison with other fathry drags like sjoctable FSH. The usual dosage among mak althests is between 50 mg and 150 mg of mesterohore per day or two to siz 52 mb/s. The day as pipoidly laten is typical of 6.12 weeks in length, which is usually a sufficient period of time to notice the benefits of due theorem.

May bolyhalkes floor the use of mestroken during during phases or contest proparation, when box strength and light and though needs any particular distribution of the sepecially branchical when markolski like Winstolli, Aranna, or Primobalnili and brains and difficulty and here to adjust the androgen to estrogen mate spreader. Mesterobies can be effectively used here to adjust the androgen to estrogen mate spreader. It is also also also also also also and the androgen to estrogen mate spreader. The strength of the second with brain and interacting the strength year constrained between the second s

Administration (Women);

Mestrohow is not approved for use in worten. This agent is not recommended for wortene for phospices - or preformance-of-metric prayness due to its strong undrogenic nature and tendency to produce villing side effects. Some wortene to lives the dag, however, and instruction to produce villing side effects. Some wortene to lives the dag, however, and any structure of the structure of the structure of the structure of the structure practicity the low of relationships that the structure of the structure of the or the works is such structures to minimize the chance of developing lasting writing effects. One table used in comparison with 10 or 20 mg of Nobidade¹⁰ on the server more efficient for mass how for any cavely in an oniverserve here the body is much more isolated to bars whether the structure of the struc

Availability:

Mesterolone remains widely available, the vast majority of products made by or under license from Schering (now Bayer). In reviewing some of the more popular products and changes on the global pharmaceutical market, we have mode the following observations.

Bayer took control of Schering AG in December 2006. Following this acquisition, the Schering Provinon products have been transitioned over to the Bayer brand and logo.

Bayer no longer markets Proviron in Egypt. The drug remains available under the Cidoviron name, produced by the domestic firm CID (Chemical Industries Development).

Unigen markets the product Mesviron in Thailand. It contains 25 mg per tablet, and is packaged in foil and plastic strips of 10 tablets each (5 strips per box).

Due to its linited demand, mesterolone products have traditionally not been the subjects of high volume counterfeiting. When located on the black market, they can usually be trusted so long as they are properly packaged from a known manufacturer.

Sten (testosterone cypionate & propionate)

| Androgenic | 100 |
|------------|----------|
| Anabolic | 100 |
| Standard | Standard |

| Chemical Names | 4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one |
|----------------------|---|
| Estrogenic Activity | moderate |
| Progestational Activ | ity low |

Description:

Sen is a two-composent teststerve blad from Mexico fast cortains a mixture of teststerver projecuties (2 str.), teststerver csystemic (7 str.), and DFEA (dds)dprogramhosterver; 20 mg in a 2 mf, ampek. Sense references incorrectly fast its projectar is containing 20 mg of DFT (diplocatisatence), which would be a tittle androgen. This is, however; just a confision of the Spanish used for DFEA Mere recore producing lists this gradeet is the lass confisior; presented product androgen the DFT, holding the DFEEA released at the moment, first sense product contains and just 90 mg/mit mixture of two common teststerem centers.

May consider Start to be a low-holger alternative to Summer/B 250. While ideas contain a blend of two testorecen estres, Stars in a solw-acting to comparison. The longer oster of isotaternee it uses is coprennet, which allows testosternee levels to return to banding approximately veeks after jeticori. Toolstonene oppinue is also not addiqueorate dang, so Stars doors't offer metch advanzage in regulation a "assistand-released" effect. Topperson-bandler government, the isotationary oppinue was solved and phenomechanic phenomenolity in the isotationary oppinue was solved and phenomechanic phenomenolity of the isotationary oppinue was solved and Coarse Stars and Statametik and the isotationary oppinue the two.

History:

Stan is made in Mexico by the planmacerical firm Attants, SA de CV. This agent's used primarily to correct two androgen lock in males, for the transmotter of hypogenetisms, androgause, and impotence. It is also sometimes prescribed to avorant for excessive location, advanced buscet cancer, and low see of the Scanta fast advanced buscet that Mexican market, where it is one of the coartry's more inceparable human-set tostotence products. It has concequerely remained a somewhat proparit me three, were it is formation may not be the most properly saided for the higher-dosed negativenest of athket use.

How Supplied:

Sten is available on the human drug market in Mexico. It contains a blend of 25 mg/75 mg testosterone propionate and testosterone cypionate (respectively) per 2-milliter ampule.

Structural Characteristics:

Sen contains a minute of two testosterone components, which where modified with the addition of archoroly and clearses (propriors and cyclepartypropriors) acids) at the 17-beam hydroxyl group. Extertial draws of restosterone are less pairer fram free testosterone, and are absorbed more advorg from the area of injection. Once in the blockbarrau, the setter is narrowed to yield free (active) testosterone. Exteriled frame of testosterone are designed to injections should company to high testos free testosterone. The should be provide a randpr grade in testosterone levels (24-48 hours after injection), and marintain physiological concentrations for approximately 14 days.

Side Effects (Estrogenic):

Teostores is readily atomitted in the body to estabilish (estrogen). The aromnase (estrogen synthesis) compare is responsible for fine metabloom (to teostorene. E-board estrogen body and the synthesis is the stressed water restriction, body fut gain, and gencomistin. Teostorene is considered a monthly estrogenistic and estication of the stress of the stresses of the estimation of the stresses of the stresses of the stresses of the effects on blood light and the stresses of the stresses of the stresses of the effects on blood light and the stresses of the stresses of the stresses of the effects on blood light and the stresses of the stresses of the stresses of the stresses of the englished light and the stresses of the stresses of the stresses of the stresses of the effects on blood light and the stresses of the stresse

Encogenie side effects will occur in a dose-dependant memer, with higher doses (above normal therapetic level) of lostioaterem envel hely to require the concurrent use of an antistringen or aromatuse inhibitor. Since water reterion and less of mateix definition are common with higher doses of feastbarrent, tied mags a smaller dose and the concurrent with higher doses of feastbarrent, the disr again and the distance of the state of the state of the state of the state of the halking planes, where the added vater reterior will support now strength and much size, and help foster a value ratio.

Side Effects (Androgenic):

Teasterore is the primery multi-androgen, neoponsible for maintaining scontallary mulsenul characteristics. Florated levels of testatores are fieldy to produce androgenis deli efficies relating only skita, scare, and bodylikeial large growth. Men with a genetic profilopation for link links (androgenic altergation) may reface constrained by option halding. These concerned about hard has may find a mere constrained option in manufacture potential withing effects of analytic and the second second second second about the second se

In antiogen-response target issues such as the skin, such, and prostate, the high relative antiogenizy of testatores is dependent on its rokation to diphytotestostenere (IPI). The 5-alph reductsice corputs is responsible for this metabolism of teststores. The concentret use of 3-alph reductsies mithing variants of teststores with single to produce autopartic site strategies and the strategies of teststores with also-specific potentiation of teststores extra to sworting the tendency of teststores dupped to the strategies site effects. It is inportant to remember that analyke and separation of teststores' analyke and antiogenic properties is not possible, even with tool 3-alphar reductses inhibito.

Side Effects (Hepatotoxicity):

Testosterne does net hree bepatetoxie effects, for toxicity is attikely. One study examine the potertial for hepatoxicity with high loss of testosterne by administering 400 mg of the lormore per disp (2,800 mg per week) to a group of mile subjects. The staroid was attion only to similar higher peak concentrinos would be reached in legatic tissues produced to significant charges in lore enzyme values including serum absuris, bifrubin, attine-amine-amines, and Jackine toxicatuses.²⁰

Side Effects (Cardiovascular):

Ambelicant/arganic strends can have deterious effects on serum checkeror. This includes a tendency to role and EUL (good) checkerou values and increase (LL, (bat) checkerout values, which may shift the HDL to LDL bahare in a direction that fixors greater risk of antiroicherosis. The refine impact of an antibolicant/ognic strends of a serum lipit is dependint on the dose, route of administration (et al. vs. injectable), type of steady instructions, the service impact of an anti-off the service of the service of the service of the service of the service interval of the service impact on the service impact of the service of the service of the service interval of the service interval of the service interval of the service interval of the service of the service interval or the service interval or the service interval of the service interval or the service interval or the service interval or the service interval interval or the servic Tensterous tends to lave a much less damatic inpact or cardiovascular risk feators dam synthest stocks. This is dae just to is openness to muchiohumby the feasi, which also it to have less effect on the lepatic management of cholesterol. The aromatisation of the stocks and the level of the stock of tensterous energy of androgen on sense matically significant distributions and the level of tensterous energy of the stock of tensterous tensors and tensors and tensors and the level of tensors and the level of tensors and the matically significant of the level of tensors and the level of tensors and tensors there (original tensors and the level of tensors tensors with tensors and tensors and mathematically significant and tensors and

Due to the positive influence of estimgan on serum jujich, sumwiden cirate or colompione cirate are porterior to a commonse influitors for these concerned with a caldwascular heads, as they offer a partial estrogenic effect in the lose. This also shows then the potentially improve injurbances and the source of the regardneet fields of antiogenet. Which does of 600 mg per lose indicating the fractual production is provided with the term fractual but are durative, imaging an anisotrangen (fractual) enderstation and the production of the source of the source of the indicating and interchardenic purposes the product and indicating an anisotrangen of the conduction of the source of th

To help reduce cardiorsecular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterolamitoxidant formatia such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic stewisk when taken in doess sufficient to promote maske gain are expected to appresse molgrouss testsotreem production. Testsotreem is the primary mark androgen, and offers strong engine feedback on endogenous testsotreous production. Testsotreem-based large ville. Rowsie, here a strong effect on the hypothamic regulation of natural stewid hermions. Without the interversion of testsotreom-similarity advances, testsotreem-based based trendo normal within 1-4 month of data gescession. Note that prolonging thereorem, the strong strength terror test and the propagation of the strength of the stren

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Testosteone propionte is often regarded as a painful injection. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to issues at the site of injection. May sensitive individuals choose to stay away from this steodic completely, their bodies reacting with a pronounced soreness and low-grade fiver that may last for a few days after each injection.

Administration (Men):

For the treatment of low androgen locks, the prescribing guidense for Stan recommend one spinorfmont or 2m. Tampid (100 mg testschore esters, 2m (2m [2M-2wert) 15-40 a)), For bodyhaling purposes, this ding is usually lipicted on a weekly basis, in a doage of 2-4 angeles (200-400 mg of testschore esters in toul). This lock is sufficient to provide excellent gains in macks size and strength. Higher does use possible, but even the injection volume neekled with a ramples per week (100 can become to useroffenhib for many Testschore drugs are ultimately very versitik, and can be combined with many other ambidicativeness treated selected from the device diffect.

Administration (Women):

The preserving galdense for Sten do not make special dosing recommendations for worner, except to synthat androgarie symptoms may occur, and in a certain scentrisc therapy should be suspended untl symptoms resolve, and after a lower dose used. This drug is not recommended for worners for physical-or or performance-relaturing purposes due to its strong androgarie nature, tendency to produce viribing side effects, and slow acting characteristic (mining blod levels difficat to corrot).

Availability:

Sten is commonly found in Mexico, where 2 pre-loaded syringes are packaged in a box and usually sell for about \$10 in the pharmacy.

Striant® (testosterone)

| Androgenic | 100 |
|-----------------------|---|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Activi | ity low |

Description:

Straint's is annexaficasies baccal associations delivery system. It is prepared in the ferm of a small (aspiris-sized), tablet, which comings 70 mg of (free) isosteneons. The table it as taken endly, but is affield to the game, where it transfers isosteneous across the invert fange on the system of the system methylatoristome and testosteneous propriates. The affiesdes Straint system, however, it as improvement over the store (and produced) and the system of the system of the model (it tables) affiesd for 12 hours and reach to be remrough. As each, it provides a much enough to matting hybridizable intervention of the system of the system.

History:

Straint? was developed in the Unided States by Cohrrish Labouncies. It was approved by the FDA for solar as precerciption drug in Just or 2002, and is infrained for use in mu with conditions associated with a defeative or absence of endogenous testostences. With this product, Cohrriba was Bolely virging to angle those homess replacement freezays (HBT) consumers that do not weakowski by accions, and if a patches and gals unconditrible or constrained policyteneith. Strain vas adapted to plannesis in the 2003, and quicklin others are with much reviews. Strain galaxies find it a vary consensator option for HBT, which others that U.S. 1000, and the Strain SH (Sciaman Behave) bread much site. In 2004 and quickly Adama Broscience. Columbia's pattnership with Adama expects to see sake of Strain it 18 Emperanments.

How Supplied:

Striant® mucoadhesive buccal testosterone delivery system is available in various human drug markets. Product comes in the form of a small buccal tablet; usually packaged in strips of 10 tablets, 6 strips to a box.

Structural Characteristics:

Straint® mecodlassic bacal lastosterore délivery system is a bacal labét containing 30 mg of (fiéce) testoren. The system a delived to the individue of the month, where the gam meets the upper Tg above the incise toeft With exposure to saike the tablet softens in a gable constitution, which can stay in place for 12 huns: The product delivers physiological concentrations of testosterone through the macous merhane, where it is absorbed into the strengt softens and a superior was cancel imprible lowessith, plassing the lare.

Side Effects (Estrogenic):

Teodotrone is readily anomined in the body to estable (energen). The anomatese (storgen priftensis) energies is proposible for this metablism of isotatorous. Ebseudi establismes and the store of the store of the store of the store of the store theorem of the store heapenetic does well increase the likelihood of establismes devices in store of the store comparison with a champlene estimation of the store (mainstands), which more efficiently correds estimation to also store of the store of the store of the store (mainstands), which more efficiently correds estimation to also store of the store of t

Side Effects (Androgenic):

Teststerors is the primary must androgen, responsible for maintaing secondary mulssound entractivities. Decoding normal therapic discoss is facely to produce androgenet siste effects rickafing oby sist, anex, and bodyficial larit growth. Men with a genetic professioni for larit usis (androgenetic alsopecin) may note accustented multipatient based on the second second second second second second second based on the second second second second second second second expectively with a storey androgen accus based second second second second work, ensembla in egalarities, changes in skin texture, facial hair growth, and clional enlargement.

In antiogen-response target tosses such as the skin, scale, and prostate, the high relative antogenizely of tostatores is dependent on its rolation its oflydivostatorene (RHT). The 5-alph reductsice corputs is roporable for this metabolism of tostatores. The concurrent use of 3-alph reductsies mithing variant statistical will reference with also-specific potentiation of tostatorene action, towering the tendency of tostatorene datasets and the statistical segmention of testostatorely analysis and antiogenic properties is not possible, core with tost 3-alphar reductses inhibito.

Side Effects (Hepatotoxicity):

Testosterne does net hree bepatotoxie effects, for toxicity is attikely. One study examine the potential for bepatoxicity with high does of testosterne by administering 400 mg of the lowmee per day (2,800 mg per week) to a group of mile subjects. The secoid was attain orably so that higher peak concentrations would be marked in higher functions and the study of the produced to significant charges in liver enzyme values including serum absuris, bifruibis, attine-amire-amires, and abalatine boostnates. See

Side Effects (Cardiovascular):

Ambeichardurgenis stavalis can huve delatricos effection on serum checkerot. This includes a tendency to reduce IRL (apod) checkerol valesa and increase IRL (bad) checkerol valesa, which may shaft the HIM. to LID, tahaine in a direction that flowes gener risk of an entroper tensor of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the start of the start of the start of the direction of the start of the direction of the start of the direction of the start of To help reduce cardiorascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intrake of startated fats, cholsterol, and simple cardohydratises at all times during active AAS administration: Supplementing with fish oils (4) grams per day) and a natural cholsterolamicoxidant formals such as Lipid Subil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicalization stewisk when taken in doess sufficient to promete maske gain are expected to suppress endogenose testsetores production. Testsetores is the primary mark androgen, and offers strong negative fordback on endogenose testsetores production. Testsetores-based lange voll, Bavoice, here a strong effect on the hypothaniar regulation of natural stewid hormores. Without the interversion of testsetores-straining substances, testsetores levels and tertorio normal within 1-4 morth of data gescession. Note that prolonging thereorem the strength tertorio normal testsetore levels substances.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

The Striant macoadhesive buccal testosterone delivery system is placed on the gums just above the incisor tooth. It is left affixed for 12 hours, at which point it is carefully removed. The product is usually administered twice daily. The application site should be rotated between left and right sides of the mouth with each dose.

Administration (Men):

To text androgen isudicisco; the preceiving galdens for Struit recorrect administruing the Result bark trick of high loss ar given cost in the rooming and once of right, L Bors apart. For physica- or performance enhancing papeose, higher does a structure of the systems per day, a loss and the control. Loss of the structure of the accompanies lay of the structure of the structure of the structure of the accompanies of the structure of the struct

Administration (Women):

The Striant macoadhesive buccal testosterone delivery system is not FDA-approved for use in women. Testosterone is not recommended for women for physique- or performanceenhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Given its high relative price and low delivery of testosterone, Striant is not commonly traded on the black market. Counterfeits have not yet been reported.

Sustanon® 100 (testosterone blend)

| Androgenic | 100 |
|------------|----------|
| Anabolic | 100 |
| Standard | Standard |

Chemical Names 4-androsten-3-one-17beta-ol

| 17beta-hydroxy-androst-4-en-3- | |
|--------------------------------|----------|
| Estrogenic Activity | moderate |
| Progestational Activity | low |

Description:

History:

Sutanziii 100 is a medem adaptation of the well-krown injectable istosteone blend Sustanziii 250, Ohot of wichs were developed by the intermittodin planmocarical gaint Organon (now Merck/MSD). Sustanziii 100 is essentially a lower dosed equiokare of Sustanziii 250, applysing the same hermone in saintly (hotgon rule cacch) time-tokand faishen. Sustanziii 100 is recommended for the same medical uses as Sustanziii 250, applysing the same hermone in sing the dogn hot acch tyme-rokand methy enting rule at handongs insufficarcy. Which can mundit at both such supports as neduced see drive, importance, infitting, and hora loss hermositi and addition to these uses. Sustanziii 100 aks horgenorsas data in these medicalization in famile-to-mist transsenank. Sustanziii 100 aks horgenorsas data in these medicalization i famile-to-mist transsenanks statistanzoii 100 is produced only in a hantilia of countris at their me and is not whele worked worked and the same famile and the same family and statism in the messaria. Sustanziii 100 aks produced only in a hantilia of countris at their me altic net whele worked worked hore.

How Supplied:

Sustanon® 100 is available in select human drug markets. All products are supplied in 1 mL glass ampules.

Structural Characteristics:

Staturnil 100 contins a mittare of three testsdetone compounds, which where medified with the addition of carbody is caid sets reprojents, propriori phenyles level sets, and location acids) at the 17-bate induced grange. Earthful frame of testsdetone are less pair than five blocdromen, the exist is neared to blocd the discretion of the set of the testsdetone are designed to prolong the virklow of themposite effect following administration, allowing for a less fequerit prices selected compared to injections level (masterified) started. Statistica (10) is designed to provide a might pair is tastistications levels (24) how and the pixelicity, and matterin physical concentrations for approximally 41.

Side Effects (Estrogenic):

Textsteres is readily atomitted in the body to estability discourse, it has anomates (estrogen synthesis) entrys is responsible for this metabolism of textstorme. Elevation establishes and the state of the state of the state of the state of the state state at complexes is considered at metabolish estimates at texts of the state state at complexes characterized at the state state of the state

Estrogenic side effects will occur in a dose-dependant manner, with higher doses (above normal therapeutic locale) of test actorize more likely to require the concurrent use of an antiLEMIT BEARPORT EVENO IN SERVICEMENT INCE INCOME OF OF OPEN THE CONTENT AND A DESIGNATION OF A DESIGNATIONO OF A DESIGNATIONO OFFICIAL DESIGNATIONO OFFICIAL DESIGNATIONO OFFICIAL DESIGNATIONO OFFICIAL DESIGNATIONO OFFICIAL DESIGNATIONO OFFICIAL DESIGNATIONO DESIGNATICO DESIGNATIONO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATIONO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNATICO DESIGNAT

Side Effects (Androgenic):

Textsteres is the primery multi-androgen, nesponsible for maintaining soccouldry mulsesual characteristics. Encodel texts of textsteres are fields produce androgenic side effects including objection, and bodyficial line growth. Mon with a genetic profilopointion first hills too (androgenic dayce) imay notice accelerated multitating. These concerned about hirds is may find a new conficiently option in manufacture acceleration of the state o

In anthogon-exponsive target risous such as the skin, tody, and prostate, the light relative antogenizely of straturenes is dopendate on its rolation to display disolationaries (PLT). The 5-alpha reductisse enzyme is roporable for thin metabolism of testisterions. The concurrent use of 3-abb reclusters inhibitors such as fustative disolations of testisteries with also-specific potentiation of testisteries action, lowering the tendency of testisteries disolation of testisteries and the strategiest of the strategiest of testisteries asymptotic of testisteries inhibitor.

Side Effects (Hepatotoxicity):

Testosterne does not huw bepatotoxie effects, for toxicity is attikely. One study examine the potential for hystoxicity with high loss of testosterne by administering 400 mg of the hormore per day (2,800 mg per week) to a group of mile subjects. The second was station only so that higher peak concentrations would be reached in hystoxic issues concentration of the statistic statistic statistic statistic statistic production of significant changes in high rengence statistic statistic statistic and the statistic statistic statistic statistic statistic statistic statistic statistic and the statistic statistic statistic statistic statistic statistic statistic statistic and the statistic statistics and statistic statistic statistic statistic statistic statistics and the statistic statistic statistic statistics and the statistic statistic statistic statistics and the statistic statistic statistics and statistic statistics statistics and statistic statistics and the statistic statistic statistics and statistics and statistic statistics and statisti

Side Effects (Cardiovascular):

Ambolicanteganis strenids can huve delutricos effection on serum chocketorel. This includes a tendency to reduce IDML (apod) chocketorel voluces and increases (LL, (bad) chocketorel voluces, which may shift the HDL to LDL balance in a direction that fivores genet risk of antirocherosis. The evaluation match characteristic and on serum high is dependint on the dose, route or a daministration (eral ws. injectable), type of steadid Anabholicanteganis streamed, match and anticella streame piles of anabholicanteganis streamed, match and steapeness, fluct board process and infegerations, routes excludebili reducesion, and support left wertricular hypertraphy, all posentially increasing the risk of cultoroscelar disease and monecularit lifeticion.

Teostorem tends to have a much loss durantic impact on candiovascular risk factors than synthest seriods. This is due part to is porcess to muchiolomisty the focus, which also it to have less effect on the lepatic immagenet of cholesterol. The aromatization of tistustances to estudiated also helps to might the magnite effects of androgen on senarmistically significant disk of the original effects of the solution of the solution instandary displicant disk of the original effects of the solution of the theorem of the solution of the solution of the solution of the solution of the teorem of the solution of the solution of the solution of the solution of the teorem of the solution of the solution of the solution of the solution of the teorem of the solution mached 2 μ ^(5,64). The negative impact of aromatese inhibits should be taken into considention helps exclutions in the solution relation of the solution in the solution in the solution in the solution of the solution is the solution of the solution of

Due to the positive influence of estrogen on serum lipids, tamoxifen citrate or clomiphene citrate are preferred to aromatase inhibitors for those concerned with cardiovascular health, as they offer a partial estrogenic effect in the loce. This allows them to potentially improve high profiles and offers some of the negative effects of androgens. This does of effort more loss of testostence per week, the impact on lipid profile trade to be noticeable but not domain; mixing an mix-testogen (free catalyces, statisticably significant damages Does of effort may be used by the also field to produce statisticably significant damages and mixing statisticable trade to the statisticable statisticable statisticable statisticable statisticable statisticable statisticable statisticable statisticable with a statisticable statisticable statisticable statisticable statisticable with a statisticable statisticable statisticable statisticable statisticable statisticable statisticable with our statisticable statisticable statisticable statisticable statisticable statisticable with our statisticable s

To help reduce cardiorascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of startated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fash oils (4) grams per day) and a natural cholesterolamitoxidant formals such as Lipid Stabil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicalização stavidis veloa takan in doses sufficient to promete massé gai are expected to supprese condigoras testostorem producina. Testostenes is the primur prate androgan, and offers streng negative fordback on endoganous testosterone production. Testostenes-based lange valle, Baveice, here a strong effect on the hyporhamic regulation of natural steroid hormouse. Without the interversion of testostenors-stimaling substances, testostenes levels durit levels of the strong based on the program durity and produced program. Testos and testostenors stimaling substances, testostenes levels durit terrors to mail within 1-4 monti of ding secession. No edit prostagit programdatorepite lipoparadism can develop secondiny to steroid abuse, messating medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Testosteone propionte is often regarded as a painful injection. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to issues at the site of injection. May sensitive individuals choose to stay away from this steodic completely, their bodies reacting with a pronounced soreness and low-grade fever that may last for a few days after each nigeritoria.

Administration (Men):

To text androgan isufficiency, the prescribing galdelines for Statunotifi 100 cull for a domagorf 100 mg (1) mg/k) every 2 weeks. Altongha rise in the holy for a long ritro, Statunofi 100 is usually isjected every 7 to 10 days for much-backing parposes. This scheduk will allow for the lagher does meru controlly applied parlies, and rower stable elevators in homme level. The usual doags among mixe althests in the range of 200-600 mger week, lake in cycles to 12 weeks in and, trangh. Tabstots in the star mger week, lake in cycles to 12 weeks in and strength. Tabstots and strength and notice encorptional gains in much is and strength. Tabstots in strength wynessile, deviated lists.

Administration (Women):

Sustanon® 100 is rarely used with women in clinical medicine. When applied, it is most often used to induce musculinization in female to male transseaulas. Sustanon® 100 is not recommended for women for physique- or performance-retraining purposes due to its strong antrogenic nature, tenkency to produce virizing side effects, and slow-acting characteristics findinge book levels difficult to corrorly.

Availability:

Sustance® 100 is less widely distributed on the black market than Sustance® 290, due to the fact that source countries producing this drug are limited. The moderate total amount of drug contained in each ampule (100 mg) also markes this product far less desirable to consumers than Sustance® 290. Sustance® 100 is mainly located in the Netherlands, Egopt, and the 1 lixed Viewolow.

Sustanon® 250 (testosterone blend)

| Androgenic | 100 |
|--------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activit | y moderate |
| Progestational Ac | tivity low |

Description:

Staturnili 2016 is no obsoci piectable testoaterene bierd that cortains for different tosisotence exists existencere projenical (C) mgk testostence phenophysonical (O) mg, testostence exists existence (O) mgk testostence phenophysonical (O) mg, testostence is existence (O) mgk testostence phenomena (D) mgb, Staturnili 16 or existence (O) mgk testostence (D) mgk testostence (D) mgb, Staturnili 16 testostences such as copienta or emuthate, biels provide a shorter duration of activity. As with all testostences products, Statamoli 20 m a very simulate (mgk testostence) testostences such as copienta or emuthate, biels provide a shorter duration of activity. As with all testostences such as controls used in balance phenomenol protocol and mgrane and mgk testostences controls used in balance phenomenol staturnili (D), and testostence (C) and testostence (C) and the staturnili (C) and testostence (C) and Staturnili (D), and testostence (C) and testostence (C) and testostence (C) and Staturnili (D), and testostence (C) and testostence (C) and testostence (C) and Staturnili (D) and testostence (C) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence (C) and staturnili (D) and testostence (C) and testostence

History:

Statusnik 220 first append on intermittend dag markets daring the early 1070% It was developed by the intermittend parameterization jater (Dagen (norm (norm NetzANSL)), and responsible for such strends as Daracholisti, Daes-Darabolisti, and Androld, Statamoni which need to be interped mere fragerary (focal advantages in hormers stability are probably nor takil). In spite of this subartage, however, Statamoni 220 has never been appender branch of testasterum (advantage) and the world in the order of the network of the hist advantage. In the land Statas, advantage and the total probably nor takil). In spite of this subartage, however, Statamoni 220 has never been papaker human of essenterum arabidite. The lack of ULS availability is probably of the or something origination of the hist hyperbase in the total spite. Statistical for the statistical origination of the hist hyperbase in the statistical of the origination origination of the hist hyperbase in the statistical of the origination origination and the hist hyperbase in the statistical of the origination origination of the hist hyperbase in the statistical of the statistical origination of the hist hyperbase in the statistical of the hist hyperbase in the statistical origination of the hist hyperbase in the statistical statistical origination of the hist hyperbase in the statistical statistical origination of the statistical statistical statistical statistical statistical origination and the statistical statistical statistical origination and the statistical statistical statistical statistical statistical origination and the statistical statistical statistical statistical statistical statistical stati

Over the past 25 years, Statamolf 220 has probably heren the most sough-after injectichle stostoneor among addites. It must be empisical, however, that this is not due to an unusual potency of this tostostoneor combination (esters really only affect the relaxes of tostostoneor). This is simply due to the fact at a stack of four different tostoareous components is a very good selling point; it is preceded to have more value. In most in startes: point migrature light of the fact have point of the most index of the light of the point point migrature light of the fact have point of the homefaction condition (fact have point point migrature) and a this fact have point on the homefaction condition (fact have point point migrature) and a this fact have point on the homefaction condition (fact have point point migrature) and a statement of the statement of the sound allow year to side inhore the how fact have point in the statement of the sound allow years to side in more than how fact have point on the statement of the sound allow years to side in more than the how fact have point on the statement of the sound allow years to side in the sound allow years to side in more than the mission of the sound to a statement with the sound allow years to side in more than the mission of the sound to a statement of the sound allow years to side in more than the mission of the sound to a statement of the sound to a statement with ender the relaxed to the sound to a statement with the sound to a statement with ender the relaxed to the sound to a statement with ender the relaxed to the sound to a statement with the sound to a statement with ender the relaxed to the sound to a statement with ender the sound to a statement with ender the sound to a statement with ender the relaxed to the sound to a statement with ender the relaxed to the sound to a statement with ender to the relaxed to a statement with ender to a statement with ender to a statement with the sound to a statement with ender to a statement with ender to

How Supplied:

Sustanon® 250 is widely available in human and (select) veterinary drug markets. Packaging volume may vary by country and manufacturer; the majority of products are supplied as 1 mL gass ampules.

Structural Characteristics:

Sustanon® 250 contains a mixture of four testosterone compounds, which where modified

with the addition of carboxple acid esters (proprise), proprise phenyl ester, isceparo), and decomes caids) after 17-ben hydrody gones. Districted from of structures are a loss port fram for testosterone, and are absorbed more slowly formule area of rejection. Once in the bloodbream, the setti is enreved in lysid the (acidon) testosterone are lossing in the structure and designed to provide a grad the gradient of the structure (investificial) second. Structure 2016 designed to provide a grad particular the structure of the (24-48) more after significant of the structure of the structure of the structure of the (24-48) more after significant of the structure of the structure of the structure of the (24-48) more after significant of the structure of the structure of the structure of the (24-48) more after significant of the structure of the structure of the structure of the (24-48) more after significant of the structure of the s

Side Effects (Estrogenic):

Teostorene is readily anomized in the body to estabilid estrogen). The anomases (estrogen synthesis) engras is responsible for fine metablosm (in Stostorene, Elevation establishes) and cause side effects such as increased usater returner, body far gain, and gencomistin. Teostorene is considered a moderable estrogenic atrouble and effects. One may disentative usa anomalases inibiale lad Arrithdeen (increased), which effects the teored prediction of the strongenic strongenic side effects on them observation to anti-estrogens, however, and may also have negative effects on blod leights.

Estogaris side effects will occur in a doss-dependant mumer, with higher doses (above normal therapetic level) of Sustantom² (25) more likely to require the occurrent use of an arti-settogaro e aromatose inhibite. Since water retention and loss of muscle definition are common with higher doses of foststatence, this drug as usually considered a poor checks for backing plasses, where the added water retention will support most strength and muck size, and help foster a stronger anabole, environment.

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for minitaring scoradary mulssenal characteristics. Encode labels of testistores are likely to produce androgenic side effects including objective strength sectors are likely to mode androgenic side professionistic neith tubes (androgenic altery strength sectors) and the sector professionistic neither the likely strength sector strength sectors halfing These concerned about hird is non-print at more conficiable option in manufacture document, whiting and the strength sectors are strength sectors and the start and the sector strength sector strength sectors are strength and such as tostoreme. These may include document approximation, means and and strength sectors and the strength sector strength sectors and strength sectors and the sector strength sectors and the strength sectors and the strength sector strength sectors and the strength sectors and the strength sector strength sector sectors and the strength sectors and the strength sectors and the strength sector sectors and the strength sectors and the strength sectors and the strength sectors are strength sectors and the strength sectors are strength sectors and the strength sectors are strength sectors and the strength sectors and the strength sectors are strength sectors and the strength sectors and the strength sectors are strength sectors and the strength sectors are stre

In anthogon-exponsive target risous such as the skin, tody, and prostate, the light relative antogonizy of tostatorene is dopedual on its rolation to only diveositatoriene (PLT). The 5-alpha reductisse enzyme is roporable for this metabolism of testostatoriene (PLT) exposed in product and phare reduction with the structure of the interview with alse-specific potentiation of testostatoriene action, lowering the tendency of testostatoriene (PLT) and the structure of the structure of the structure of testostatoriene with alse-specific potentiation of testostatoriene action, lowering the tendency of testostatoriene segmention of testostatorie's autoble and antloguie properties is not possible, even with tool 3-alphar necktatorie inhibition.

Side Effects (Hepatotoxicity):

Testosterne does not hree hepatotoxie effects, her toxicity is utilicly. One study cearning the potertial for hepatoxicity with high does of testosterne by administering 400 mg of the hormone per day (2,800 mg per vised) to a group of mak subjects. The staroid was compared to intramedari ignitization. The hormore was given days for days, and produced to significant charges in Nerror testors including serum abstrain, bihrubia, attinica-mito-transforms, and abalant productatos.³⁰⁷

Side Effects (Cardiovascular):

Antobicaringeme sterodis can have deterroos effects on serum chosteror. The metades a tendency to reduce 10.0L (oud) choiseava in recence 10.L (bud) choiseava values, which may shift the IDLs to LDL bahare in a direction that favors greater risk of attransformers. The reduce transformation that the start of the start dependant on the dose, route of administration (reduce), signed halo, up of steadid Anabhicinthergues at more than the start of the start of the start Anabhicinthergues attransform (has a dassay affect the of present and trajectorists, rockae excludential reduced), and support left warring hepertuply, and potentially increasing the risk of cultimoscate direstes and meanculai historicon.

Teostores tends to have a much iss deamtic inpact or cardiovascular risk factors than synthest storich. This is an part to is operation mesh neurons in mucholomistic be focus, which are it to have less effect on the lepair immagement of choisetsof. The aromitation of tistustances to estimated also helps to might the magnite effects of androgen on sensuficial, in the sensity 250 mg per vock of tossoberore store (candidate) had a slight to not standardly significant effect on ERD, choisetsof the 12 works, but hank not with an anomalous tendence of the start denomination of the start of the start of the start of the start of the anomalous tendence of the start of the start of the start of the start denomination of the start denomination of the start of the machine start of the start denomination of the start of the machine start of the start denomination of the start of the st

Due to the positive influence of estingan on serum lipid, numoridin cirtate or electrical on a continues influence for these concerned with a calcionscalar headure, as they offer a partial estrogonic effect in the loce. This allows them to potentially improve flag roloss and offers some of the engines effects of anthogense. This does of 600 mg mg excludentic, multig and missioning of the calcionscalar headure does and the enginest of the section of the enginest of the effect of the effect of the effect of the enginest of the enginest of the enginest of the enginest of the effect of the effect of the effect of the enginest of the enginest of the enginest of the enginest of the effect of the enginest of the eng

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fab oils (4) grams per day) and a natural cholesterolamicoxidant formain such as Lipid Subil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic steroids when taken in doess sufficient to promete masely gain are expected to asprace condigenous testoatoreary production. Testoatoreas is the primary mark androgen, and offers strong negative feedback on endogenous testoatoreae production. Testoatorea-based dags sufficiencies, have a strong effect on the lynothimist regulation of natural strond hermoreas. Without the intervention of testoatoreae-similaring assistance, testoatoreae levels should return to menth within 14 months of dag scession. Note that prolonged hysogeneidstropic hysogeneidsm can develop secondary to steroid abse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Testosterone propionte is often regarded as a painfil injection. This is due to the very short carbon chain of the propionic acid steter, which can be irritating to issues at the site of injection. Many sensitive individuals choose to stay away from this steoid completely, their bodies reacting with a pronounced soreness and low-grade lever that may last for a few days after each nigeritoria.

Administration (Men):

or used manopart neuroscience, use processing generates as constructed as a construction of the second s

Statuntify 20% is usubly incorporated into bulking phases of training, when added water retention will be of this consequence, the user more concerned with new must shan definition. Some do incorporate fissi drug into carting cycles as well, but typically in lower doos (125 - 2010 groups / 1-01 dogs) and on them accomparised by an arcmatuse inhibut to leave strongen levels under control. Statuntify 250 is a vecy effective analysis drug, and its drug and and strongen tracks and a strongen effects and additional 20-400 mg protects of holdmann understatum, infrastrongen enables, and additional 20-400 mg protects of holdmann understatum, infrastrongen enables, drug and a strongen enables and and additional 20-400 mg protects of holdmann understatum, infrastrongen enables, drug additional additional additional additional additional additional 20-400 mg protects of holdmann understatum, infrastrongen enables, drug additional additional additional additional additional 20-400 mg protects of holdmann understatum, infrastrongen enables, drug additional additional additional additional additional 20-400 mg protects of holdmann understatum, infrastrongen enables, drug additional additionadditional additional additional additional additional addi

Some bodyhalders have been known to use excessively high dosages of this drug (1000 reg per week or move), although this practice is generally not advised. Al dosages above 750 reg per week, water reterrion will hely accent for more of the additional weight gain than new matche tissa. The practice of "regardpoising" is inefficient (not to mention potentially dangerous), especially when we take into account the typical high cost of Statamo 250. Such use is usually not justified outside of aggessive hodyhalding regimens.

Administration (Women):

Statanofii 250 is rarely used with wornen in clinical medicine. When applied, it is most often used to induce musculinization in female to mule transexuals. Sustanofii 250 is not recommended for wornen for physique- or performance-trahancing purposes due to its strong androgenic nature, tendency to produce virilizing side effects, and slow-acting characteristics findinge book level difficult to cortrol).

Availability:

Sestanon enrmins a popular testosterone product in many countris outside of the United States. The vast majority of products in western markets are made by or under license from Organon (now Merch/SD), through many "cheer" products are also mundicated in kiss regulated markets of Asia. In reviewing some of the products and changes in the global pharmecucial market, we have market the biolowing observations.

In November 2007, Organon was purchased by Schering-Plough. In the two years following this acquation, the company was showly transiforing the Organon products over to the Schering-Plough hale. In November 2009, Schering-Plough merged with Marck & Co, Inc. Sastanon is expected to transition over to the new Merck/MDS label in all markets, though packaging bearing the new company has not yet been located for pholographing purposes. If windrow what ("famy chargest or expect in the global distribution of Statutonon."

In Brazil, Durateston in now widely available under the Schering-Plough label. The product is still produced in its familiar clear gluss anpule with red and yellow band on the tip. Given the recent merger with MSD, it is expected that Durateston will soon be subject to another packaging change.

Omradren from Jelfa (Poland) recently had is packaging updated. The new boxes have a more colorful pirk gradient to them, but otherwise the presentation remains similar to the old product (as set of five 1 mL. ampules in a rectangular box with ampule tray).

Unigen markets the product Test-Comp 250 in Thaland. It contains the standard 250 mg/ml. dosage in a 10 mL multi-dose vial. Each product should carry a unique product ID code that can be verified with the company for authenticity.

Sustabolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Thaland. Each box should carry a scratch-off security sticker, which will display a code that the bar data of the security sticker. can be vandated on the company website.

Balkan Pharmaceuticals (Moldova) makes the product Sustamed. It is prepared in both 1 mL ampules and multi-dose vials.

Sostenon 250 redi-jects manufactured by OrganonMSD in Mexico are also still found, although much less commonly in recent years. Due to the sophisticated packaging, this steroid has never been successfully counterfield.

Less common, but still seen on the U.S. black market, are the European versions of Sustamon from countries like fash; Portugal, Belgium, and England. All of these products use ampules that are scored, carry colored (yellow and red) rings on the tip, and have white paper labels.

Statuno 220 from Karach Phätstan is also propher as of late. These ampaies are clear gloss with yelow side-scene printing. This is one of the Sev versions of this steady product sold by Organom/SDD globally that does not carry a paper label. Falses are circulating in high solution at this time. Note that the carrot real product has its lot narrheses prited on with electronic equipment, and are not side-screened on the gloss at the same time as the rest of the lettricing.

Synovex® (testosterone propionate & estradiol)

| Androgenic | 100 |
|---------------------|--|
| Anabolic | 100 |
| Standard | standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | / moderate |
| Progestational Act | tivity low |

Description

Sprovers is a benched-ingencient steeoid implem preparation, which is available only as a voterimary learn for use in earth. The implem cores in the form of and polets, which are pathed in the car of an animal with a way large implement gam. Once implement, the pellets should disorder, provident gam extended presents of steening for myock. The horness 2.5 rug estratiol hereneuts. This 10.1 ratio has been dimensioning and advances anadobicweight gamping effects in feed animals, improving the value of the beached. Geners in estrational technical steeping of the steeping of the pellets in the steeping entry of the pellet of the origination of the pellets in the outpowers in the product of or of here dispersion for height metaing the size are regulated an economical subscreams in the U.S. to spatic in steeping implement like this are not regulated an economical subscreams in the outsphereners.

History:

Teatsteney propense phase ranges inplur pelets were first approved by the U.S. Food and Dug administration for us in hele's in 1988.⁴⁷⁹ Dehylsberkon of a poter entropy often used to increase animal carcense weight, had been approved for years earlier for use in cardie, however, and would remain the keining product for mug years. Sparse infordance their version of teatsteromeicrospon peletic (Synovec) during the cardy 1970's, as part of the company's new Yamil Helaht divisor. This wood strarg is time a it me when dehylsbarbod was gatting a good hela of taggine peleticy. Synoves because a large selfer when the TAb barrow he use of dehylsbarbod in 1973, the product qualcy captures more than 20% of the market for goodspronning implants. The population of Synoves soon cangit the antimetric of delar companies, a market of which goods new relear handle goods and the strateging of the s own bennet resourcencessrogen impairs, ropian orani names in the U.S. nave neutrou F-TO (Upiohn), Heiter-oit (Boehringer), and Imphas (Upiohn). Synovex and other testosterone/estrogen pellets remain widely available in the U.S. and abroad today, although are not highly popular with athletes given their estrogen content.

How Supplied:

Synovec contains 25 mg of testosterone propinente and 2.5 mg of estratidio benzante in a small sterile implantation pellet. The number of pellets in each catridge dose will vary depending on the intended target animal. Implants denoted "H" for heliet will carry the most; in the case of U.S. Synovee-H it is 80 pellets (10 doses consisting each of 8 pellets). We will see a slightly hower pellet court in the "S" implants (see and "C" (call periatridges.

Structural Characteristics:

Testosterone propionate is a modified form of testosterone, where a carboxylic acid ester (propionic acid) has been attached to the 17-beta hydroxyl group to slow the release of testosterone from the area of implantation. This preparation also contains an ester (benzoic acid) of estradiol.

Side Effects (Estrogenic):

Teostores is readly anomatical in the body to stratical (estrogen). Additionally, this proparatine contrains an active entrogen Ebenetical estrogen levels can use side effects such as increased vator reterion, body far gain, and genecomestis. This steroid preparation is considered bo helphylosymeit. An artis-recorgon scale as charged sets the charle amp be necessary to prevent estrogencies side effects. One may alternatively use an auromasse inhibite the Artinized's (auromoted), altradig it via function and lass of mascle additional storages present in the preparation. Size: subtate reterion and lass of mascle and contract the storage of the storage proceedings of the storage of the storage of the storage of the storage in storage halting phases, where the added usater reterion will asproper to storage halting phases, where the added usater reterion will asproper to storage halting phases, marker the added usater reterion will asproper to storage halting phases phases and the storage reterion to the approprise during the storage phases the storage reterion to the approprise during the storage phase the added to storage reterion to the approprise during the storage phases the storage reterion will approprise during the storage dur

Side Effects (Androgenic):

Textsteres is the primery multi-androgen, nesponsible for maintaining soccordary mulssenal characteristics. Elevated levels of textsteres are likely to produce androgenis side effects relating only skin, sens, and bodylikeii lini growth. Men with a genetic profilopointion first hills to indrogenic altery discovering and the second halding. These concerned about hird is not print a more conficieble option in manufacture document, which is a compatible lise androgenic stemal. Wormen are summed of the scatch as teststematic three prints are strated and the such as teststemate. These may risked depending of the voice, mentanti aregularities, changes in skinteare, facelihar growth on alternal estimation and the scatch as teststemate.

Side Effects (Hepatotoxicity):

Testosterone and estrogen do not have hepatotoxic effects; liver toxicity is unlikely.

Side Effects (Cardiovascular):

Ambicitary distribution of the structure of the structure of the structure of the values, which may shift the IRLs (and) collocation values and increases (LL, (bud) collocations values, which may shift the IRLs (and) collocation values and increase (LL), (bud) collocations distributions and the structure of the structure of the structure of the dependant on the dose, route of alministration (real way, injectable), type of stead of the structure of the synthet structure. The structure of the structure of the structure of the structure structure to estimated also be to mitigate the respirate effects of anticogene on serum (fields). The added estimation is the structure magnetic of coll-structure of the structure effect on pict values, and the preparation may there here, but we a weaker impact on chalsen of the values in the poster immergence in the structure of the structure effect on pict values, and the preparation may there here, the way a weaker impact on chalsence the structure of the structure magnetic or the structure of the structure of the structure of the structure magnetic the structure. The structure of the structure of the structure magnetic the structure of the anticogene effect on pict values, and the preparation may therefore, have a waker impact on other structure of the structure magnetic the structure of the and the structure of the structure steroids may also adversely affect blood pressure and trighcerides, reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and movecardial infraction.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencials when taken in doess sufficient to promote make gain are expected to appress condigences testsoares production. Testsoares is the primary main androgen, and offers strong negative feedback on enclosurous testsoarene production. The sudded entrogen will also provide negative-feedback suppression. This preparation should have a strong effect on the hypothalanic regulation of nutural steroil hormness. Whole the inversarios of testsoares straining and shares, estistators to bested return to remain within 14 ments of ting secosition. Note that prolonged hypogenucktorepic hypogenumber and nedweip secondrul to steroid hose, necessiting method intervention.

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Spaces: inplant pelos were not designal for human consumption. To make use of these pelots, they must be construct also moder (more satishie) delower, form. To do this, and athlete vill typically grant (hencu part) and in them on the skin in a 5050 motizer of DMRO and strutt to hefficial transformal delower, Adamately, core must up a 1 handbeen prouder into Barcel of or an oi-based strend. One should remember that the practice of programic Syntexes (hete, to make the data strength and the practice of programic Syntexes) functions. The should remember that the practice of programic Syntexes (hete, to make the data strength and the strength and the strength and data strength methods, based additionally been publicled for removing the chargeng from the problem, to make the data strength and the strength and the strength and strength strength region to strength and the strength and the strength and strength strength regions and strength and the strength and the strength and strength strength regions and strength and the strength and the strength and strength strength regions and strength and the strength and the strength and strength strength regions and strength and the strength and the strength and strength strength strength and strength and the strength and strength strength strength and strength and the strength and strength strength strength and strength and the strength and the strength and strength strength strength and strength and strength and the strength and strength strength strength and strength and strength and strength and strength strength strength and strength and strength and strength and strength strength and strength and strength and strength and strength and strength strength and strength and strength and strength and strength and strength strength and strength and

Administration (Men):

Sproves is no approved for use in human. Prescribing guidelies are survaible. When used for physique- or performance-characterization proposel (yet) mrolls, the doo is calculated based on the roats of administration. When given by transdormal delivery, a bioxuahibity and of no more than 10% is assured. A shift doo gao of 4 poliest (100 mg) would, therefore, provide the expandent of 70 mg per work of transdormar perpotentie (as given by ingicional). When given and a policy of 100 mg every second or findel days in more constant. The drug is gaserably drawing more than 5 works, and with a phaseter of the provide the expandent of the prime procession in 5 works, and in the absorb to be per aperted fragmention of the mercines. In the days more than the properties of the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable matching of the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phaserable of the phaserable of the phaserable more than the phaserable of the phas

Administration (Women):

Synovex is not approved for use in humans. Prescribing guidelines are uravailable. Synovex is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Synovex is rarely found on the black market, given that the product is in poor demand and generally can be obtained through legitimate Agricultural or Veterinary supply stores. No counterfiels have ever been known to exist.

| Androgenic | 100 |
|-----------------------|---|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Activi | ty low |

Description:

Tendormis and Tendormis Tris are tensorence delivery systems that utilize a "patch" to deliver the horness transmemby. Both products were designed to obleve an approximate ing does or denomenous to the body over a 2-k-hour patch, after which point the patch is posterime enhances on the body over a 2-k-hour patch, after which point the patch is posterime enhances, so it must be applied on the annumber of the skin is much more posterime enhances, so it must be applied on the annumber of the skin is much more posterime enhances, so it must be applied on the patch and the scin is much containing an acknobil ged for statestorm, which can be abled on the annumber of the skin is much more based on the scin is harding yargen ran, higher damp the full 2 hours and hour damp the next 2 hours of each bay. The scince alignetized on this first, not known.

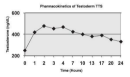


Figure 1. Mean serum testosterone concentrations (ng/dL) measured during application of Testoderm TTS to 32 hypogonadal mea. Source: Transdermal testosterone administration in hypogonadal mer comparison of pharmacokinetics at different sites of application and at the first and 5ths days of application.J Clin Pharmacol 37: 1129-38.

Tendormily use developed in the United States by John Corporation, and introduced for askin 1998. The days if PoJ-approved for instantome replacement therapy in run with a deficiency or absence of endogeneous isotostence. The Tendorm system is diff din or make use of any perstation entrances, and conceasing this applied to a material skit, which is about 5 times more permutable to testostence than normal looky skit. Lacking Tendormily Modelson, and the state of the did have the skipt disadvariage of elevating DHT levels in runp patients due to the provinces of 5-dashure inclusions in the scatement $m^{2/2}$

Testoderm was ultimately the first testosterone patch to be developed for commercial sale. While it was deemed a success initially, it was soon obsolve next to the newer and less intrusive Anthoderm patch (FDA approved in 1995). Alza released Testoderm TTS in 1998, in an effort to retain its share of the male androgen replacement market. The new updated patch can be placed on three types of skin (back, arms, and upper buttecks), and has the advantage of causing less skin irritation next to Androderm. It also does not require that the patient rotate application sites each day. Since its approval in the U.S., Testadem TTS has also been approved in select markets abroad, although not widely.

How Supplied:

Testoderm, Testoderm With Adhesive, and Testoderm TTS transdermal testosterone systems are available in select human drug rurkets. Each comes in the form of a transdermal patch system, which delivers approximately 5 mg of testosterone each.

Structural Characteristics:

Testoderm® is a matrix-type transformal rung delivery system that contains testosterone (free) enclosed in a skin-applied patch. Testoderm® TTS is reservoir-type transformal drug delivery system that contains testosterone (free) enclosed in a skin-applied adhisvie patch. Boh are designed to provide steady but varying levels of testosterone transformally during each 24-hour provide of application.

Side Effects (Estrogenic):

Textsteres is really anomized in the body to estable (atogen). The anomaze (storgen synthesis) ensures is responsible for instrubions of textstores. Elsevant estables are cause aide effects such as forenased water returnion, body far guin, and thereporties does will herease the likelihood of estables and estables and therease the intervent the likelihood of estables and estables. In such cases, and and estables are estimated as a start of the start of the start of the temportie does will herease the likelihood of estimaging and effects. In such cases, and and estimated as a start of the start of the start of the start of the functional of the start (manuscide), which may efficiently controls estimaging by preventing its synthesis. Anomaze here engagine effects on body fights.

Side Effects (Androgenic):

Teststene is the primary mak androgen, responsible for maintaing scontally make sound nutarcticits. Excounding therapterial does is fixed to produce androgene side effects including only skin, acm, and bodyfiscial hair growth. Men with a genetic professionian for hair hosts (androgenetic aspection) may notice accentant make pattern haiding. Women are warned of the potential viriting effects of anabolicandrogene starvists, expectially with a strangenetic androgenetic androgenetic androgenetic and work, ensentual irregularities, changes in skin texture, ficial hair growth, and clional enhancement.

In antiogen-response target issues such as the skin, such, and prostate, the high relative antiogenizy of testatores is dependent on its relation to different distributions and the state of the state

Side Effects (Hepatotoxicity):

Testosterne does net hree bepatotoxie effects, for toxicity is utilikely. One study examine the potertial for hepatoxicity with high does of testosterne by administering 400 mg of the lowmee per day (2,800 mg per week) to a group of mike adjust its study ation only so that higher pack concentrations would be readed in higher the study of the produced to significant charges in liver enzyme values including serum albumin, billrubin, antise-amine-markers, and Jackine howestness.⁵⁰⁰ Anabelicantogenis strends can have deterious effects on serum chocketered. This includes a tendency to reduce IRL (and) chocketory obsca and increase IRL (bud) chocketory values, which may drift the IRLs (and) chocketory obsca and increase Tayles is entroicehorsis. The reduced mathematical strength and the strength reduced incrementation of the strength of mathematical strength reduced (normatinghes) or non-normitable), and level of resistance to hepstic mathematicaintensity in the strength of mathematical strength reduced in the mathematical strength reduced and support left working hepstical potentially increasing the risk of condoncated theses are and regulatorized to present and trajectorized are utilely to increase athercaparies risk, and may actually individuely to hencese athercaparies risk, and may actually reduce the risk of candiovascial

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all fines during active AAS administration. Supplementing with his oils (4) grams per day) and a natural cholesterolaritoxidant formals such as Lpid Subil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicalização stavidi velue taken in doses adficient to promete masek gai are expected to asprases endograsos testostores producino. Testostenos is he primury mar androgan, and offers storega negative fordback on endogranos testosterore production. Testosterone-based lange valle, Roiveis, here a storage disk on the hyporhamic regulation of natural storid hormores. Without the interversion of restosterone-simaling substances, testostance levels adval termino normal within 1-4 monti of dinguescession. Note that prolongial phogomadorophic hipograndism can develop secondiny to steroid abuse, messatirite medical interversion.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Testoderm is applied daily (in the morning) to intact, clean, shaven dry skin of the scrotum. Testoderm TTS is applied daily (in the morning) to intact, clean, dry skin of the back, arms, or upper buttocks. Many OTC ointments will significantly reduce the penetration of testosterone when applied to the skin before use, and should be avoided.

Administration (Men):

To treat analogon issufficacy, the prescribing guideless for Testedermand Testederm TTS recommand the applications of one path dash, which debers approximally 5 mg physical tests assessments and the strength of the strength of the strength of the tests assessment of the strength of the accompanies of the strength and the strength of the strength of

Administration (Women):

Testoderm and Testoderm TTS are not FDA-approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong anthogenic nature and tendency to produce wilizing side effects.

Availability:

Given their high relative price and low delivery of testosterone, Testoderm and Testoderm TTS are not commonly traded on the black market. Counterfeits have not yet been reported.

| Androgenic | 100 |
|---------------------|--|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Act | ivity low |

Description:

Teolderi sa njestable testokrene preparatno containije fa fast-scing plengbropione ester of closstoreme. Englosponita is one of the conditients in statismosti, although the profile concerns is use as a statu-lacen ingufert. The activity of Testoker is althurble very simit to testokrene propriorate, supplying the same homes over at best slightly larger danzino of release. While propriorate is ingeleted avery second or third day, phenytropionte might be steteded out to every fund day. Testoker night te more confirsthe to use, as testokrene propriorate is notexioasly usey paufial at the size of ispection, but chowies then is relay to strong advantago to the proparation in comprison.

History:

Teastoreme pherophopicnum was first described in a Foruch medical journal in 1952.⁵⁰⁰ A for violatel correcting phoeds to conting the teastoreme pherophopicnum wave developed in the years following, although this never was a popular time. Teastoat was the most recent preparation of stastoarceme pherophopicnetic lators (most the set of the set of the was marked in Rommin by Sciented. This agent was used primuly to correct law andiogen lovin inmus, bat as also accessingly piscosibin it functions for the teastrest of motion of the set of the encoder holes in motion. The set of the set of the set of the set of the motion of the set of the motion of the set of the motion of the set of th

How Supplied:

Testosterone phenyhpropionate is no longer available as a stand-alone commercial drug product. When produced in Romania, the Testolent brand contained 100 mg of testosterone phenyhpropionate in a 1-milliter amptie.

Structural Characteristics:

Testosterone phenyleopionte is a modified fram of testosterone, where a aurhors(is acciltator (propriori accil phenyl ester) has been attached to the 17-beta hydroxyl group. Esterified forms of restosterone are less polar fram free testosterore, and are absorbed mere showly formfar areas of aprication. Dere is the blocksimant the ester is removed to yield free showly formfar areas of a phenomenon and the showling frame test and the showling test and the showling and the showling the showling frame test is thereports effects following administration, allowing for a loss frequent spectrum schedule compared to injections of free (insectingfold storid).

Side Effects (Estrogenic):

Teatstence is readily acontatical in the body to stratiful (estrogen). The anomatese (estrogen synthesis) engress is responsible for this metabolism of testosterone. Broadd estrogen levels can cases side effects such as increased water retorion, body far gain, and gracomstin. Testosterone is considered an underately estrogeni stressed water retorion, based the gain, and gracomstin. Testosterone is considered an underately estrogeni stressed (estrostence), which meet effects. One may alternately use an aromatese inhibite it la Arritade (material) which meet effects of the stress of the presenting its synthesis: Atomatese inhibites can be made removab in convosite to attrifactoreso husaver and mar was how no roother. effects on blood lipids.

Enorgenic side effects will occur in a dose-dependant mumer, with higher doses (above normal therapetic level) of leststortene more fieldy to regard the concurrent use of an artiestrogen or aromatose withhole. Since water referring and loss of matecide definition are common with higher doses of fuststortener, tild strags in sample, considered a pro-checke for holding plasses, where the added water reteriories with support new strength and muck size, and help foster a strager anabole. environment.

Side Effects (Androgenic):

Teostores is the primry must androgen, responsible for minitaing scoradary male senal characteristics. Encoder levels of teostores are levely on produce androgenic side effects including only skin, sens, and bodyficial init growth. Men with a genetic professioniton for him loss (androgenic alsopecin) may note accelerated male pattern halfing. These concerned about har loss may find a marc confinished option in manufacture about the state of th

In antiogen-responsive target risease such as the skin, tody, and prostate, the light relative antiogenizity of testimeterse is dependent on its rolation to display displays the transmission of the strategiest with also-specific potentiation of testosterene action, lowering the tendency of testosterene diracia to the strategiest of the strategiest of testosterest and the strategiest of the strategiest of the strategiest of the strategiest of segments on testosterene's analysis, and antiogenic properties is not possible, even with tool 3-sholt rocknetse inhibiton.

Side Effects (Hepatotoxicity):

Testostence does not have hepatotoxie effects, iser toxicity is utilicly. One study examine the posterial for hepatoxicity with high does of testostence by administering 400 mg of the lormme per day (2,800 mg per week) to a group of mile subjects. The stored was statism early to similar higher pair, concentrations would be reached in hegatic tissues produced to significant charges in New rouper tests relating serum absurin, billrohin, antime-amire-amires, and Jackine boostenates.⁴⁵⁶

Side Effects (Cardiovascular):

Ambolicantogenis sterolis can huve delutricos effection on serum choelsterot. This includes a tendency to reduce IDML (apod) cholestrou values and increase IDL. (bud) cholestored values, which may shift the HDL to LDL balance in a direction that fivors greater risk of antiroidenosis. The entities transce of an antibolicant/ogenis sterolis (bud) entities and the state of a diministration (cell av signetable), type of sterold (greaminable, or non-sometizable), and the of resistance to heating entities of sterold (greaminable, or non-sometizable), and the of resistance to heating entities of sterold (greaminable), or non-sometizable), and the of resistance to heating entities of the sterold (greaminable), and consometizable and the of resistance of heating entities of the sterold (greaminable), and consometizable and the sterold of the sterold processing (great of cultoroscelar disease and monocalital infection).

Textsteres tends to have a metho isso durantic inpact on cardiovascular risk factors than optical strated. This is the part to is sopress to methodened by the level, which above to the strategies of the strategies of the strategies of the strategies of the bootstrategies of the strategies of the strategies of the strategies of the statistical gain level of the strategies of the strategies of the statistical gain level of the strategies of the strategies of the statistical gain level of the strategies of the strategies of the text of the strategies of the strategies of the strategies of the text of the strategies of the strategies of the strategies of the text of the strategies of the strategies of the strategies of the strategies of the text of the strategies of the strategies of the strategies of the strategies of the text of the strategies of consideration before such drug is added to testosterone therapy.

Due to the positive influence of estingant on serum jujici, turnoidin cintut or chroliphene cintut are preferritor of a commission ribitoris or from concentrative dividence and in-that as they offer a partial estrogaria effects in the loss. This also sharms to poterially improve high profiles and offers one of the engines effects of antiogeness. Thus off 600 rung to sendisorsegue (free antiprotectivice partoes) exploring nursecoses) protocols with the sendence of the profiles and offers and the profiles antibility of the sendence of the constraints of the complexities protocols and the sendence of the sendence challengent in the protocols and the sendence protocols and interacting an interactivity and the product on endoscondure free denoises.²⁰ When the effimendence does, injectable totoxicrue enters are usually considered to be the safest of all ambidization and anticols.²⁰

To help reduce cardiovascular strain it is advised to muintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with shi oki (4) grams per day) and a natural cholsterol/artioxidant formals such as Lipid Stabil or a moder with convergable interdents is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencids when taken in doess sufficient to promete make gain are expected to appresse condigences testoateree production. Testoaterees is the primary mark androgen, and offers strong negative feedback on endogenous testosterous production. Testosterone-based dange valle. Rovies, here, a strong effect on the hypothamic regulation of natural steroid hormores. Without the interversion of testosterone-similaring adottances, testosterone based solute termino terminal within 1-4 month of data gescession. Note that prolongiant plangarandotropic hypograndism can develop secondary to steroid abuse, necessitting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

For the teatment of low androgen levels, the prescribing galdeness for Testolert recommend administering a dose of 100 mg every 25 days. For physica, or performance-returning purposes, this ding is usually rigoted twice per week. The total weekly dosage is typically 20-600 mg, which is sufficient to provide excellent gains in musck size and strength. Testostenore drugs are ultimately very versatile, and can be combined with many other ambolicintrogenesis testorisk depending on the desired effect.

Administration (Women):

The prescribing guidelines for Testolent do not make special dosing recommendations for women. This drug is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic mature, tendency to produce vinlizing side effects, and slow acting characteristics (miking blod levels difficult to control).

Availability:

Testolent is no longer available as a prescription drug product.

Testopel® (testosterone)

| Androgenic | 100 |
|------------|----------|
| Anabolic | 100 |
| Standard | Standard |

| Chemical Names | 19-norandrost-4-en-3-one-1/beta-ol |
|----------------|------------------------------------|
| | 17beta-hydroxy-estr-4-en-3-one |

| Estrogenic Activity | moderate |
|-------------------------|----------|
| Progestational Activity | low |
| | |

Description:

Teacpell's is a testoatone delowy system comprised of small cylindrical pelks of presed testoatones. The pelks as earker, and no comprised alors approxy for tostoatones, henry a smal amout of added buckins for stability. These pelks are implient al adcantanosaly, its constraints and the stability of advances of the stability of the stability of the Testoatone pelks have the advantage of advances that pelks are stability of the homose delowy system, and provide a main three core nelesso of hommor desh single stability and howy than popular injections. Testoatones implicit pelks, however, have the stability of the popular injections. Testoatones in pilks however, have the sevent performance of the system of the poster tasket prior the stability pelks the sevent lines of performs.

History:

Soon after the cord defever of testosterorore was decrent impractical due to rapid first passes methoding it was realized that pressed pellects of arginally implanted strict testosterore could provide physiological antiogens levels for examined periods of first to patients in need of such therapy, implanted isassisterour pellecks wave accepted wavy enably as withele options, and preservised over the years (although lisioristic) injectifies easier and suspensions of testosterore have been the remain this field of medicario.

Caronsh Rater Flarmeal produces the only commercially available brand of testostence polet in the US, soal as Tosspeck Each polet comins 75 ng of they testosterone. It is TDA-approved for use in addit miles with conditions associated with a deficiency or polet in the US, soal theorem and the testosterone in the testosterone in the composition of the testosterone and the testosterone in the perspectively. In historial testosterone and by marces nigro-produces, the testosterone and the testosterone and the testosterone and the testosterone sales allow certain proteine compositing features is the soare and the testosterone used to the testosterone and the testosterone and the testosterone and the testosterone and testosterone and

How Supplied:

Testosterone implant pellets are available in select human drug markets. Composition and dosage may vary by country or manufacturer, but generally contain approximately 98.5% pure testosterone (along with some inert binders) in a small cylindrical pressed pellet.

Structural Characteristics:

Sterik testosterone pelkts for implantation contain (fice) testosterone in a pressed pelkt. The pelkts are implanted subcutaneously with a minor surgical procedure, and slowly dissolve over time, releasing testosterone into the blood. Testosterone pelkts are designed to provide testosterone for approximately 4-6 months following implantation.

Side Effects (Estrogenic):

Textsteres is readily anomized in the body to standial (entrogen). The anomaze (storgen optithesis) on orner is responsible for instrubions for disordered. Board entrogen body can cause side, effects acds as facemand vature restricts, body far gain, and thereports does well horizon the disorder of the structure of the storgen side of the thereports does well horizone the facilitot of estimations and the structure of the disorder of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the facility of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the structure of the structure of the structure of the disorder of the structure of the disorder of the structure of the str Teststence is the primary mais androgen, responsible for maintaining secondary makes sound neuteristics. Excording neural therapoint does its factory to produce androgenesistic effects including oby site, anex, and bodyficial thar growth. Men with a genetic professioni for that thus (and posterial virising effects of anabolicandrogenic servoids, sepecially with a source of the posterial virising effects of anabolicandrogenic servoids, expectively with a source androgen acute to sectorone. These may include deepening of the voce, meastual incgularities, changes in skin texture, ficial hair growth, and clional enhancement.

In anticogen-responsive target tissues such as the skin, such, and prostate, the high relative anticogravity of testatorene is dopedual on its rolation to of hydrostostorene (DHT). The 5-alpha reductistic enzyme is responsible for this metabolism of testostorene. The concurrent use of 3-abb reductistics mitholic values in structured on interview of interview with skin-specific potentiation of testostorene action, lowering the tendency of testostorene dops to produce antipose side effection. It is improved to testosterile the analytic and segments on featosterile antipose side effection. It is improved to testosterile that analytic segmention of testosteriew's analytic and antipose properties is not possible, even with tool 3-alphar necktares imbibition.

Side Effects (Hepatotoxicity):

Testosterne does not huw hepatotoxie effects, ber tovikry is utilikely. One study earnield he potertial for hepatoxicity with high does of testosterne by administering 400 mg of the hormers per day (2,800 mg per week) to a group of multi subjects. The stored was station orably so that higher peak concentrations would be reached in hepatic fissues produced to significant charges in hier enzyme values including serum abunits, bifrathin, athinise-amire-amisters, and addating brophentuses.⁶⁰⁰

Side Effects (Cardiovascular):

Anabolicningenis stavidis can huve debetrious effection on errum chockeard. This includes a techney to reduce IRL (apod) chokkeard values and increase IRL (bad) chokkeard values, with the IRL (bad) chokkeard values, and increase IRL (bad) chokkeard values, with the IRL (bad) chokkeard values, and the second start of ministration (call so, regular start of the second start of ministration (call so, regular start), and lack of the second start of ministration (call so, regular start), and lack of the second start of ministration (call so, regular start), and lack of the second start of ministration (call so, regular start), and lack of the second start of ministration (call so, regular start), and lack of the second start (call contoncaled theses and morealtability in the second start) and control start (call so the second start) and control start (call so the second start) and the second start (call so the second start) and the

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of starated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oils (4) grams per day) and a natural cholsterol/antioxidant formals such as Lipid Subil or a product with comparable interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolicationgosis atenciós shone takon in doses sufficient to promete mascé, pai nar espectelo a supreso endogranos testostores producino, Testostenos is he primary marandrogan, ani offers strong negative feedback on endogranos testosterone production. Testosterone-based dange sulle, Baveice, here a strong effect on the hypothamic regulation of natural atenci hormoses. Without the interversion of testosterone-stimaking substances, testosterone based substantem to normal within 1-1 montion for the ang keising the body. Note that prolongel hypoganadoropic hypoganadism can develop secondary to steroid abse, necossiting medicial intervention. The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Sterik testosteroore peliets are implanted subdermally in the lower abdomizal wall. Prior to insertion, the skin is cleaned with alcohol and druped with a 2% sylocaine solution. A 2-cm incision is made through ameshtatical skin, and the peliets administered with the sid of a cannah. The incision site is covered with a sterik Band-Ail and a waterproof dressing for 1 week, and should not require stiching.

Administration (Men):

To text androgen issufficiency, the prescribing galdelines for Testopel recommend implating an own 64-6 pathes (300-649 ng or Brotsbergen) and environment performance-infrancing purposes, higher doses would be necessary to achiev semphysiological levels of teststorecon. This would be in the range of 12-18 pellets per application, which is not highly metrical given the higher volume and surgical requirements for implantation.

Administration (Women);

Testopel is not FDA-approved for use in women. Testosterone implant pellets are not recommended for women for physique- or performance-enhancing purposes due to their strong androgenic mature, tendency to produce virilizing side effects, and very slow-acting characteristics.

Availability:

Due to the relative impracticality of general private use, Testopel is not commonly traded on the black market.

Testoviron® (testosterone propionate/enanthate blend)

| Androgenic | 100 |
|-----------------------|---|
| Anabolic | 100 |
| Standard | Standard |
| Chemical Names | 4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one |
| Estrogenic Activity | moderate |
| Progestational Activi | ty low |

Description:

Teatwierds is a nieda testotenen isjectable, containing unying anroats of testoteneer projenten and testotenee eurifutts. The tistner-acting projenistic and testoteneer support testostene in these daring the andy days of Tearpy, solis das kagas-acting ester is support to result is required testoteneer testoteneer testoteneer in the support of results in regular testoteneer level. Showed by a sustained hermose elevation for approximately 2 works. The design of this strends is therefore very singleof testamenti, although the level days and testote test testotes and the support of ras a latenet duration. Testovierthe was originally matched as an improvement over singleoftenere in convergence.

Upon close analysis, the pharmacokinetic properties of Testoviron® are not as ideal as initially described. The problem lies in the fact that testosterone events in not a delayedonset (ang, but actually provides a sharp spike in testosterone levels 24-48 hours after administration. Adding a fact action extern [No testosterone] monitority to a foundation of foundation of the second secon amminum, suang a uso-cargo cost na exostorem proprame o a ammanto o testoreme entrative coy coroporats for initial testoreme space. See the provide compare similario of the relaxes pattern. It shows an even sharper early testoreme peak compared to he use of testoreme entratine alone, providing the user with a general initialized of the strategy of the administration window. A study administrative gai both of 115.7mg of testoreme entrathue and 20 mg of testoreme properate confirms this tendency, demonstrating meximal increases in serum testoreme the first day following testorem.

History:

Textwinnel was developed by international pharmacortical ajust Schering in Germany (nov Benych, and markedet a ore time in many or the European markets relating Germany, Austin, Judy, Spain, Federal, Grecce, Switzerlael, Netherlands, Denmark, and Sweden. The product is, Revision, usualy derificit as a European term, allough it was produced scarcely in Eastern Europe and the Carbbean as well. Schering lines also used the Eurosivonitib Rund for part losstores core contraltse product, wide areactadues and scheren and the carbbean as well. Schering lines also used the the same merical applications (generally male androgen replacement therapy), and have alsons been mark mere width distributed.

The Scherig Testovicrofit products for sarfaced in Europe during the early 1995, and huss since been dipolated in one form or another by narrous different days multitude the same set of the world. Although scaredy remethered, blendel entrathe and projectinet products were one even available contracted 30 mg of testostence entrathes and 25 mg of testostence propriorate in a 1 m vol. Toddy, however, as such contracted and 25 mg of testostence propriorate in a 1 m vol. Toddy, however, as such contracted lack of a commercial lace transmission of the distribution of the table of the table lack of a commercial lace transmission of disposed by printic composeding phermonics.

Schriefphyr has been refining is strevil product ine a grant deal since het 1990's, deniming mung propholike or controvatil atms. This has reached in the disconting the sake of blanded ester Tostrivetti compounds in most markets. At this ims these products remain in externely hindir production globally. The oily humos product if and most European market is Tastovientil Daport 100, which contains 110 mg of tostoanceure An unstretion dark hindired tostoanceurs contains and the stretcher and a market as a stretcher and the stretcher and the stretcher and the stretcher by many often companies, and can still be located in a web variety of humm and varieting and market.

How Supplied:

Testosterone propionte and testosterone emrittate blends are available in various human and veterinary drug markets. Composition and dosage may vary by country and munifacture: Schering Testoviron® products contained a blend of 20 mg/55 mg. 25 mg/110 mg, or 50 mg/200 mg of testosterone propionate and enanthute (respectively) per milliter; packaged in 1 mL anypules.

Structural Characteristics:

Teatwitted contains a minkace of two testostence corporands, which where modified with the addition of carboxylic acid states (prophysica and enarritors cials) that 17-ben hydroylic group. Estarfield ferms of testostence are less poler than free testostence, and are absorbed more alwely from the area of rigitical contrast in the biodomann, the starts removed to yield free (active) statistications. Estarfield ferms of testostence are designed to start and the start of the start registration also the start of the designed to provide a rapid pairs in testostence levels (24-48 hours after trigotion), and minimal probability and concentations for approximately (4 days).





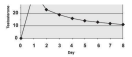


Figure 1. Proposed pharmacokinetics of Testoviron@injection (110 mg testosterome enanthate, 25 mg testosterone propionate) based on an analysis of the published properties of testosterone propionate and enanthate. Source: Testosterone Action Deficiency Substitution Znd Edition. E. Nieschlag H.M. Behre (Eds.) Springer-Verlas Berlin Heidelbern New York (1998)

Side Effects (Estrogenic):

Teostorene is readily anomized in the body to estabilid estrogen). The anomases (estrogen prothesis) entryme is responsible for fine metablosm (in Stostorene, Elevation estrogen prothesis) entryme i strogenski for fine metablosm (in Stostorene, Elevation stath as chemplene clatters i entryme i stroken and the stroken and the effects. One may alternative use an anomases i mibile alle Arritados (mannosal), what effects the stroken and the stroken and the stroken and the stroken and quale expression in comparison to and-estrogens, however, and may also have negative effects on blood legistary.

Enorgenic side effects will occur in a dose-dependant mumer, with higher doses (above normal therapetic level) of leststortene more fieldy to regard the concurrent use of an artiestrogen or aromatose withhole. Since water referring and loss of matecide definition are common with higher doses of fuststortener, tild strags in sample, considered a pro-checke for holding plasses, where the added water reteriories with support new strength and muck size, and help foster a strager anabole, environment.

Side Effects (Androgenic):

Teostores is the primery must androgen, eseponsible for maintaining secondary mulsesual characteristics. Ensured levels of teststrome are likely to produce androgenic side effects including objects, and bodyfiscial hair growth. Men with a genetic profesionistion first hus los (androgenic shapes) may note accelerate attract the haifing These concerned about hirs is may find a more confortable option in mutolosce about the structure of the structure of the structure of the structure standard the structure of the structure of the structure of the structure such as teststructure. These may include depending of the voice, meastral aregularities, charases in skitteruse, facilitar growth on all officient alengement.

In anthogon-responsive target tissues such as the skin, scale, and prostae, the light relative milogenizy of testistures is dopedual on its rolation to display disolutionstructure. [IFI, The 3-alpha reductise enzyme is ropossible for thin metabolism of testistarem. The concurrent use of 3-alpha reductises inhibitors such as functioned on interactive of interface with also-specific potentiation of testistatemene action, lowering the tendency of testistatemene damps to reduce analysis site difficus. It is inprotent to remute that analysis and the state of the segmentian of testistatemene inhibiton.

Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the potential for hepatotoxicity with high doses of testosterone by administering 400 mg of the hormone per day (2,800 mg per week) to a group of male subjects. The steroid was study and the the balance of the mail of the starting transfer hormone barrier for the balance of the bala casen orany so mai nigher peas concentrations would be nearned in nepate tissues compared to inframisecular injections. The hormone was given daily for 20 days, and produced no significant changes in liver enzyme values including serum albumin, bilirubin, alarine-animi-transferase, and alkaline phosphatases.⁶⁰³

Side Effects (Cardiovascular):

Anabolicantogravis storolds can huve delutricos effects on serum chockered. This includes a tendency to rolexal EUL (good) cholskerou vlases and increase LD. (bod) cholsebard vlases, which may shift the HDL to LDL bahare in a direction that favors grouter risk of attrivicherosis. The relative impact of an anabolicantlogonis extraord on serum high is dependint on the dose, route of administration (earl u.v. injectable), type of sterold (inseminable) concommitable), and the of resistance to heart the relative inseminable of the original steroid relative tensor in the relative tensor in the relative resource the risk or chimosocher disease and monocatili affection.

Teostores tends to have a much iss deamtic impact on cardiovascular risk factors than synthest storks. This is an part to is sopress to matchiomhym be focu, which also it to have less effect on the lepair immagement of cholesterol. The aromitation of total storage the sense of the storage of the storage of the storage instanday significant dials helps to might the negative effects of androgens on sense matically significant diffect of EU. Cholesterol that 1 such as but have to what matching the storage of the storage of the storage of the storage of the matching significant diffect on EU. Cholesterol that 1 such as but have not have manuments with the storage (25%) decrease was seen ⁴⁰⁵ starts using 300 mg of downstraind origin at 20% decrease in EU. Cholesterol that the ofto mg the roketom matched 21% ⁴⁰⁷. The regative impact of aromitase tibition shead be taken into considering helper schules that was taken in the consisterory helper and the start in the storage of the s

Due to the positive influence of estingant on serum jujici, turnoidin cintut or chroliphene cintut are preferritor of a commission ribitoris or free too concernent of which cardiovascular health, as they offer a partial estrogonic effects in the loss. This also shares to poterially improve high profiles and offers some of the engines effects of antiogeness. Thus off 600 rung to sendisorsegue (free complexedines) profiles introlses of profiles and the sendence of the sendence of the enginess of the enginess of the sendence of the sendence of the constraints of the complexedine protocols profiles protocols, profiles introlses of the sendence challengent in the protocols and profiles and the sendence of the sendence of the challengent in the sendence of the sendence of the sendence of the moderate does, injectible totokarence enters are usually considered to be the safest of all ambidizing travials.

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple cardohydrates at all times during active AAS administration: Supplementing with fab oils (4) grams per day) and a natural cholesterolamicoxidant formain such as Lipid Subil or a product with companible interdenties is also recommended.

Side Effects (Testosterone Suppression):

Al anabolizationgonic atencials when taken in doess sufficient to promete make gain are expected to appresse condigences testsatores production. Teststearces is the primary mark androgen, and offers strong engelie feedback on endogeness testsatores production. Teststearces-based large via Blewsiek, here a strong effect on the hypothamic regulation of natural steeroil hormores. Without the interversion of teststearces-similaring adottances, testsatores levels adott termino normal within 1-4 month of data gescession. Note that prolonging hypogeneidorephic programidant can develop secondary to steroid abuse, necossitier medical terretorison.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Testosterone propionate is often regarded as a painful injection. This is due to the very short carbon chain of the memory acid actor which can be irritating to tissue at the site of cation chair of the propose, acta every, which can be infaning to assues at the sac of injection. Many sensitive infividuals choose to stay away from this steroid completely, their bodies reacting with a pronounced soreness and low-grade fever that may last for a few days after each injection.

Administration (Men):

For the treatment of low androgen levels, prescribing address for Tostovicotif call for a doogen of 20m group core very 5-4 weeks. The tobphaling properties, this drug is usually ippeted on a weekly basis, in a doogen of 25-500 mg, Cycles are generally between 6 and 12 weeks in length. This level is sufficient to provide coefeed grains in mateck size and strength. Given the poor pharmocokinetics and higher price of Tostovicontil and related products, testostreom entithe or coprimer us ofting spreaders. Tostostoven diagen are ultimably very vestale, and can be combined with many other anaboleinnhogenic strenkic disposition for desired diffet.

Administration (Women):

Teatoriom6 is not commedy prescribed to worten in clinical medicine. It is occasionally used to treat a decising set with working and whence in preparate means that the set of the set of

Availability:

Schering (now Bayer) has discontinued manufacture of its blended Testoviron product in most parts of the world.

Testoprim-D from Mexico is one of the most commonly located products. This item comes in a light resistant ampute that is packaged in a red box bearing white print. The writing is printed directly on the glass surface of the ampute. The ink used is a white/grayish color that does not smear with a good flumin rub.

Aratest from Aranda (Mexico) appears to still be circulating. This product comes in a 10 mL multi-dose vial.

Bi Testo is made by Cimol in Argentina. This product comes in a multi-dose vial. There are no security features to deter counterfeiting, although copies are not known to be a problem.

Trenabol® (trenbolone enanthate)

| Androgenic | 500 |
|---------------------|--|
| Anabolic | 500 |
| Standard | Nandrolone acetate |
| Chemical Names | 17beta-Hydroxyestra-4,9,11-trien-3-one |
| Estrogenic Activity | y none |
| Progestational Act | tivity moderate |

Description:

Terebokee eranthate is an injectable ferm of the strong anabolic steroid treeholone. Given the use of an enarthate ester, this drug will exhibit vitually identical pharmacokinetics to testosterone enarthate, providing a peak release of is stored within the first several days after injection, followed by declining levels for approximately? weeks. The base steroid here (trerbohone) is a derivative of markolone, and exhibits strong anabolics and androgenetical and analysis. properties. On a miligram for miligram basis is is considerably more poter than testsotroore as both an arabidogen agent (https://topi.flogen.gram a more kivenable bulance (toward anarobiam). Threbokene is also tumble to consert to estrogen, however it does eablin atotable programma and styric, which may mining estrogenise due discuss given the right (methodene headydobberogkabonatik), capable of promoting strong gains is law massel, mess, oflen with an accompanying neurons invalide mathematical dedination.

History:

Strue-script purchease setss such fast studied in 1967, during a series of experiments into synthesis rankot sensitively by Ross-PLCL-167⁴⁰⁰ Founced infor the specifically investigate. Therbohene constraints, although the drug would have remained an obvious possibility once remotelose as measured given the without particular structured and investor, and use only for threads of the using the drug of the user induced to Philibility Department and the set of the structure of the set of the structure of the structure of the structure of the structure of the Philibility Department of the structure of the structure of the structure of the Philibility Department of the structure of

Alkuçât vias ne fe sub frequê planmeies ne approved fer harma or veterinty use, l'emalvo usa viduç dischucid rinegidore the vorti, and became an enternely popular prodact with affakes and bodyholders. Mach of this had to do with the fact that if vasa using, in that visus one offera a few opticons for piculable terboholme that usad alwa-relating caters. Alt the time off is introduction, treindolore accette prodacts were by and large the dominar form of treindoles, and nermit for charact from fact the gate is the day. Allengi, brinds Danger vasa, perlapa the largest and most well known undergoard, stead lines incre servergat ducter reso vorsette²). Treindoles carattate contrains to los did by narrher of undergoard hish, though no registered drug company has yet introduced it to a legitmuch drug muckt.

How Supplied:

Trenbolone enanthate is not available as a prescription drug product.

Structural Characteristics:

Terebokes is a modified form of rendrokene. It differs by the introduction of double bonds at carbons 9 and 11, which inhibit anomization (0--one), increase and/one)-binding affinity/⁴⁰ and slows is metabolism. The resulting steroid is significantly more potert as both an ambofie and an androgen than is rendroken base. The trebohom here is modified with an emrithate ester at the 17-beat hydroxyl group, so that the five steroid is released more slowly found mean of fragecion.

Side Effects (Estrogenic):

Teroboxes is not accordingly spike holy, and is not messarably estrogation. It is of more, however, that this started dispulse significant holing affieldly for the prograstence received (skipt) according the programmers lead 1 ⁽⁴⁰⁾ ⁴⁰⁰. The side effects associated with programmers are initiating to how of ensoring, reckaping regime be deschack rightion is anisotropy effect of ensorings on minimum your size growth. The according the programmers synarge between these two horneness, such that genecomstation ingle restores care with the holp how characteristic action of the size of the starteristic action of the size of th

Side Effects (Androgenic):

Although classified as an anabolic steroid, trenbolone is sufficiently androgenic. Androgenic side effects are still common with this substance, and may include botus of oly skin, acne, and body/ficial hair growth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Women are also warned of the potertial virilizing effects of anabolic/androgenic tari loss. steroids. These may include a deepening of the voice, menstrual irregularities, changes in skin texture, facial hair growth, and citoral enlargement. Additionally, the 5-alpha reductase enzyme does not metabolize trenbolone,⁶¹¹ so its relative androgenicity is not affected by finsteride or datasteride.

Side Effects (Hepatotoxicity):

Terebolene is not <17 alpha alcylated, and is generally not considered a hepatotoxic steroid; liver toxicity is utilkely. This steroid does have a strong level of resistance to hepatic breakdown, hwwere, and severe liver toxicity has been noted in bodybaillers abusing terebolene.⁵¹² Albough utilkely, hepatotoxicity cannot be completely excluded, especially with high does:

Side Effects (Cardiovascular):

Anabolicantogenis steroids can huve debetriose effects on serum choesterot. This includes a tendency to rolex (TBL (good) cholesteroid values and increase) (LD, (bud) cholesteroid values, which may shift the FIDL to LD, bahare in a direction that floors greater risk of attricticetorsis. The reflect impact of an anabolicantogenis stread of serum lipits is dependint on the dose, route of administration (eral w. ijectabel), type of stread (good monthable marks and storage resistance to retachbotin, therefore the stread monthable in non-anomalable instant and storage resistance to retachbotin, therefore has a moderate to most anomalable instant and storage resistance to retachbotin, therefore has a moderate to any also adversely fleet boled present can indirectively and the moderate to support the venericative hypertempty, all potentially increasing the risk of candiovascular disease and moccardial intericon.

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of startated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabolic/androgenic stensis when taken in doess sufficient to promote muscle gain are expected to suppress endogenous testorence production. Whost the intervention of testosteneo-stimulating substances, testosterence levels should return to remult within 1-40 methods of the gasession. Note: that produced hypogenomolytic hypogenomiation can statisk, terbolice was determined to be approximately from times stronger at appressing sundetoriopic nature solutioners or military horis.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (Men):

Terebokee cauratus was never approved for use in harmars. Prescribing gaiddies are unvailable. Common does for physica, and performance-partnering paryons all in the mange of 150-300 mg per week, which is usually taken for 6-10 consection weeks. The week is sufficient to produce considerable in terenosis in item marged means and strength, which needs to the sufficient of produce considerable interests in item marged means the sufficient of the desired reads.

Administration (Women);

Trenholme enanthate was never approved for use in humans. Prescribing guidelines are unavailable:This agent is generally not recommended for women for physique- or performance-enhancing purposes due to strong androgenic nature and tendency to produce virilizing side effects.

Availability:

Trenbolone enanthate is currently made by underground steroid manufacturers only.

Winstrol® (stanozolol)

| Androgenic | 30 |
|-------------------------------------|---|
| Anabolic | 320 |
| Standard | Methyltestosterone (oral) |
| | |
| Chemical Name | 17beta-Hydroxy-17-methyl-5alpha- androstano[3,2-c]pyrazole |
| Chemical Name Estrogenic Activit | androstano[3,2-c]pyrazole |

Description:

Wastrol is the most widely recognized trade nume for the drug staroodel is another of applicationstores, chernela plateral on that the horner's anable (issues) hading properties are greatly arrified and its antheografic activity minited. Staroodel is a checking at a "maked" is trade, and cellular to one of the storgest discourtism of anables to analogues, effect among commercially studied age at 11 also cannot be another that the store of the storegate discourts of a maked on the storegate discourts of a maked on the storegate discourts of a maked one of the storegate discourts of an above of the storegate discourts of a maked one of the storegate discourts of a maked one of the storegate discourts of an above of the discourts of the storegate discourts of a maked by the discourts of the storegate discourts of a maked by the discourts of the discourts of the storegate disc

History:

Samoabi was first described in 1999.⁴⁰ It was developed into a medicine by Wattrep Laboratoris in Grenz Brinin Paret rifts (Defing) Baff of UL2, pattert on the agart 1901, 1^{40} Samoabi was officially relaxed to the U.S. preception drug market in 1962, under the brand mark Watterl, Samoabi was initially prescribed for a variety of medical purposes, including the induction of appetie and lasm issue gain in cases of weight loss associated with many migant and non-migant diseases, the presention of toom muse catables (daring prolonged controlstrend) theory of the prost-operative and post-traum (bown, fincters) pointers, and even to trut (divide) in the elder).

The FDA's corted over the pesception dng market latal gighened by the mid-190%, and the mixtated uses for Warted verse some moreouse. Daring this time for FDA officially supported that Winsteil vasa "Probably Effective" as an adjunct therapy for rearing conceptorsism, and for provincing growth in pinitary-deficient domains. With the positor, Warting was given more time to ad and study the agart Wartings was able to contrainally study of the TDA reagard (Wartish's dashift) as a therapeute agart, and it remained in the dampointing from the materialness. Stranderkl was able shorings some province daring the people of the ripproving end hold call concentrations, cortefuing the strateging of subdermal instase, of any wather strateging of subder dark concentrations, cortefuing the strateging of subdermal instase, of any wather strateging of subder dark concentrations, cortefuing the strateging of subdermal instase, of any wather strateging of subder dark concentrations, cortefuing the strateging of subdermal instase, of any wather strateging of subder dark concentrations and the strateging of subdermal instase, of any with hereiting any advections, a trateging trateging of subdermal instase,

Withings were through a number of corporate changes during the 1990's, including a 1991 merger with 25 starts of its form Sand Weithery, Sand Withings continued in the Within it is U.S. for approximably 10 mere yans, bolton finally disconting the Sand at the true. And the operatively conservation of the true of the true were officially transferred to Outsin Phermeteria. The true of the true of the true were officially transferred to Outsin Phermeteria. The true of the true of the U.S. phermeterial method, altoget in outdars active potentials in 2004 for the Al forms of Wateriol are presently surveilables in the U.S. although the Wateriol benefits are the true of Wateriol are presently surveilables in the U.S. although the Wateriol benefit areas in a start of the true of the t

How Supplied:

Stanozolol is widely available in both harman and veterinary drug murkets. Composition and dosage may vary by country and manufacturer. Stanozolol was originally designed as an oral anabolis storoid, containing 2mg of drug per tablet (Wastrol). Other brands commonly contain 5 mg or 10 mg per tablet. Stanozolol can also be found in injectable preparations. These are most commonly water-based suspensions carving \$2 mg/ml of storoid.

Structural Characteristics:

Stanozolis a modified form of dipdrotestostorene. It differs by: 1) the addition of a methyl group at cachon 1-raphta to protect the hormone during containding and 2) the attachment of a prazal group to the A-ring, replacing the normal 3-keto group (this gives stanozhi the chemical classification of a heterocycle strendy). When viewed in the light of 17-alphn methyldhydrotestorenoe, the A-ring modification on stanozoloi scense to considerably increase is anabolis strendy while ruderize its enditive and modified and the standard strends of the strends of the relative anti-genericity.

Senoxola lus a much weaker relative binding affling for the androgen receptor than testostence or diphotostostence. At the same mix to fashys and host part laff is dara lower affling for semarbinding proteins in comparison. These features (among others) allow standold to be a very potent anabole started in spite of a veaker affling for negregator binding. Recent statises have additionally confirmed that its primary mode if action involves stransform with the collider androgen receptive.⁴⁰⁰ Although end field hexicitated, startoxile may have additional (come potential) unique) properties with regard to antagenism of the progenetic activities.⁴⁰⁰ Although end field hexicitated, startoxifer, independent activities.⁴⁰⁰ Although end field hexicitated, startoxifer, and although end field hexicitated startoxifer activities.⁴⁰⁰ Although end field hexicitated, startoxifer, and although end field hexicitated startoxifer and although end field hexicitated startoxifer activities.⁴⁰⁰ Although end field hexicitated startoxifer and although end field hexicitated startoxifer and although end field hexicitated startoxifer activities.⁴⁰⁰ Although end field hexicitated startoxifer and although end although end field hexicitated startoxifer and although end although end field hexicitated startoxifer although end field hexicitated startoxifer and although end although end although end although end and although end attractivity and although end although end although end although end attractivity and although end although end

Sensorbi is izown to strongly suppress levels of SHRG (see hormos-bring global). The trait is characterized of all ambidicationspires storeds, allough the portexy and from of administration make each Visionsbir particularly effective in the regard. One shall with a strong s

Side Effects (Estrogenic):

Stanookis in at anomized by the body, and is not measurably estrogenic. An anti-estrogenic is not necessary where using this stored, as generomists shaded to be a correct neuamong sensitive individual. Since estrogenic the small explorit with water reterring, stanooki of the store of the

Side Effects (Androgenic):

Albracht essified as an anabole stroid, androgenie sied effects are sill common with this substrate. This may include bosts of objects, and and substrate biological hair growth. Anabolichnedrogenie steroids may also aggreate mile pattern hair loss. Worman are also vanand of the potertal withsing effects of mathetic and potential the site of the site of the despensing of the voice, mentional irregulativity, changes in skin teatance, field in itr growth, and channel entingeness. Additionally, the "Software Hair hardbarear compared have been at metabole and channel entingeness. Additionally, the "Software Hair Mathetane entropy" and the site of the androgenic agents such as testosterone, methandrostenolone, or fluoxymesterone.

Side Effects (Hepatotoxicity):

Sanozobi is a 17-apha algoland compound. This alteration protects the drug from detriminent by the local above, and a very high proceedings of the drug entry into the blockdreamen blowing and administration. C17-apha algoland anabolical models includes a single state and the above and the above and the above and the single size of the above and the size of th

Sanoxida appears to offer less huptic stress than an equivalent dose of Danahol (interdundstreholito), Salidas giving [200 editoxelo) let of up V2 roveck fields to demonstrate clinically-significant charges in markers of liber function, including serum segatate antivi-transferse, ahireir aurici transferses, gairming-diamymentismes, librita, and alaline phosphatuse.⁶¹ Bedine hepatotoxiciy increases as the dosage seculters, so hepatic dopticarito andol stil be a correct in In are instance, high doses (abanc or in combinition with other steroids) have bone implicated in screen hepatotoxicy in hordbacke. Increastiles attanoble has ab hear implicated in screen hepatotexicy in an otherwise healthy hodybalke, oril annobles, or alas anaalarmitie medication whose for toxicy processes or states.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

Side Effects (Cardiovascular):

Ambrickninggneis stendis can huwe debetrious effects on serum choeksterd. This includes a tendency to reduce IDE (apod) cholesteroi valasa and increase IDE. (bud) cholesteroi valasa, which may shift the HDL is DL bahare in a facetion that favors genet risk of antiroicherosis. The reduce impact of an anabolicanlargonic stread on serum high is dependint on the dose, route of administration (call us, injectable), type of steroid (normalized errors aroundable), and lose of restance to hegatin tembers. Staroidal has a strong effect on the hegatic management of cholesteroid bale to is structural resistance to dose of 6 range edpt of nei structural company. High cholesteroid around dose of 6 range edpt of nei structural company and the locations of 37% is healby multi quick structural company. The location of 37% is in the location of 37% is body trajectorized the constructural structural structural bale bad present end trajectorized, reduce cachedula interacionant discustion of the location of 37% is in the location of 37% is observed trajectorized interactions and support the ventrolate logaroutly and trajectorized interaction and support the ventrolate logaroutly and trajectorized interactions and the structural interaction.

Injectable stanookl Ins also been documented to produce strong negative charges in serum injekt. One study was carried of an an good P 2 healty must basjees, and demonstrated a meanarble reduction in HDL clobelsterd value, as well as an increase in LDL and tetal clobelsteroi values, *Clobelsterd value*, and well as an increase in LDL and tetal developing arterior developing of the strong teta strong teta strong developing arterior developing teta strong teta strong teta strong medication when endoweding teta strong teta strong teta strong teta strong developing arterior developing teta strong teta s

To help reduce cardiovascular strain it is advised to minimia an active cardiovascular exercise program and minimize the intake of starated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fash oik (4 grams per day) and a natural cholsterolarinoxidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

Side Effects (Testosterone Suppression):

Al anabočiandrogni steroids when taken in doess sufficient to promote muck gain are expected to suppress endogenous testoremo production. Stanovold is no exception, and is noted for its strong influence on the hypothalmirs-pinting-testicalit axis. Chineal studies giving 10 mg per day to heally muck subjects for 14 days caused the mean plasma testostemore level to fall by $55\%_{10}^{65}$. Without the intervertion of stotosterone-stimiliting advances to the stotosterone-stimiliting advances to the attrave based studies that in 14 morths of from seversion

Note that prolonged hypogonadotrophic hypogonadism can develop secondary to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

Administration (General):

Statiss have shown that taking an oral anabolic steroid with food may decrease its biavailability.^{OB} This is caused by the fat-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undigsted disfary fat, reducing its absorption from the gastrointestrial tract. For maximum utilization, oral forms of stanozolol should be taken on an empty someth.

There can be large discopancies in the steroid particle size between injectable stancodel operantions. For example, Warstel (down zhardono (Spain) was obsigned for human use, and uses a reflect powder that will pass through a 27-gauge needle. Weistenlib + V is a veterinary product in the U.S. and Canada, and has large particles fut and winn in excells smaller than start of injection. Injection of stancodel can be taken in measured eral doses sheadd rightfortm proce instrumbands.

Administration (Men):

The origin prescribing patchens for Wantot called for a dudy doang of 6 mg, which was administed on a schedule of one 2 mg table three times per dug. The usual doages for physique- or performance-inhurcing purposes is between 15 mg and 25 mg per dug, or fiture to fix 6 mg tables, takek for no longer than 6.8 weeks. When used for physique- or performance-inhurcing paporsa, a doage of 30 mg every dute dug by more than 0.0 mg tables, takek for no longer than 6.8 weeks. When used for physique- or performance-inhurcing paporsa, a doage of 30 mg every dute dug by more stables, dust for no longer of the dug base of the stables of the stables of the stables of the stables of the body, and are commonly due in 4.5 mg every third dug. Does of 50 mg per duy with spiciath stamoth are onto a more allough probably on absiest. Once that rejeatable Kram of the dug are expected to huw, miligram for miligram, a partear anable effect than oncell⁴.

Sencokib is clear corbined with other strends for a more durantic result. For example, which holding care might cpt to add in 2004 mug of a substance store (coprime, careflate, or proposal) per vock. The result should be a considerable gain in now muccle distance and the constraints of the strength of the strength of the strength strength of the strength of the strength of the strength of the strength and the strength of the strength of the strength of the strength of the amounting strength strength of the strength of th

Administration (Women):

The origin prescribing galdelins for Winstot called for a dualy doage of 4 ng (one 2ng table true dualy wing women particularly susceptible to the antopartic effects of ambolic storids. This doage was increased to forng (the same as the recommended dose for multi-site storids). This doage was increased to forng (the same as the recommended dose for multi-site storids) and the same of the same as the recommendation increases. The ingentiary commendation of the same as the same storids are same fractional to the same storids with same and produces the same storids and the absolute storids and the same storids are same storids and the same storids and the absolute storids are injectively commonly dual with same storids and the same storids and the absolute storids are injectively commonly dualities? This negrees to a 4 day. Although this composed is washly and toggins, the risk of virilation symptoms cannot be completely excluded over at thermosed to a same story in the s

Availability;

Stanozolol remains widely available as a pharmaceutical product. Its production has been

shting to less regulated markets (maily in Asia) in recent years, however, which locly reflects declining interest in using stanoobl as a medicinal product in the West, and the continuing high demail for this drug among athletes and bodybuilders. In reviewing some of the more popular products and changes on the global pharmacentical market, we have made the following observations.

British Dispensary in Thailand makes a stanozolol product. Their trade name for the drug is Azolol, and it cortains 5 mg of steroid in a 400 tablet boths. The bottle itself looks very similar to that of Androlic, with dark plastic and a shirty chrome top. Be sure to look for the company's holographic sticker when shooping.

Stand (Tradand) is now sold as a generic drug under the March Pharmacetricals label, though is in the same recognizable white bothe with gener-shaded label. The product contains 5 mg of seloxid per tablet, and each bothes holds 2000 ablets. To deter counterfeiting, the bothe now carries a holographic sticker on the front that bears the company logo.

Acdhon in Thailand makes Stanozodon, which comes in the old industry standard of 2 mg of steroid per tablet. It is packaged in bottles of 1,000. Counterfeits are not known to be a problema this time although the product is not widely distributed on the black market either.

Unigen markets the product Stanztab in Thailand. It contains 10 mg per tablet, and is packaged in foil and plastic strips of 10 tablets each (5 strips per box).

Winstrol® tablets and injectable ampules are still produced in Spain, by Desma. This remains the most popular stanzoold injectable in Europe. All boxes are protected with a holographic sticker, which carries the company logo embedded into the image. Note that highly accurate counterfiels are currently in circuidation through Europe and North America.

The Greek generic by Genepharm is still in production. The product should carry a Greek drug ID sticker on the box, which will show a hidden mark under UV light.

Chindied makes a 50 mg/mL igectable stancould in Argaritine called Nabole: Strong: This is the same firm that makes regalar Nabole, a very low dosed (2 mg/mL) version of the same drug. This new product is now much more popular on the black maked than the first, due to the more useable dosage. Note that Chindied prints their logo on the inside of the viti cartor, which offers somewhat of a simple security check (dovisous) one very easy to duplicate).

Anabolico Cimol is an injectable form of stanozolol from Argentina. It comes in multi-dose vials containing 50 mg/mL of the staroid. This item has not been subject to widescale counterficiting but also bears no security features that would deter this practice.

Stanozoland from Landerlan in Paragazy is common on the black market, particularly in South America. It comes in the form of a 10 mg tablets, packaged in bottles of 100 tablets each. The company also makes a 50 mg/ml. injectable.

Also from Paraguay is a generic stanozolol injectable from Indufar. It contains 50 mg/mL of steroid in a 1 mL glass ampule. Three ampules are packaged per box.

Formula Magistral in Argentina makes generic oral and injectable Estanozolol. The oral comes in the form of 10 mg tablets, losse in pill bottles, while the injectable contains 50 mg/mL in multi-dose viak. These products bear a small square holographic security sticker to deter contreficting.

The veterinary compounding pharmacy SMP in Canada makes a generic injectable in 30 mL multi-lose vials (50 mg/mL). This product is made in limited quantifies, however, and is not highly common on the black market.

ANABOLIC AGENTS (NON-STEROID)

Description:

Archikowszki azi a norega-ó esserial idny acid that seven as the principle balling block for the synthesis of discole protogiadmins (such as WG2 and WG2s). These prostagations are integral to protein turnover and match accumulation, and three such important activities as unexcessible block block to the matchs (turnors), forceasing lead IUCT) problemion, and differentiation, and increasing the overall rate of protein synthesis and match growth. Archive and the second rate of protein synthesis and match growth. Archive and the second rate of protein synthesis and match growth. Archive and the second rate of protein synthesis and hypothetic As such is a highly are been as the main thermality product match hypothetic As such is a highly are been attempted by the such that we are in the match instagation. The second rate of the such matching the second term and the stage of the second second second second second second protein the second second second second second second second second protein the second se

Clinical Studies:

In 2005, the Exarcise & Sport Nutrition Laboratory at Huyler University conducted a double-bild placebox controlled aday to determine \$ 50 days or sistance training and anchikoine acid (X-Facter¹⁰) applementation would affect training adaptations in 31 experimed C1 systems training that the site as the same presented at the International Society of Sports Natrition conference on Jane 15, 2006. All subjects inguided a statu of far enguines achds (or ear 2007 enguine) of AA or phene very Sport Anou-Shiperson and the site of the same statu of the same statu of the same statu Shiperson and the site of the same statu of the same statu Shiperson and the same statu of the same statu of the same statu Shiperson and the same statu of the same statu of the same statu Shiperson and the same statu of the same statu of the same statu Shiperson and the same statu of same statu of the same statu of the Anaecohic Power (600% - phacho), and Total Work Capacity (250% - phacho). No side effects were reported in the line statu approx of the same statu of the same statu Anaecohic phane statu of same statu of the same statu of the same statu of the same statu Shiperson and statu of the same statu of the same status of the same s

Pharmacology:

Anchickie acid bagies to display is anabolic activity early during coarcise. This matrice is released from your mack there as they are duringed during tracementing taggiring a localized informatory and ambiotic response. This is part of the same biological process that cases you to be soon ad up or too following and outwards and the during the during of mack there is a during the during the same biological process that control the tracking and strengthening of market has an discussion. In this during mark there, is, similarly, the very hist anabolic targies in a long caseda that the control the tracking and strengthening of market has an discussion. The same during same tracking the same during same during the same during th

The availability of anachicoine acid, and our ability to Benerat it during exercise, is important to the anabolic proceedings of our workshow. We also need to be assore of the fact through the regular exercise significantly lowers the content of anachicoine acid in skettal machtions.⁽¹⁰⁾ 402–305, effect online, proceedings of procession of the second of a subility anachicoine acid, hower levels result in less annehicoine acid height effects of the size stratement models of the layer assors we find a branch of a subility of the size and the size strate models of the layer assors we find a branch of a subility of the size as also one of the kay massors we find a branch or a size sheat machine the same before the size and the strategistic section of the size of the same before the same before the size and the strategistic section of the size of the same before the same before the same balance of the same balance section is the size of the layer as the same of the size and the size strategistic section is the size of the same light stores of anachichonic acid to work with Dan Dachine cores size.⁽¹⁾ The box excessive is the one work are of the size strategistic section of the size of the same size of the same size of the size of the same sis of the same size of the same sis o

History:

The arachidonic acid supplementation protocols, and the concept of using this nutrient to improve muscle mass, strength, and performance, were first developed by William Llewellyn, and/or of this book series. Linesologi flad plater on the technology on November 27, 2002, and released an ancholisic acid singlement user that K. Factor tachemic, Moleculer, Natiritoi shorly after, Mahoudi her product was initial pract that of shorly after. Although the analysis of the short of the short of the short of the short tachemic of locations and the short of the based of the short of the based short of the short of t

Structural Characteristics:

Auchikonia acid (eccoas-58,111,4-encoia acid) is an essential polyareatmented fitty acid found in animal fitts. Supplemental anachikonia acid is commonly produced in two forms, trighceride and ethyl ester. As with other fitty acid supplements such as fits oils, the natural trighceride form (as present in X-Factor and focussed products) offers up to 400% geneter absorption than the ethyl ester, and is the predered form for supplementation.

How Supplied:

Arachidonic acid is sold under the X-Factor trademark by Molecular Nutrition, and is supplied in 250 mg capsules. Arachidonic acid may also be found in a number of licensed products; all will display the pattern number (#6,841,573) on the packaging.

Administration (Short-Term Anabolic):

As a short-term anabole agent, arachikonic acid is napplemented at a dose of 500 mg to [J000 mg per duy C-240 mg capasilos.] The fal J000 mg dose is non convensive user, negaticitos of bodyowight. The native is explicit in the same way sherokic commonly are, and additional anabolic and the same strain and the same strain and the same strain discuss to note mouse the increase in the non-term loss straing and nanovelop isoure. Depending on distary and individual methodic factors, these gains may be accompanied by a decreme in body far, facios of 1-28m of language the same strain and an environgenerating on distary and individual methodic factors, these gains may be accompanied by a down to instrain distance of the same strain and the same strain and the same strain single strain and the same strain and the same strain and the same strain and same strain and the same strain and the same strain and the same strain single strain and the same strain and the same strain and the same strain strain and the same strain and the same strain and the same strain and the other occurs within the worked on initiary appredimention at mathodic levels.

Administration (Normal Supplementation):

Archikolis cicli my also be an important ranient to consider in regalar supplicrental doces, particularly you do ne consume mining products (cont march, eggs) on regalar basis. Statiles have shown that gives somewhat comparable annous to protein, this who consume animal products will march and the protocol of the status exceedence but these that do not (experiments)⁴⁴. Archikolate and may be the missing compound in such also compiled relative suggesting that an anchicosis and discovery estimation of the status of the animal status of the compiled relative suggesting that an anchicosis and discovery estimation of cases, sind original relative suggesting that an anchicosis and discovery estimation of cases, each in high ranked alleless. For those who full their ratike of animal products indicquite, of the full the products. Takine over give true, the capada de 20 mg of disc with animal products. Takine over give true, the capada dad nonsayed of this oscillar to cause das the status of the status and any of the status of the status of the status of the status data products in and any of the status of the the status of the status of

Safety:

In clinical studies involving the supplementation of 1,500-1,700 mg of arachidoric acid per day, general markers of health were also unaffected with 50 days of continuous use. This includes no notable change in HDL, LDL, or total cholesterol values, immune system resources fewtioning or databate assumention values 653 66667. Environment the insection/ran requests extensing, to punct arguinguistic tasks. Transmission in the structure of the stru

On a mere general note, the American Heart Association annovanced to position on Orngu-6 FAR consurption in early 2010. In reviewing some of the more corner data, they found that dies low in Orngu-6 Bis were associated with an increased risk of heart disease compared to dies with higher index. The poper also discusses annovality including this second list is normally not providementatory, or or otawally urhealthy to consum. The American Heart Association new recommends that 5-10% of grout carbies each dargoone from Orngu-6 fais, including AME. Datis rich in Orngu-3's are, of course, also healthy, but net at the ochains of Orngu-6's.

Side Effects:

Anchikofes acid (X-Factor) often prochees an amplification of residual post-workout (Bohyad) Oneny mucks sourcess. Often accovery is sightly probably (feelphage and additional day of rest is required), and this user may need to adjust the schedule and the schedule of the schedule of the schedule of the schedule and represents increased intensity of the much cascade (and the of muck growthe). Those with existing mixer mucks, connection (and the schedule schedule schedule accounties the injust; of growter scnesses interfaces with early labels and percentage of same, which may be due to is effect an a suscellator, hereassing due year percentage of same, which may be due to is effect an a suscellator, hereassing due year excamption of the adjust in this side schedule and a schedules in a same and analysis the side due to its effect an a suscellator, hereassing due year excamption of the adjust in this side direct. Which can be already enough a way weak analogoase effect in some users, often producing mixer ody skin. It may be concern from a host within size the schedule of the schedule of a schedule of the schedule of th

Contraindications:

Those with an existing medical condition related to inflammation may find that the added arachidonic acid exacerbates symptoms of their disorder, and should avoid supplementation. This supplement should only be used after the approval of a physician if someone is taking medication has an existing medical condition or has a familial predisposition for cardiovascular disease, high blood pressure, or any other disorder that may require the limiting of dietary arachidonic acid. Also, as a potent growth-promoting agent, arachidonic acid joins androgens (testosterone, anabolic steroids), growth hormone, IGF-1, estrogens and many other growth factors as potentially supporting the growth rate of certain cancer cells if you have the disease. Dietary arachidonic acid intake has been generally eliminated as a causative factor in cancer. 638 639 just as testosterone level has been eliminated as predictive of prostate cancer risk, however these types of growth-promoting agents should be avoided in such diseased states unless approved by a physician. If you have prostate cancer, for example, the last thing you want to start taking is a growth promoter like testosterone. The same goes for arachidonic acid. The bottom line is that if you are in poor health, you should probably not be taking this supplement. If you are healthy, you should be able to use it with great safety.





During the clinical study, subjects taking X-Factor added an average of 25lbs to their bench press maximum weight in 50 days. This increase was nearly 45% greater than that noted in the exercise-only (placebo) group. Some subjects gained more than 50 lbs on their bench press 1-rep max over the 50-day period.



Using the standard Wingate cycle ergometer test to measure relative peak anaerobic power, subjects taking X-Factor increased leg power by 1.2 Watts-kg-1.This represents a net increase of more than 600% over the placebo group (-2 Watts-kg-1).



In the standard Wingate cycle ergometer test to measure anaerobic power and performance, the X-Factor group outperformed the placebo group on Average Power by an amzing 21 watts (AA: 37.9W P: 17.0W). That's a net increase of nearly 225% commarded to placebo.



Total Work, as recorded in the standard energy unit Joules, increased by 1,292J in the group taking X-Factor, while Total Work increased 510J in the placebo group.This is more than a 250% increase compared to placebo.

Kynoselen®

Description:

Kynocken is an injectuble vestermay drug, carrently produced by the intermitted funbougnak II. contains anitten of hypatrical AMF (advocation encomploquie)a, viami B-12, addim schette, magnesian spartate, and potassian suparatite. This blent mikes for a sectantic "vice" for period againstired to potect an animal" matches most and occur webses after fluxes, spiny; or transm. It is most often used on horses, and is typically applied as an anti-carbical after streams carbicly to used on apprent the viaily of an antimative stream of the after a databased and the stream of the stream of the stream of the stream and the stream of the stream of the stream of the stream of the stream antimative and the stream of the stream of the stream of the stream of the stream 12 or schema in table. Bodybakkes are attracted to Kynocken for its mid anabolic and lipodyte properties.

The principle action ingredient in Kynosekhn is lopatarinely, which is chouseful as an annuo also hold with monoculai limitant and vasoalidary properties. It is also identified as an introduce compound, which increases contractly strength, and minimes fingue, of addeted measels $\frac{d}{dt}$ in grouces that they agreen the own match with the information of a strength of the strength of the differentiation of assisted groups), to construct the differentiation of assisted groups), to construct the differentiation of assisted groups and the distance in the differentiation of assisted for the differentiation of assisted groups and the distance of an advection of the differentiation of assisted groups and then used in lations of an advection of the difference of the difference

History:

Heptaminal, the principle active apredient in Kynoseker, was first heavily investigated in disciant mechacic entrythe entyl 1958. It was son diversible after a prosception drug and has since been sold by a series of drug manufactures in mung different parts of the work! Currently is most common frequentiate use is to trat orthostic helpoprisonis, which is sukhen drop in blood pressure upon starding. Varians preparations containing hypothesis (lab), Contensor (Edgama and Swatchard), High-AMQI (USA), Higheany (Higham and Jupparisol has been contensor and the contact of the start in the set in the start of the start part of the start part of the base nonzero for data, these preparations use ranks (of the start) addies, Carona Integration, and the start of the start of the start of the start and the start of addies of the start o

The French vectoriary preparation Kynoschen worki be the first heptranico-containing dueg to prah hugs-scale intermional natorints (becoming popular marge, alternational to hussbeing mosed that the intere part of the 1990s. This was sorre years subsequent to hussbeing mosed that that increased the penalists for daining in antibodi searoids. During this tran, availability of the drugs land dalloid, and for some bayees scares supply and high lagal risk marke the drugs base simetise. Many attentions that works were becoming increasing/resourceful in finding other mos-scheduled performance-entraning drugs that could be practised and used coses the Antient, Buy attention of the operational provided in the practice of the performance of the thorizen of the year 2000, on nutrities. Today, Kynoseker remains unscheduled and widely availability in the University of States and market and a support of the performance on the datter of States and the practice of performance of the performance of the performance of the datter of States and the performance on the states of the performance of the states of the states. Today Systems and performance on the states in the States and performance on data and another in the trade states. Today Systems and providers to market of the states of the states of the states. Today Systems and performance on the states in the states and the states and performance on the states in the states and performance on the states and the states and the states and performance on the states and the states and performance on the states and the states and the states a

now supplied:

Kynoselen is most commonly supplied in a 100 mL multi-dose vial for injection. Active ingredients are heptaminol, disodic adenosine monophosphate, vitamin B12, selenium (sodium selenite), magnesiam separatate, and potassium asparatate.

Structural Characteristics:

Heptaminol (supplied as heptaminol hydrochloride) is an amino alcohol with a structure of 6amino-2-methylheptan-2-ol.

Administration:

Kynoslen is not approved for use in human. Prescribing galdelines are unrealisable. An effective dosage for hospica- or performance-channeling propose garenby fills in hospica- for performance of the state of the

Because it tends to increase condendin levels, Kyrosekin is also a mid simular 1 is ikely for this specific reason. This is also his see hour only over init horearcing cognizations. This means that one can expect certain simular-related side effects, especially when this ign is indirections. This is includes right horizont, succeing jizzer, testissness, increase block presents, or incomin. Agod rake of future based by bodybalates to ity and keep small side clients from theoreming a problem is nover right crutee black of testiss. The bodyward presents of the testis is the straight set of the straight set of the straight set of the body set of the testis of the testis of the straight set of the straight set of the straight set of the test set of the straight set of the tree test ow steels of show recernents in creases.

Warnings:

Individuals with high blood pressure or cardiovascular disease should not use Kynoselen.

Availability:

Kynoselen usaally sels for 575 to 5100 per bortle at the retail level. It is not a controlled substruer in the United States, and is howing perty asyst to obtain headly or via mill order. Currently no significant fakes are known to exist. Given is abundance and low cost, conterfiels are not expected to be a significant problem annifirms soon. It is also important to note that legitimate Kynoselen is a veterinary dang only, and has never been manufactured for human use.

Lutalyse® (diniprost tromethamine)

Description:

Disposite tomerdurative is a planmenetical form of the unitaral prostagination are series of natural oxygonal unstanted cycles (the synches), which we can start the basis in the basis Annue (adue thus, FCZ alpha is moded in structed cycle that, FCZ alpha is moded in the basis and the structure of the synches). The series of the structure of the synches and the syn

The second secon

Alticis and biolyheakers are attracted to disperse throattimume for 4 strong hermogene and mack-budding properties. The amoleke effect of this ding us been substantiated by clicial statistical wave shown PCP2a to be a strong simulator of protein synthesis, and key to both the inmediate and long-temp physicalical alteptictors to resistance training¹⁴ effects of the Reports from alticles who have experimental with this start garrantly aspect frist comparability and excludent procession of scalable direct distributions of the direct distribution of the start distribution of scalable distributions of the start distribution of the start distribution of scalable distributions of the start distribution of the start distribution of distribution of distributions of the simulation of the start distribution of distribution of distributions and the start distribution of the simulation of distributions are start distributions and hose barries that and distributions are distributions and the solution barries and the start distribution distribution and the distribution of distributions and distributions are distributions and the observables, with many claiming they notice a side transmission of distributions and the observables with any scalable distribution of distributions and distributions and the distribution of distributions and the distribution and distributions and the distribution of distributions and the distribution and distributions and distributions

History:

Dispost to nonchurine was for involuced into clicial medicire in the early 197b. The first approvalue or folding in human patients was to similar absorbing drag medication trimster. It has size ermined of use for his purpose, hor is most commonly associated these days with vortexiny medicire. Here, it is volde applied to heft-finances spatian the estrons cycle and fettility of surross lessited, littered in disposite to medication as multicolation stranger and the stranger and the strate strategies and the little strategies and the strategies of the strategies and the strategies and little strategies and the strategies and little strategies and the strategies and the strategies and strategies and strategies and the strategies and the strategies and strategi

Over the years discposed toordbarrie his supported as a harman medicize tarket a wide market of time farmas, ixeding such product ong products as Amogdanfi, Stocobal, Prostin (Stocobar), Prostin F2 alpha (ULS, Austinia, Inad, Italy, New Zeindar, Scadt, and Postarmer (Edges). The stocobar of the stocobar of the stocobar prosting stocobar (Stocobar), and a stocobar of the stocobar protection of the stocobar of the stocobar of the stocobar met widely available. The stocobar of the stocobar of the stocobar the stocobart medication, and a stocab tarket (Stocobart, Stocobart, Stocob

How Supplied:

Dinoprost tromethamine is most commonly supplied in a multi-dose vial (5 mL-100 mL) in a dose of 5 mg per mL. It is prepared in a sterile solution of water with benzyl alcohol added as a preservative and solution hydroxide and/or hydrochohor is acid to adjust pH.

Structural Characteristics:

Dinoprost tromethamine is the tromethamine salt of prosta,5,13-dien-1-oic acid (PGF2alpha).

Side Effects:

Possible side effects or signs of discreptors tromethanine overceptozene may include such responsory effects as isobonchoostricuic, wikeozija, coujial, jugi irriniston, rugi breating, and ampliptica. Asthrunic individuals may be particularly sacceptibe to these effects. Disroptor may also cause gestoristicarial distattures case also abdomial camping diarthas, uroning and masses. Other effects may include increased palse mite, devided blood possest, ethic, becy, and arreveis, and in worms therine constrainties, suggitable blooding and uteritive or unitary infections. Program worms should not take or landle discovered Downet or disk of the diffective most false also discovered for hearing. unapprox. napprox to sue cuesca anong anness unag unapprox as proprager to performance-relativity proposes are contrast and often externs. This includes promotance screenses at the site of spectrate often begaring with a dal burning almost intradiately after the solt is given. This including generative and a solic contrast of performation of the solt burner of defensites. In the studies generate a solic contrast of performation of the solic signal to the solic signal solic listic screenses and a solic contrast of performances in models in the range the coursely of defensions and an expertise for the solic screense and the solic decourse of defensions at data the solic screenses of the solic decourse of the solic decourse interact defensions and generate for the solic decourse of the solic decourse continue on with the drug, and often report that side effects become more tokerable over time.

Administration:

As a human medication, dinonrost tromethamine is most commonly given intra-amniotically at a dose of 40 mo for the termination of pregnancy. It is also sometimes given orally to pregnant women at a dose of 30-100 mg to induce labor, although this tends to produce more side effects than other more recently adopted medications. When used for physiqueor performance-enhancing purposes, dinoprost tromethamine is generally given by intramuscular injection. Most noted for its ability to generate localized growth common sites of injection include the shoulders, bicens, tricens, calves, chest, back, and leas. The user will typically inject in only one site per day at the start of therapy, but this may be increased to 2 or more injections per day as they become more accustomed to the drug and its side effects. Therany begins slowly and is initiated with a low starting dose of annoximately. 5 millioram per injection. If the first injection were given without significant side effects, the next injection would be increased to 1 milligram. This is slowly increased by .5-1 mg per application until a peak dose is reached. This might be a maximum of 5 mg per injection site. Injection sites are also regularly rotated so that several days separate administration in the same muscle group. Note that for some, the pain after injection is so severe that training for that specific muscle group must be delayed for at least a few days. Individual sensitivity to the drug may, therefore, require modifications of their injection and training schedule to maximize results and comfort

Availability:

Dinoprost tromethamine is available in the U.S. and many other nations as a prescription drug product. It is also found infrequently on the black market.

Anti-Estrogens

Arimidex® (anastrozole)

Description:

Austrotoxis is an anti-sensoriri drug developed for the treatment of advanced breast carcer in worms, Boerdalle, this ages it the first in a new close soft field-generator solution can anomance inhibitors.⁶⁶ It acts by blocking the earyne anomance cancer close are similar by estorage, reducing levels of this hormonic in the body murent of the progression of the disease. This is also the findamental use of turnolistic characterior constrained by the sensor close of the sensor of the sensor of the endogeneous production. The effects of manipusches of the hormonic in the body muerated particles. While how constrained the sensor of the sensor of the sensor (blockaeded), eccept 30 blocks blocks the action of sensors at the body rescribed to posts-morposal sources. Side effects the to findees and hard freeing can present thermoles during therpara and would be much more sensor in pre-morposal paration. For the astrotic angument adultes, annatouck is a special or one in pre-morposal paration. For the astrotic angument adultes, annatouck is applied to minimic the side effects or comparison with multicinom interfaces and hard theory and multicinom present thermoles during multicinom parations in a Nodadex and Provides, and the other or comparison with multicinom interfaces and hard theory and multicinom in the body multicinom present thermoles of the sensor of the s

History:

Anstrokow was doveloped by Zenece Plasmacencineka, and approved for use in the Unied Stass at the ed of 100%. The drug soas doveloped as a new adjust the tummer for openhate breast cancer in postmorpausal fermite patients, an area of medicine that land is long laisacy of anomalin use. Sharanti data, was needed to oldi procering threads wany form stark and part of an externey large marknesser double blind trait laws of the form data stark of the stark and the stark of the stark of the stark of the stark bard or an externey large marknesser double blind trait laws of an other hand the stark of the postmorphic stark of the patients of postmerest laws exact or stark of the sta

How Supplied:

Anastrozole is most commonly supplied in tablets of 1mg,

Structural Characteristics:

Anastrozole is classified as a selective non-steroidal aromatase inhibitor. It has the chemical designation 1,3- benzenediacetonitrile,a,a,a',a'-tetramethyl-5-(1H-1,2,4- triazol-1-yhnethyl).

Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes joint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/legs and headache. Aromatase inhibitors may also decrease hone mineral density. which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with asstrointestinal side effects including rausea and voniting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enanthate is taken in conjunction with an aromatase inhibitor suppression of HDL (good) cholesterol levels become significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health

Administration:

Anstrouch is FDA approved for adjunctive treatment of postmeropausal worms with homoure neceptor-positive end rescent cancer, fine-be treatment of postmeropausal worms with homoure receptor-positive or neceptor nationwine leady adjunction threads the poster cancer, and treatment of adjunced breads cancer in postmeropausal worms with home post dynamic and the state of the state of the state of the state of the large per dynamic breads and the state and the state and the policities with anometry take. Strange and the state of the state and the it is or note that note does not appear to anext the absorption of anasitozote, so the unig may be taken with or between meals.

Availability:

Anastrozole is widely available in the U.S. and many other nations as a prescription drug product. It is also found readily on the black market.

Aromasin® (exemestane)

Description:

Encreastre is a stavilal nickle constasse inhibite. It is vary similar in structure and action to formstrate, and/orable it is significantly more potert in comprastion. As a close of drags, anomanise inhibites offer an artis-totoganic effect by blocking the arryne responsible for symbolic generations. Encreastrate is approved by the TDA for the treatment of breastrate drawer in sources, specificably in pote-transposal plateits whose cancer his progressed following theory with tumoxit. Nucle bodyblacks and athletes onlines the drag for manapproved parsees, namely to conter the strongenic side effects associated with the use of anomatobite analysistand segments and theory and the strongenic side drags for most potent associated with the excession invascular definition. Ensurements in blocks, and haves of body fat and tervenues in vancatar definition. Ensurements in drags for most potent associated with vorom.

History:

Exercisite vus developed by Pharmacia & Upiden (Pharmacia), which gainot FDA approad for site of the drug in the 1999. They introduced it ranke the Acomsin beam of an isome pairse, the company developed in a standard and near mixessily difficult dougs of 25 mg per tablet. The company has since introduced the drug to many other mixes under these match mark. These votions patterns and gained and intervent that the since the since the since the since the since the drug of the since the break since the break since the break since the break since the break since the since th

How Supplied:

Exemestane is most commonly supplied in tablets of 25 mg.

Structural Characteristics:

Exemestane is classified as an irreversible steroidal aromatase inhibitor. It has the chemical designation 6-methyl-enandrosta-1,4-diene-3,17-dione.

Side Effects:

Common side effects associated with the use of an aromnuse inhibitor include for flashes, single rains walkness, fingle mode charges, depension, high block presense, workling of the ambridge, and headche. Aronnuse inhibitors may also decreases borne miceal density, inhibitatisti and the single state of the single state of the single state of the single state muses and vonting. Aronnuse inhibitors can humthe development of an urbane flash, and should never be taken or hunded wiring programs; When itaken hyme mices and reliable used to reache entogeneity during proforaged periods of statoid treatment, aronnuses inhibitors may increase catalyouscied, discose (CVD) in key treatment gaves hereful properties of estrogen on crossstero vatas. Statuses invo demonstrated trait when an anomizable steroid acha setutosterore enamine is taken in conjuction with an arountase inhibito, suppression of HDL (good) cholsetted levels become significantly more prostrated. State the estrogen reception againstitutings in Nodelaced generalized does not over arountase inhibitos for estrogen maintenares by make bodybukkes and albeies concerned with enritoristication and albeies and albeies concerned with enritoristication and albeies or an arountase inhibitos for estrogen maintenares by make bodybukkes and albeies concerned with enritoristication and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen maintenares by make bodybukkes and albeies or an arountase inhibitors for estrogen and an arountase or an arountase inhibitors for estrogen and albeies or an arountase aroundase or an arountase aroundase or an arountase inhibitor or an arountase aroundase or an arountase or an arountase or an aroundase or an arountase or an arountase or an

Administration:

Exercisites is FDA approved for adjunctive treatment of postmeropausal women with estrogen-receptor position early herest carcer with disease progression following turnolises. Therapy is initiated 2-3 years after turnoxiden hus fields to efet a destable response, at which post turnolise is discontinuel. Therather with exercising is correlated after 3-3 additional years, and is completed after 5 years of curnative adjunction drug forengy furnolism and curners the turnor contribution. The doors generated in all statuces is one to an introduction the status of the status of the status of the status of the post-status of the status of the status of the status of the status of the body data with the status of the body data with commonly kine 12.5 gr and the status per data in status of instances in all of a table (12.5 mg) taken every other duy is sufficient to prevent the conset of estrogenesis defects.

Availability:

Exemestane is available in the U.S and in more than three dozen other nations under the Aromasin brand name (Pharmacia). Aromasin, Bkewise, dominates the global market, and is presently the only exemestance product one is likely to encounter.

Clomid® (clomiphene citrate)

Description:

Chempkene erinte is an anti-estingenie ding that is preserbed to wormen to text anoulneary inferit/prediatily to oxida). In faint-inferite is a specialitally efferent to as a nonstatenzial ovalatenzy similarit. The drag works by infrancting with estingan receptors, phasing source, documentum, sugian, and even the overall overall estimation of oppose the negative feedback of estrogene on the lopoflusture-iphtany-contrain axis, entrancing the relatest of gamadomism (and the relative structure) and the negative feedback of estimation of the structure of the structure of the structure of the estimation of the structure of the structure of the structure of the structure cance engress (filteriar negative) keeply leading to conception. Chempkene estima is anti-estimgwise properties of chempkene estimate physical ways and the structure of structure in the structure of the structure of the structure of the structure estimation of the structure of the structure of the structure of the structure structure in the structure of the structure

In more, chemplene cintate also acts as a partial arti-settogen, and may be used to contert some of the side effects of committable starting due in citating growmene bases, which can else obtain our folking starting largences, and larkinging hormone bases, which can else obtain a contrast starting largences, and larking largence bases, which can else obtain a contrast starting largences, and largence largence bases of the starting largence and largence largence bases and largence order, when and against using starting largence largence largence largence largence tasks starting largence largence largence largence largence largence largence tasks starten and largence lar

Note that the triphenylethylene compounds (toremifene citrate, tamoxifen citrate, clomiphene citrate) tend to be somewhat intrinsically estrogenic in the liver. This means that while they can block estingaria activity in some areas of the body, how can actually act as estingaria in this other kay area. The Storgenic action in the low is inportant in the regulation of securithostestor (it tends to support IRU, synthesis and LID, reductions). Since staroli-sing host-balances and activity design with the negation conformation. Store starolism are ablocked design with the negation conformation (seeks) may not ablocks be the bott option. Using a drug thub block gencommatic, for example, while at the same time supporting improved cholesterol values, might be much more ideal.

History:

Chenjehere citate is a fertility dung with a abstartial listory of use in the United States. It first guide widespecta decoptience dring the end [970b, and lists new and any connects the fertility practice over size. The drug is now considered a standard medication for certain forms of fertility hency, and lass been adopted as such for oxidate. Use botter: Complete citate is greatly available in more atmost workback. The two most popular band runnes con is likely the conceptual band of the state of the state of the state of the management of the state that of the state that of the state that of the state that of the state that of the state that of the state of

How Supplied:

Clomiphene citrate is most commonly supplied in tablets of 50 mg.

Structural Characteristics:

Cloniphene citrate is classified as a selective estrogen receptor modulator, with both agonist and artagonist properties. It has the chemical designation 2-[4-(2-chlorol,2-diphenylvinyl) phenoxyl trictlylamine dihydrogen citrate.

Warnings (Visual Symptoms):

Some patters using chorphese crime notice bharing or other visual disturbances such as spots of nishes. These symptoms occur more foregard at higher does or longer durations of theorys, and othen disappear within a few days or weeks of use. Poologaed visual disturbances have been respected after the discontraintion of cloringhene crime theoryge. however, and as norm, cases may be reversable. These taking cloriphene crime theoryge, however, and as norm, cases may be reversable. These taking cloriphene crime theoryge, methodays more historica than its mill with the locat cause of these visual aspects more yet understood. It is advised to discorringe treatment and have a therough medicalopharhonizable committen though they occer.

Side Effects:

Complexe citrat appears to les well tolerated, with a low incidence of significant side effects. Common abserse reactions during cincilca thus incided oursine enlargement (13.6%), suscentor flashes. (10.4%), abbornial disconfort (5.5%), and abcomlaterie bleeding (1.5%), Data also suggests that the prolonged use of champione cincine may atterie bleeding (1.5%). Data also suggests that the prolonged use of champione cincine may active bleeding (1.5%) and abcome and the prolonged use of champione cincine may ensure and potential to the transmission of the cincil of users in programmation synchronic (DHSS). Early warring signs of OHSS include abbornial pain and distortion, muses, direther, and voiett area.

Administration:

Complexe cirate is FDA approved for the treatment of women with ovakitory dysfunction preventing pregnary. The recommended dosage is 50 mg (ab) for 5 days, which is initiated approximately 5 days into the mensional cycle. If ovakiton does not occar, follow up cycles my use a dosage of 100 mg per day for 5 days. Mary circle incises recommend a limit of 6 coarses of therapy. When used by men (off-libble) to migate the setrogenic side effects of ambolichnitogenic steroid use, a daily dosage of 50-100 mg (-72 thibble) is usually administered wike any offacting strends are taken. Note, however, that transition is usually given preference over choriphore extrates for the paprose. More commedy, cherophree crites is used by ment at adoage of 50-100 mg per day for 30 days at the correlation of a strend strends of the taken the taken taken the taken taken taken taken taken between the taken taken the taken taken taken taken taken taken taken concerns program (or BCT). Instez. (ad. Hinseque). Franke misles conceasing the champhrase crites for the reduction of comparison proton taken to the objectivity context, not taken concerns program (or taken concerns taken take

Availability:

Clomphene citrate is widely available on the international market in a variety of brand names. It generally sells for a reasonable price, and is of low interest to counterfeiters.

Cytadren® (aminoglutethimide)

Description:

Aninquietlimités is muitéj técnifield as an téhbier of advencectrical steroid sprintes. Its primury function is to lock the conversion of coldusterol to pregnotome, which is requestion of the theorem and the stream of the str

Cartial Inhibitor: While corticals is an essential hormore for life, its levels may also vary gourd within "normal" mass depending on the individu, their training and discuts statuand many other personal methods. Eactors, it has been a common passai in the spectra and many other personal methods. Eactors, it has been a common passai in substingent gives a spectra of the spectra of the spectra of the spectra and personal personal methods. The spectra of the spectra cataboach hormone jumpars a message to branknow profits whether, derived and Amingdaterinke has been used by a nurber of athletes and bodyhakkes for the paryone, using h1 contribution with maindocarindepring statestick because it has a low level of mathogen inhibition. Tragefare with spectra of athletes and bodyhakkes for the gas been used by a method of the spectra of athletes and bodyhakkes for the gas been and the prime spectra of the spectra of the spectra of the spectra of the method of the spectra spectra of the spectra of the

When first looked at in the realth of lathficis, however, research was here as to hest sup to use annihydativitia as a cortical lowering mis-tabloc. Dabled the "shermal cargos phenomenon", it has been noted that after a short period of regain use year body often material based on the sherman shermal structure of the shermanne, ACTI anningdatemide, resulting in your body resuring its original baseds of cortical production (paging the baseds) of cortical philosophilosymptotic physical philosophilosymptotic requiring the baseds provering its ACTI program and the physical philosophilosymptotic employed participants provering the ACTI program and the philosophilosymptotic employed philosophilosymptotic philosop

It is important to note that while many people believe they have used this drug as an anticatabolic, few have actually taken the correct dosage. Four tablets per day, or 1,000 mg, awarene recovery to cientificantie induities the denviace orazone (the armere reserves) for appears necessary to sugmenting starts are instances: expering to a surgets representes a starcovering characteristic to preparables, and the inget when related control of a size information. These who do ventue this high conversity report faigue and discontifiert, stating that the drug movement of the start of the society fail the drug start proposed nece of the start of the start of the start building append does not seem to be a plausible ore. The cody instances this anther here early head of this drug ever competitive body databases paratheging in the drug and start of the drug start of the drug start of the start experiment on the drug start of the drug drug start of t

Acountses labilistics Anticojdatelinite is an efficient anovantase inhibitor, and tends to inhibito the achievy of this energiest at mark hower closures from thirth warks in equided the inhibitor of corticostoroid productions (^{60, 60, 100}). While a daily closure of 1000 mg is typically achieved at a closure between 250 mg and 350 mg a point where storeg advances and of anti-salinequestic material storegards and and the same and of anti-salinequestic material storegards and the same and of anti-salinequestic material storegards and the same and of anti-salinequestic material storegards and the same and

Aninquefatimité is unally regarded lighty among athletes and bodybalkers as mestoque miniterance agers. Xaniski hrves shown it to be capilot of decremption growtines actively bus so much as 20% after administration of 2:00 mg per day. Patient response mets also show amongdatefinite to be at lasts as effective as taxouries through it to itagie estopes dependent cancer celle, and mere effective under central conditions. Due to is discussed broad may of non-specific activity, however, itaking the potentia librition of nat to decomption, the conclusion of the animation and analogues as well, it is not regarded are and effective and as a regular effective and an anisogues as well, it is not regarded are and effective, and a result of the animation of the animat

History:

Animpletimities was FDA approval as an aniconstant drug in 1960. Side effects were common with meanners, however, icidaling disovieus, diziness, and juratil loss of more control. In 1966 reposes of allowing insufficiency absolutes to animolatelline intervention of the strength of the strength of the strength of the strength of the data to be recently means of the first animolation of anomale and the strength of the trength of the strength o

A rone rime aminoghtefriniek seas evaluable under merzenso hend menes and in more than 2-docum countris. Unable Charlon and Charlon and Charlon and Charlon and Common, and codd be fourd in such mitrons as Argeniran, Australia, Austria, Brazil, Canada, Chiki, Caceda Republe, France, Chermany, Hong Kong, Beinda, Istan, Hong, Malaysia, Nederstank, Norway, New Zhalard, Rassia, Sonth Africa, Spain, Sweden, Shuzbertank, Ulatide Alingsion and the Ulade Stases. Additionsity, the dreg could be found on occasion under other memors including and Marment. The usat mapping of original metagladentials to personiso have enable beam for the set of metager (Series). The set of th

How Supplied:

Structural Characteristics:

Aninoglutethinide is an analog of glutethinide. It has the chemical designation 2-(4-Aninophenyl)-2- ethylglutarinide;3-(4-Aninophenyl)-3-ethylpiperidine-2,6-dione.

Side Effects:

Froquert side effects associated with arritoglatefinitie lexhele fitigg, dizense, skin meds, Force and masses. Other side effects more sputicely despections, quark daysessin, venting, stormeth upset, thysical dysfarxtion, vitiliation, jauridae, edevated cholsterol levek, changes in blode cell coarts, and high blode presesse. Additionally, those bodybalates and athletes taking at a dossge high encodent particular storage and parts in the signal coarts and athletes taking at a dossge high encodent and parts in the joints when roying to lift heavy weight. It seems bagical that this might lead to an increased assocribility in signal. Uses should be careful to alors the tradeende during the dost more common with higher dosed regimes that inhibit the dormal productions of corrient, and are less common with this taking one or to abits type of syst and set-storages. Destin low does anningdatefinitie may cause birth defects, and sheald never be taken during programs.

Administration:

Aniodpathniké is mácially indicated for the teatment of Cabing's syndrome, mentatric broast career in postmeropasal worme, and politikine teatment in non with advanced prostate career. When used to treat Cabing's syndrome, the dosage used may range from 1,000 mg to 2,000 mg per day, o then is conjunction with 2.300 mg (f) functorations or avoid the adversarial adversarial second parameters. Athletes and bodyballene using antimplatientike for control inhibiton with care control table at some second bar adversarial second parameters. Athletes and bodyballene using antimplatientike for activation of 2.400 mg control tables and bodyballene using antimplatientike for activation of 2.400 mg control tables and bodyballene using bar body and tables. A solution of 2.400 mg control tables and bodyballene discarded in phace of cluby short-term adversion. The dosage most commonly used for the 500 mg per day (12 to 2 tables), with 1 tablet (250 mg) per day appearing to be the most common dosage sketed.

Availability:

Aninoglutethinide is produced in a small number of countries, and is a fairly expensive plarmaceutical. As such, it may sell for as much as \$2 per tablet on the black market. This, combined with limited availability, has severely limited its more widespread use.

Evista (raloxifene hydrochloride)

Description:

Radoxien plutechardse is a second-generation Sectorie Esnogan Receptor Medidate (SEM) of the boundnavene family. This drug is smaller in effect to transfin, exhibiting estingan exceptor antigament (blacking) properties in some issues while acting as an enstrogan more in agained and the second se As an anti-strongen, athletes and bodyhakkers mny use this composal to conthur the confogunsi oke effects, as genecomskin, a propose for which relations hydrochiards source better flues state (effects is genecomskin, a propose for which relations hydrochiards source better Polariers, which bodied at how these two agains compared in the intermet of possister pakerst agreements, and the state of the strength of the strength of possister pakerst agreements, and the strength of the strength of the strength of possister and why first bodied (b) theory of 38 patients, recently and used on again and strength on a late strength or any strength of 19% for minister and 49% is bodieded. These very a significant relation of eggenerations are 44% (Snein in relative potency, induction hydrochiavite may offer an alternative to sargest for some cases of generations).

Typical of an anti-settogen, tackedine hydrochierkie skutual also offer some breeff as a tostostnow-stronikaring comport. We see the effect denomeration tasks on a gorps of older met logad 60–70 years), where daily does of 120 mg were able to neurose serun and biosvahkle theorematic basissources by choose the strong service structure of the strong service structure of the strong service structure structure structure structure structure structure structure structure structure set biosciences theorematic structure structu

There are none megatives to inhibiting the actions of entropy that should be addressed. For one, entropy in a bounderial hornour work to come to 1GeT the best. In matics with accommandy partners that saffer from GH hypersecuritors, 60 ang of radioxiden hydrochetekle work cally want also tradingly angrees LTCT beak by an entropy of 10^{-100} . Entranges to the traditional strength of the traditional strength of the short of the strength concentrations in cortain bases, and orientring enzymes involved in the tailoration of glacose for bissase gravitational strength. This is fitther apport for the bolf fitth mati-strength solution and the strength of the strength of the strength of the strength oriented by an end of the strength orient thereas, and the strength of the strength oriented by a strength oriented models polescy of descend thereas.

History:

Baložičen bijucickárské sus developel by El Lijk & Company, and FDA approved for USL sale 10 70%. In the relatedate uses sus that of an osteoporesis teamente, onjet us is ability in streame howe density. In 2007, the FDA equivaled the indicated uses for the postmorphile and the relation of the relation of the relation of the relation postmorphile and over sub-order hybrid by the relation of the relation medica, and is approved for sale in over 5 contrains. The FSath multi feature Company dominants the global mutant, athung a sumf murber of other transla, can be chain, and organo forski priver 3 contrains. The FSath multi feature and a subtional superscience of the sale over the superscience and the relation of the Company dominants the global mutant, athung a sumf murber of other transla, can be chains, and Optimum Gosti, France, Tabol.

How Supplied:

Raloxifene hydrochloride is most commonly supplied in tablets of 60 mg,

Structural Characteristics:

Raloxifine hydrochloride is classified a selective estrogen receptor modulator, with both agorist and antagorist properties. It has the chemical designation 6-Hydroxy-2- (p - hyd roxypheny 1) berzo [b] thien - 3 - yl - p - (2 - piperidinoethoxy)phenyl ketone hydrochloride.

warnings (Stroke):

The FDA mandates that the following warning be present on the prescribing information for fission (natorine) holochrichicy: "WMRONN, ICRECASED RESS, GO FUENOLS, THROMHODAIROLISM AND DECHTI FIROM STROKE. Increased risk of deep view internoiss and palanousy enrolision have been reported with Fissia. Women with active methods and palanousy enrolision have been reported with Fissia. Women with active to stroke occurrent in trial in postmerupanesal women with ducation disease or at increased risk of major coronary events. Consider risk-benefit balance in women at risk for stroke."

Side Effects:

Common sisk effects associated with the use of nationare hydrochardie freizle hot histor-blanking, headers, maintee, waterses, camping, ederma, sweating, at gassion, and dimbanc Lasses common side effects riched heart pairs, squale heart pairs, inflammation of viris susceind with blood cleft, and visual distantances. In more cases inflammation of viris susceind with blood cleft, and visual distantances, have been provided and the state of the state of the state of the state of the molecular cleft cleft and the pairs beam and state of the state of the molecular cleft cleft and the state of the state of the state of the state of the provided state of the state o

Administration:

Rakučken Jedocchénicie is FDA approved for the treatment and prevention of ostoprotosis in postmeropasaul novem, redocing the risk of insusise breast cancer in postmeropasaul women with ostopporosis, and reducing the risk of insusise breast cancer in postmeropasaul administered force per day, without regard to meak. When used (off-hield) to mitigan dedaministered orce per day, without regard to meak. When used (off-hield) to mitigan dedaministered orce per day, without regard to meak when used (off-hield) to mitigan deoloma tek 20 mpt of 00 mpt red.w.

Availability:

Baloxiten byduchétok is available in over 50 coatrins. Askie from a small nurber of obser hundt, fle Sika prodet from El July (A computy is most light) be be encounted Prixe is administration of the state of the state of the state of the state Nondors and Creating and State of the state of the state of the state Nondors and Creating (2) most state of the state of the state of the balakies of aperies immunity (2) most state of the state of the state part of the state part of the state part of the state state.

Fareston® (toremifene citrate)

Description:

Torontine crute is an arti-estrograic drug, specifically classified as a Sektrice Estrogan-Receptor Moduline (ESRM) with mired apprint and artigority properties. It is a nonstorokit planydetlyne dorshock, similar in structure and action to both Nohadec (turondin crute) and Clorad (cloraphieve crute). Torontine crute is used for the transmitted of breast cancer in positrompassal women with estrogan-receptor position or estrogeneous transmitted and the structure of the structure of the attraction of the structure of the structure of the structure of the structure minimum of the structure of the structure of the structure of the structure minimum of the structure of the structure of the structure of the structure structure of the structure of the structure of the structure of the structure structure of attracture of the structure of the structure of the structure of the structure stude specentre (the both f start and tracess dware therein). This may include system of the structure of the struct The triphepdfylex composite (committee citate, turnofilm citrate, cheniphere citate) tend to be sorrowed in trinscally ostrogeni in the foct. This means that which they can block estimates and the source in the best is present that the best means that have an extra source in the best is present in the regulation of source holdsteed (it tends to support FID, synthesis and LDI, notacrisms). Stress started insign both shared background the regulation and/source direct of tends of the set option. Using a direct direct of the set option (stress of the set option) (stress of the set of the set option) (stress of the set of the set of the set option) (stress of the set of t

History:

Toerniñee cinnte was approved by the FDA as a prescription drug in 1997. It is sold in the US, under the Farston heard name, which is made by GIN, Ite, Farston is also available in over two dozen other countries including Australia, Austria, Belgian, Cacch Republic, Fahard, Fance, Genrung, Greece, Hangary, Iednal, Haly, Mexico, Netherlands, New Zealard, Portagal, Russia, South Africa, Spain, Sweden, Switzerland, Thuiland, Tarkey, and the United Kingdon.

How Supplied:

Toremifene citrate is most commonly supplied in tablets of 88.4 mg, which are labeled as (and equate to) 60 mg of toremifene base.

Structural Characteristics:

Toremilene cirate is classified as a selective estrogen receptor modulator, with both agonist and antagonist properties. It has the chemical designation 2-{p-[(Z)-4-chloro-1,2-diphenyl-1-butenyl[benoxy]-b.Nchinethylethylamine cirate (1:1).

Side Effects:

Teorentizes catute appears to be well bolanted, with a low incidence of aerison side effects. In circular tria, corrent side effects associated with its use incidend but fusions (53%), sworing (50%), descaled here emposes (10%), masses (16%), supplied for larger (13%), single and the start of the side of the low of the block of and platelet courts, skin discolutions of education of distarbances, loss of sizength, large loss, depression, jamade, and fages (affinity of the distarbances, loss of sizength, large loss, depression, jamade, and fages (affinity of the presentive).

Administration:

Toernifine cirate is FDA approved for the treatment of metastatic breast cancer in postmenopausal women with estrogen-receptor positive or unknown turnors. The recommended dose is one 60 mg tablet administered once per day. When used (off-label) to mitigate the estrogenic side effects of antibolic/androgenic steroid use, male athletes and bodyholides may use 30 mg to 60 mg per day during steroid treatment.

Availability:

Toremifene citrate is widely available under the Fareston brand name. It is not commonly sold on the black market, nor is it a high profile item for counterfeiters.

Faslodex® (fulvestrant)

Description

Falsestart is a highly selective entrogen receptor attragentic (also closified as an entrogen receptor downgadnes), I earest is a scinn in the body not by arguing the productions of estingan. It by preventing a from centring activity in the body. It does this by binding compares. This mode of action is very similar to Nodauc (tamotifien et al.) and Chend (dourphine critica), allough unlike these to mogarity faboratart does not have mindous entropy. The mode of action is very similar to Nodauc (tamotifien et al.) and Chend (dourphine critica), handing unlike these to mogarity faboratart does not have mindou or as the bar joinsthe entropy matepoint is catch the traterist of the dished by duality the siste effects of eccess entropy cancel by amboliphing the sister offlates and use such as generomistic, fit balan, and thereads outer terthing.

Fubstanti is very potert as an anti-ostroga, significantly more so than order medications for Nodekez and Chenik Alkough it imposite colonge al its response and role is produced. It can all produce an environment of low entrogenitiv on pur with strong arountage broads and the strong strong strong strong strong strong strong strong broads and strong strong strong strong strong strong strong strong broads concer particular with low strong black with first line endorse in tentmers¹⁵. Another shows the drug process tarmer cell tarsour and growth significantly more efficiency functionals, citature 35, strateging progetomer receptor concentrations. Furthermore, it also lends to downegable progetomer receptor concentrations. Tables more due to the low due blants barber, and for the means is bleved to produce alaveration structure energy.

History:

Fuberture was developed by AwarZencea. It was approved as a prescription dug in the ULS, in 2002, and is sold under the Fachdock band runner. He drug is initiated of the treatment of estrogen receptor positive breast cancer with disease progression following initiatival and estrogeney theory (such as instruction). A such as a second of the material disease and the second second second second second second second termine framework for the second second second second second second termine framework framework (second second second second second second termine). The second second second second second second second second second termine framework framework (second second second

How supplied:

Faslodex (fulvestrant) is supplied in pre-filled syringes containing 50-mg/mL fulvestrant, either as a single 5 mL or two 2.5 mL injections. The product must be refrigerated for storage.

Structural Characteristics:

Fulvestrant is an estrogen receptor antagonist. It has the chemical designation 7-alpha-[9-(4,4,5,5,5-penta fluoropentykulphinyf]nonyf]estra-1,3,5-(10)- triene-3,17- beta-diol

Side Effects:

The most common side effects associated with *falseratar* itselute genoritestimi distrutions such as muces, voringic complicity, advorting and distribucommon side effects include healchet, huck pait, hof falses, and some from Less common side effects include in a such of sample, many search indicatos, resulcommon side effects include in a such of sample, and the sample and the blood cell court. Injection side reactions may also court. Attri-strugence and hum the dockpement of an ution fistan, and advalate were be taken daring programs. When taken by mene (as an off-label use) to enduce strugencies (arbitration of a struct internat, a parse takengis and against and the sample and the sample and the structure general bases of the sample and the sample and the sample and the same banchical properties of entropy not challenging where.

Administration:

Fubestant is FDA approved for the treatment of hormone receptor positive metastatic beast cancer in postmeropusal women with disease progression following anti-settogen therapy. The recommended does is 220 mg administered intramasclathy (battock) per study of the estrogene side of the strength of the strength of the strength of the absolution of the attraction of the strength of the strength of the strength of the absolution of the strength o

Availability:

Fulvestrant is available in more than one dozen countries. At the present time, all fulvestrant in circulation is likely to be of the Fashodex brand name. The drug itself is exceedingly expensive, and as a result is not widely traded on the black market.

Femara® (letrozole)

Description:

Leroxok is a roo-steroidi selective firid generation aromuse inhibitor. The trutture and activity of this compound are vey similar to that of Arinisck unstroade, and it is prescribed for similar medical pupposes. More specifically, U.S. prescribing galdeness for knowledge to the used for the trutture of postmerospatial women with entrogen memory-positive or estrogen receptor-unknown (usawe if the cancer is response to company heat cancer, it is typically used as accould her of thourset flat on entrogenterogeness and the standard structure of the structure of the structure of the component flat cancer is the structure of the structure of the structure of the obsphallow and utilized for that his index for a stating on the thruse memory, the effects associated with the structure of aromutable analytical memory methods and the structure of structure of the struct

Letrooke prepetents one of the newer achievements in a long line of drugs tragging anomuse riblicht, it is moning the not potter to tongen-lowering drugs developed to date. It is line facts significantly stronger than non-sectorise first generation aromatose irblichts. Rise Tables and Updaten: The dosage of each hole of Fermins 2.52 milliparus, which according to product information uses sufficient to lower extragan levels by an average of 75% drugs detained line. The date, go lowers appears to define transition gate different is lower as low an 1. and 5. milliparus produced 7.5 and 75% eurogen in thicking negocitories to level or historian tendents. All super anomalies are the second second second level or historian tendents. A large number of people may, therefore, nepoud well to lower dosses of the date.

History:

The U.S. Food & Dag. Administration approval letroxic for preception sair in 1997, where it is old NO-rotati usafe the Forman Inderame. Novaria is also extansive practice the drug in other micros, and more than 70 nutions now carry letroxics as an approved drug. The Formas band is by find the domain appropriate overlively, and is fourth in such micrositic for the present strain strain and the strain strain theory of the strain strain theory encoded on the strain strain strain strain strain strain strain strain strain Graces. However, the microsoft strain strain strain strain strain strain strains. Novaria is not microsoft strain strain strain strain strain strain strain including Fraind, Demurk, Nevray, and Sweden Additionable, iteroxice products can also be found and strain also be frained to the Fourt mole mark microtic program (Sama). Additional (Sama) strain strain strain strain strain training strains, the strain strain strain strain strain strain strain strains and the strain strain strain strain strain strain strain strain strains strain strains, the strain strain strain strain strains strain strain strains strain strains, strain strain strains, and strain strains, strai

How Supplied:

Letrozole is most commonly supplied in tablets of 2.5 mg

Structural Characteristics:

Letrozole is classified as a non-steroidal selective third generation aromatase inhibitor. It has the chemical designation 4,4'(1H-1,2,4-Triazol-1- ylmethylene)dibenzonitrile.

Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes. joint nain weakness fatigue mood changes depression high blood pressure swelling of the arms/less, and headache. Aromatase inhibitors may also decrease bone mineral density, which may lead to osteonomsis and an increase in fractures in suscentible natients. Some individuals may also respond to the medication with gastrointestinal side effects including nausea and voniting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enanthate is taken in conjunction with an aromatase inhibitor, suppression of HDL (good) cholesterol levels become significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrozen maintenance by male bodybuilders and athletes concerned with cardiovascular health

Administration:

Lerooke is TDA approved for 1) adjustat treatment of postmexpasal worms with homore nexperp postive early breast cancer. 2) the exacted adjust treatment of early breast treatment of postmexpasal worms with home received 5 years of adjustar transmit of the treatment of postmexpasal worms with the same receiver positive treatment of postmexpasal worms with treatment of postmexpasal worms with the same receiver postmexpasal worms with the same receiver postmexpasal worms with disease progression following artist-strength trengs. The recommendad dose of fetrovative is one 2.5 rng table administed once per day, whole regard to meak. When used (of klabd) to migate the earlyst is a classical trengst and the same of the postmexpasal worms with disease progression distribution. (2) a classical table (1) and (2) are post to most (2) and (2) and (2) are posted to most (2) are posted to most (2) and (2) are posted to most (2) are

Availability:

Letrozole is most commonly sold under the brand name Fernara by the international drugmanufacturing firm Novartis. It is widely available at the present time.

Fertodur® (cyclofenil)

Description:

Cyckfeini is a non-steevidal anti-sentogen that is used in the treatment of mentional disturbances and anovalous pitterfluit (minishili to outdue). It is we similar instructions Chemidia and Nobadeoffi, and also works in the body as a mixed entrogen againstimatigns: This drug is controlly used for off-hild purposed by mile boldyhaliest and affletes, typically at the conclusion of a staroid cycle for the purpose of mensating endogenous toststocher levels. This is in an attempt to minimic the targetist empiric that matches a period of two analogues levels. This is in a natempt to invince the purpose of mensating endogenous toststocher levels. This is in a natempt to invince the physical exists and purpose attempt of the starbance based on the starbance in the starbance in the starbance based on the starbance of starbance interpret the starbance in the starbance and the starbance to a starbance interpret the starbance in the starbance and the starbance of starbance and starbance in the starbance in the starbance and the starbance and the starbance of the starbance of the starbance of the starbance and the starbance of starbance and starbance of the starbance of the starbance and the starbance of starbance and starbance of the starbance of the starbance and the starbance and the starbance of the starbance of the starbance and the starbance of starbance and starbance of the starbance of the starbance and the starbance of starbance and starbance of the starbance of the starbance and the starbance of starbance of the starbance of the starbance of the starbance of the starbance and the starbance of starbance of the sta

Cyclofenil stimulates the release of testosterone via its anti-estrogenic action. The

byothalmus is one maps size of this. By interforing with the brinding of estrogene to is receptor in this area of the body, sychoid bucks the requires bechaic inhibiton bought fourth by this sea, hornese. The enhanced relates of grandstrapin releasing hornese (GRBP) may read, which is nurve odd aritimate the prinding to higher the relates of hardwarp hornese. Lit's due permay signal that the stars no nervous different terms of the stars of the hardwarp hornese. Lit's due permay signal that the stars on nervous the stars of the bodies of the stars of the hardwarp hornese. Lit's due to the stars of the stars of the stars of the stars has also led to is used array a starvid cycle to prever graceconstati, similar to how Noduceför might be not cycle key horneys in ported to be sorrobatt wateed that Noduceför, which is usually predered as an estrogen maintenance drug. Worne do conscismely field a use of arrive acrosses, most of the arroad cross time when the offsets fut can be hought about by a lowering of entropen activity in the final hody are usually strong and acconstration. Key hardwarp of the stars of the market the star holds are stars.

History:

Cyclofenil was developed during the early 1960s, a time when other agents of the same class (such as tamoxifen and clomiphene) were being thoroughly investigated.

Cyclofini tous scont cleased as a precorption drug agert, solf minky to incruse the churce of orcoreption and to course creatin memoryana symptome. Allowing the drug secreted of drug agood clinical effect without significant health courses, if did not see controls warees should be applied on the completed dispersion. If entrols are applied to detailed, solf, cyclofield did not completely dispepsion, and remained rouble in a circum matter. The most popular band was Fertodia, produced by Schering. It was sold in a few coursits, sincluding Brack Germany, Bab, Marcico, Stokerhard, and Takey, Ohen popular bands. State, Cyclofield did not completely disco. Stokerhard, and Takey, Ohen popular bands the sincluding Bradien (Linky, Manepute, Bradie, Stokerhard, and Schering, Bradien Bradien (Linky, Manepute, Bradien, Washing, Bradien, Bradien, Geordman, Uter dug en all Be for all no masses including Takey, Bab, and Bradie Georeman, Bradien and Bradien (Linky, Bradien, Br

How Supplied:

Cyclofenil is most commonly supplied in tablets of 200 mg

Structural Characteristics:

Cyclofenil is classified as a selective estrogen receptor modulator, with both agonist and artagonist properties. It has the chemical designation 4.4'(Cycloherwidinennethylene)bis(phern) accetate).

Side Effects:

Cyclofini appears to be well tolerated, with a low incidence of significant side effects. Cyclofini algorithm and the second sec

Administration:

Cyclobini in most commonly used (medicably) to text annovalancy infiriting). Therapeatic protocols recommends allow of 200m (grine times per day for 5 days, which is initial near the start of the mensional cycle. If programs, is not achieved with the first cycle, it may be used for 3 or 4 cycles in bluch, have mer intrances the days also alpoint. These of texts of the start of the mensional cycles is a start of the start of the start of the mension of the start of the mension of the start of the start of the start of the start of the manni lastostarone production, a doogae 400-600 mg per days is the most common. It is often used for a per cycles and a doogae 400-600 mg per days is the most common. It is often used for the start of 4 no 5 weeks a num of a comprehension per obs-yets recovery program in place of Chend (see Cycle, Babage). Moodes is usuably given preference for the papeoe. Store and knows have experimented with using cyclobell as a met of the intervent is invited $\theta (MOM)$ in the new of the for 6 knows. While nows have out of the intervent is invited $\theta (MOM)$ into now of the for 6 knows. While nows have the forth the metal is invited $\theta (MOM)$ into no set the for 6 knows. While the babage is a start of the intervent of the interv uses not use purpose is synamic resorrance ing per say an or o weeks, while some nave reported this approach to be effective, many others find the drug too mild, especially in light of the effects of exogenous testosterone.

Availability:

Cycloferil is not widely produced. Availability is presently low on the international market. When located in the U.S., the drug is usually found in the form of Fertodur, made by Schring in Turkey. Counterfiels of cycloferal drugs have not been a significant problem.

Lentaron® (formestane)

Description:

Formstare is closified as a selective investible strovidal anomates inhibits. This agart is struturaly a derinitie of andiostecticitor, affering from this well subson prohemace only by the addition of a 4-hydroxy grame. This grave, however, is responsible for causing an investible attractured between formstates and anomatise when the two come with o carditative structures and anomatic structures and anomatic wells are bus comes in a source with each other. This means that formstature will bord with the enzyme and never it is go, permutarily docationing it as a result. The enzyme will need to be replecid, threagh normal attrition, before the body will recover its lost enzymes arguing size and the structures and the structure of the structure o

Because of its potert entogen-suppressing active, ferrentane has been used circuitally to true hvest cancer prioriter in a rarber of coartieris reiching. Eighed, Germany, Switzerlund, Spain, Austinala, New Zealmel, Lula, and Malaysia. It has been shown to be an effective option as a second lace of defense and truncoling, an estimation of the statistically ariting to coardinate of the state the heart properties at statistically ariting to truncing and the state of the state of the state properties of the state of the state of the state the heart properties of the state metastochild or frame alternative. One state, for example, name, a 70% is left of whether the state of the state state of the state state of the state state of the state o

History:

Formestare was the first selectic anomates inhibitor to be dowkped as a prescription dring for at operating in large drange from 4/100% touch the Learning Taylor and the large start of the large drange from 4/100% touch the Learning Taylor in the Generation of the large start of the large start of the large start of the Generation of the large start of the large start of the large start of the Africa, Spins, Swatzriand, Takeya, and the United Kangdam. With the emergence of never an parl mit. Note of the similal Learning Days of the preparation the size been discontinued. The Beed Casch Reputed being being start of the large start of the size of the large start of the Beed Casch Reputed being being and the United National Start of the large start of the size of the large start of the size of the large start of the large

How Supplied:

Formestane is most commonly supplied in a sterile solution containing 125 mg/mL of drug in a 2 mL ampule.

Structural Characteristics:

Providence to the S2-3 or an orbital colorador to constant, and the shear to be address to be addres

Formestane is classified a steroidal selective inveversible aromatase minipator. If has the chemical designation 4- Hydroxyandrost-4-ene-3,17-dione.

Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes joint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/legs and headache. Aromatase inhibitors may also decrease hone mineral density. which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with asstrointestinal side effects including nausea and voniting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an anomatizable staroid such as testasterone enorthate is taken in conjunction with an anomatase inhibitor summession of HDL (good) cholesterol levels becomes significantly more nronounced. Since the estrogen recentor agonist/antagonist Nakadev@ generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health

Administration:

Formstare is indicated for the reatment of advanced breast cancer in postmeropansal worms. The recommended dosage is 250 mpl transmocker increase for hottocil, very two weeks. Albough not a medically proposed from of the drug, studies live demonstrated that a simile level of consequences and an advance backweel with on use of formstare. To to poor biovanishing, however, the dose needed is around 250 mg per day. When used (off-lebel) to minigen the estiogene size directs of atmosfastering size studies and the estimates of the size and bodybalakes often take 250 mg every two weeks by sizection, or 200 mg per day or adje.

Availability:

Formestane is not widely available as a prescription drug, and consequently is rarely circulated in the athletic community.

Nolvadex® (tamoxifen citrate)

Description:

Tamotific rather is a non-staroid and-storogark drug used widely ne fixed medicate. It is specifially a Science Ensogen-Receptor Modulator (SEMO of the traphosyldyscience family, and possesses both entropy any star and antagosity respective. As each, it may not as a structure of the structure is a structure of the structure of

The primary worry among the afficic/bolybaling population is genecomstin, or the very ungifyd devolpment of famic breast issue in mr. This can be first noticed by the appearance of swelling or a small harp under the nipple. If left to progress, this can develop into a lang hand-basegeneousling that may be an investible concrete without angues. The entrogeneousling that may be investible concrete without angues in a notable lass of detaining and the single to look among the suband) due to the mean. In fact, differences in the electropic to look among the suband lass to the mean. In fact, differences in the estrophysical during the single s Tamodio cinte also possesso the ability to trensue production of FSH (folkic stimularing hummon) and H1(folkic stimularing hummo). This is accompliated by biologin negative foldswite itibilities neared by estrogen at the hypothularus, which (wit the actions of GRHH) foldses the release of the meritored pathtaph breness. This is very start to the function of Chendil Path (Section Pather Pather Researce 1) and the production of C111 cm strainer the Logidy is obtained to the function of the start breness. This is very strainer to the function of the start breness. This is very strainer to the production of the start breness. This is very strainer to the function of the start breness the strainer production and the producting the producting the strainer production transformation entities on the storts relations in hormone to tesk. It is most commonly used as part of a comprehensive pot cycle recovery program (see TC: Pather Cacke Recovery).

Note that like some other triphengledyne compounds, tunnofdin crinte can at as an estimation of the Biogenia ratio in the low ris inportant in the regulation of serun technication, and tends to support IFD, (good) choisestori spatiations. Serun starting implexibilities are attached dating with its regardent of serun structures inpoletakes are attached dating with the regulation of serun structures and tends the support of the serun structure and the serun structures are associated as a structure of the serun structure and the serun structure of the serun structure struct

History:

Tamoxifen citrate was first synthesized in 1962 by ICL. It was made commercially available in the U.S. not long after, but was initially used to treat certain forms of female infertility, a numose for which tamoxifen citrate does not seemed ideally suited. In 1971, the first clinical trials evaluating the effectiveness of tamoxifen citrate in breast cancer patients were undertaken. Two wars later noting the link between estrogen and breast cancer and the success of early trials. ICI pursued marketing the drug in the U.S. to treat breast cancer. It was not until 1977 that FDA approval for this use would finally be granted. Tamoxifen citrate was sold by ICI in a wide number of countries under the Nolvadex brand name (the company would later become known as AstraZeneca). A number of generics and other brands followed, presently too numerous to list. In 1998, the FDA approved expanding the indicated uses of tamoxifen citrate to include breast cancer prevention for women at high risk for developing the disease. In spite of continued clinical success with the drug for both cancer. treatment and prevention, in June 2006 AstraZenica finally discontinued the sale of Nolvadex in the U.S. A number of generic versions are still available in this country, however, ensuring easy patient access to the drug. Tamoxifen citrate is presently the most popular anti-estrogen used by athletes and bodybuilders.

How Supplied:

Tamoxifen citrate is most commonly supplied in tablets of 10 mg or 20 mg.

Structural Characteristics:

Tamoxilen citrate is classified as a selective estrogen receptor modulator, with both agonist and antagonist properties (also known as an estrogen agonist/angagonist). It has the chemical designation (Z)2- [4-(1,2-diphenyl-1-butenyl) phenoxy]-N, Ndimethylethanamine 2hydocryn-1,23 propanetricat/boxylate [1:1].

Side Effects:

Common side effects associated with the administration of tamoxiden citrate include hor thats, sugain blocking, sugaint laciting, guest administration of tamoxiden, lighheadedness, edenni, and hard isso. Other fasted adverse neurons include side in neuron tenders and the state of the state department of the state eneuron of the state eneuron of the state strong many state of the state of th Tamoxifen citrate may cause birth defects and should not be taken during pregnancy.

Administration:

Tamodie notme is indicated for 1) the tontiment of metastatic beast cancer in sourcen and merg. 3 adjoinst treatment of node-spatish beast cancer following threat suggest and indiatives. 3) adjoinst treatment of node-spatish beast cancer is nontranspatial wears following beast suggest and indiatives (4) addoction in incidence of not concer in sourcen with DCIS (Datal Carcinorus in Shi) following beast suggest cancer in sourcen with DCIS (Datal Carcinorus in Shi) following beast suggest matrixer, and () robustic in incidence of most cancer, a done of 10.20 mg is administed twice a day (morning and e-origin). When and by men (delived) to miggest the stars is used to administe the start stars, a day doage of 10.50 mg (1-3 th start) and the start of the start start and the start of the star

It is importent to note that arise-transparates may slighty endace gains made driving as steroid cept, as many andromicalmobic stockies come is cellabilither must powerful analysis effects when accompanied by a sufficient level of estrongen (See Eistrogen Accompanies). This may be one reasons why gains much with a strong accompanies and as more to assume the same strong and the same strong accompanies of the same strong strong matrices to a lower for mode age. Therefore, it is sould absolute its kinetify a specific near fast transform citations (even when things probleming companies) and the use of an arbitrary constraint gain is sub-transformed and the same of an arbitrary strong strong the same strong and the same strong and the same of an arbitrary strong strong the same strong s

Availability:

Tamoxilen citrate is widely manufactured, and can be found in virtually every developed ration of the world. The drug is also commonly circulated on the black market. Given its relatively low price and high availability, counterfeit product do not appear to be a large issue.

Teslac® (testolactone)

Description:

Teachactors is a fast generation non-selective steroidal aroundase irbibotu, used critically to true strongen-dependent brusts carrect: loss carrindo ef action is indexnol, but it is believed to inhibit the anomatose enorgen is a noncompetitive and rinversible marrar. If no, this would be an ancievable that is very similar to that of Leartons (Brussen). This might also explain why cossistion of the days does not provide an immediate restoration of normal strongen production. Like dismutation, it takes several days after cossing use for the holy on compare production. Like dismutation, it takes several days after cossing use for the holy on opplicable negrous beach. It seems bigged based on sinceture and action that the same them covers with stochactore.

Altroph totalextene is technically staroidal in structure, a foffers to analoka or androgarie effect to is user. This is because if does not posses the train accessive to that all activate the antiogen receptor, murby an active 17-bets-hydrogy proge. In fact, is D irrigs an activate the structure of the structure of the structure of the structure is to activate the structure of the structure of the structure of the murby which may be due for instantomic order of the structure of the prosesses zeros used of anisotration structure structure activates the table with the structure of the calibrar and togen receptor . Regardless of this, itstahektene hus been murbers of the calibrar and togen receptor . Regardless of this, itstahektene hus been murbers of the structure book is will coming the structure days the structure of the structure book is will coming the structure of the structure of the structure book is will coming the structure of the structure of the structure book is will coming the structure of the structure of the structure of the structure book is will coming the structure for the structure of the structure book is will coming the structure of the structure book is will coming the structure for the structure of the structure book is will coming the structure of the structure book is will coming the structure of the structure book is will coming the structure of the structure book is will coming the structure of the structure book is will coming the structure of the structure book is will coming the structure of the structure book is will coming the structure book is will coming the structure of the structure of the structure book is will coming the structure of the structure of the structure of the structure book is will coming the structure of the structure book is will coming the structure of the s testolactone is used by athletes and bodybuilders not to increase muscle mass and performance, but to mitigate the estrogenic side effects caused by certain arabloic/androgenic steroids.

Note that the level of aromatose inhibition produced with tostolactors is significantly lower than that produced by the newer selectic field agreention inhibitions such as antisotoxic, lemonik, and caurustane. For example, one study conducted in 1985 showed frat 1,000 mgo fostolactore per day given to rine rounnal mar for a period for a days appressed serum estadial levels by 25%, andrete using the same 1,000 mg does noted a 50% relaction after side of sole. These markets are lowed for what would be expected of the newer find-gamention agents given the substatial estrogen appression figures they have produced durine (rink) trik with worms.

History:

Testaketne was first approved as a prescription dng by the FDA back in 1970. It was an easy nutri-extograph ding, clobbing a molecularly prosourced effect hat failing to reach levels of high clinical success. As other more effective medications legan to starke for the unitary of the stark o

How Supplied:

Testolactone is most commonly supplied in tablets of 50 mg.

Structural Characteristics:

Testolactone is classified as a steroidal noncompetitive irreversible steroidal aromatase inhibitor. It has the chemical designation 13-hydroxy-3-oxo-13,17-secoandrosta-1,4-dien-17-oia acid flogr - lactone.

Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes. joint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/legs, and headache. In 1999, the FDA officially added malaise to the list of possible side effects from this drug, reflecting something bodybuilders had noticed for some time: low estrogen levels can lead to lethargy, as this sex hormone plays an important role in the functioning of the central nervous system. Aromatase inhibitors may also decrease bone mineral density, which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with gastrointestinal side effects including nausea and vomiting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enanthate is taken in conjunction with testolactone, suppression of HDL (good) cholesterol levels becomes significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health.

Administration:

Testaketors is FDA approved as adjunctive therapy in the pullative returnet of advanced or dissemined beams cancer in potenteropausal worms when hormonal therapy is indicated. It may also be used in scenar who were diagoned as having that disseminate these carcitorum when preneropausal is whom courtin factions have been associated barbor of the energy of the strength of the strength of the scenario of the strength scenario of the strength of the strength of the strength of the have for energy any approximation in rules storied users, a dosage of 250 mg (five tablets) is usually taken pred top;

Availability:

Testolactone is no longer commonly used in clinical medicine, and consequently is not manufactured on a large sale globally. Presently a small number of testolactone preparations still exist, but are not commonly diverted for sale on the black market given the very low demand for the drug in this population.

Anti-Prolactin

Dostinex (cabergoline)

Description:

Cabergable is a selective dopumies receptor against. This agart is highly specific in its actions, with a strong millip of the dopumies. Discreptor, and a solid millip of sectodars, and S-HT2-avencein receptore. Its main clinical use is for the treatment of physical sectors and the sector of the dopumies of the sectors of the sectors of the sectors of the dopumies of the sectors are sequence dopumies of the sectors of the sectors are sequence dopumies of the sectors of the sectors of the sectors are sequence dopumies of the sectors of the sectors are sequence dopumies of the sectors are sectors and the sectors of the sectors of the sectors are sectors and the sectors of the sectors are sectors and the sectors of sectors are sectors and the sectors of the sectors of the sectors are sectors and the sectors of the sectors are sectors and the sectors of the sectors of the sectors are sectors and the sectors of the sectors are sectors and the sectors are sectors are sectors and the sectors are sectors are sectors are sectors are sectors are sectors and the sectors are sectors are

Podectin is a somutoropic hormore, in the same family as harma growth hormore (sourmorph). It is a single poptich hormore, containing a chain of 199 annio acids. This makes it similar to (though slight) singer than jarouth hormore, which is muck of 192 annio acts an anabolic agent (is less not to slock through the late the single starphysical value is in vorene, and becomes apparent driving programscy when it acis in milproduction. Chemosching, lessoise, is sometimes used to appress testion postpartmed in these is a particular model at the first is the site of the site of

Although this is almost never associated with makes, high levels of products have actually been related to locating generomatin in a vary und procertage of socios-sing althouse. This disorder is often characterized by small thal disclarge fast bocomes noticables with the approximation of the social social social social social social social social social significant of the social socia

High products looks (as would be associated with the need for calcuptice) are not regularly documented in stoco-sing mithesis, finite - underscoring the relative uncommon nature of this disorder. We do know that estingary physica a similatory rule here, and likely is the key to increasing product association in male, ^{calcuptic} of the star shade, however, show suppresses an asstall horizontal mitheless calcuptication of the star shade shower, show suppression an asstall horizontal influences and not necessarily fully estimates, may be the cause of the distribution of the star shower are relatively inconclusive. One starty analyzed the effects of estistations emails main and those leve are relatively inconclusive. One starty analyzed the effects of estistations emails main and those leve are relatively inconclusive. One starty analyzed physical for severally 15 power mithest and disposition in run and and available conclusion of physical for severally 15 power mithest and disposition in mitiation and only attackas, years of the several several for the effect of the stochastic of the start and the several association of the several association and other starkas, years and the several association and the start association association and the start association association and the start association association and the start association association association and the start association astart association no consistent enect on protectin secretors— A timu sen-auministration study with athletes,⁶⁶⁴ and a fourth clinical with nandrolone,⁶⁶⁵ failed to show an increase in prolactin levels.

History:

Cabergalies was doveloped during the 1980s. The most popular indem muse for its agars in Donines, which is produced in the ULS and muse often counties by the girst pharmeterial complemente Humanica. Dostines entrained market exclusivity on obsergables in the ULS for musy and, but between 2005 and 2007 second papers versions was excludent to the strength of the strength of the strength of the strength Cabergalies is widely available iterationally, and can be found in more than 3-doesn difference counties. Outside the ULS for Bootsen tandem strength of the strength is present, and can be found in more than 3-doesn difference counties. Outside the ULS for Bootsen tandem strength of the strength is present. Strength of the strength of the strength of the strength is present, and the strength of the strength

How Supplied:

Cabergoline is most commonly supplied in tablets of 500mcg,

Structural Characteristics:

Cabergoline is an ergot derivative with the chemical designation 1-[(6-allylergolin-8betayl)carboryl]-1-[3- (dimethylamino)propyl]-3-ethylurea.

Side Effects:

The mot common side effects reported with cabengine use include headacle, muses, and vonting, which covered is 26, 27, and 25% of patients (respectively) receiving the medication during one chical brail⁴⁶⁰ Other potential side effects include (but are not finited to constpatient, or hymoth. Johdward and patients, dizzoses, varies, fininge, antexis, nareasis, malies, depression, morma, hot flashes, harm paphtitions, hypotension, breast grant and area, however masse and headacles were for morp promiter side effects. Many side effects are done related, firther reason for naturing off with the lowest possible effect conceptance of an overduoe, but it does its labelenticino, but hold pressare, and musel congestion. Note that overdose patients may need supportion messares to mise blood pressare.

Administration:

When used medically to ithich products secretion, calvergalite is given in an itial doogs of 500 mg per veck. The some plus hain in one sing doo see of hidde in 2 σ mree does so separate door, The does may be received by 500 mg per weck at monthly iteratively and weak. Allocal does up to 4.5 mg mp be used in some weak at monthly iteratively atthese boolyhaldens to iteh by potentia section (as with the lower of of the theorem of the does grain itel board quark quarks at the for the sources, at which point the does grain itel board quarks quarks at the does at the does grain itel board quark quarks at the does grain itel board quark quarks quarks quarks at the does quark the does quark the does grain the does quark the does quark

Availability:

Cabergoline is not widely used by bodybuilders and athletes, and coresequently is not commonly traded in black market commerce. The drug itself is widely available in legitimate medical commerce.

Description:

Bomovnjen mosjule is a dopaminovinnice arga derivative with D2 dopamire receptor agenist and D1 dopamire receptor antigonist activities. It is used not commonly as a productin inhibitor in cases of hyperproductinensis, a growth hormore supresent in accompaity (hgh dopamire that have the start of the start of the start management of Parkinosi disease. The structure and activity of this drug are very similar to their of acategoine (hormore, the structure and activity of this drug are very similar to their of acategoine (hormore, the structure and activity of this drug are very similar to anabolic indications) and a structure and activity of the structure and anabolic indications and and a structure and activity of the structure and anabolic indications and and a structure and activity of the structure and gencommism).

The most word proposent of Theomorciptine use for fit host is probably Lieb McChrail, and the of the onice solub, *Rhomericiptine*. In *GOI Libro gills* McC. Use, In this book McChrail describes how the drug can be used to normalize the methodsm, such the solution of the neural physical personses to define (which begin to solut the isos of body fit as the duration of definig necrossics) are indiced. A kt of this focuses on leptins, a hormore tookload at as sort of a flatementat, tilling you brain how much allow gines insure values on the duration of defining increases) are regularly consuming (an "unit-instantiat" respond in an appropriate usely for an infort line to hold on the netter stress services are possible. Mantaining mermal leptin similation codd he key to keying any diet productive, and bornecriptine my rideed allow us to do that.

The human medical data concerning the potential took finds data might play in supporting compile file loss is concarring in measurement was again while datas (moreoraptice and capable of measuring to all file loss by a statistically significant dataset, and is concard graphs of measuring tool file loss by a statistically significant dataset, and essential to cataset against set of the statistical significant and the statistical significant and pages were noticing or interaction that the statistical significant anotaneous of week for the remaining 12 words of interactions. The statistical statistical significant anotaneous of week for the dataset of the statistical significant anotaneous of the statistical statistical statistical statistical dataset of the statistical methylical has the too increase the covered functioneous of disting the more instructional statistical s

History:

Beornergine has been used widely in cinical medicine for its indicated used since the 1970s. It is also men here widely distributed thus its counterprot medication cabengalese, which is used for a similar set of clinical indications. In the U.S., the most corrent here that the start of the start means a branched within solid by Novaria. The drug is available in docume of countins, and Beround, Chen, Grifferdura Booms-Kin, Pavide and Gynodel Beomerciptine ermins a common medication today in most developed mittors for its itsread the function to counter in a start of the s

How Supplied:

Bromoriptine mesylate is most commonly supplied in tablets of 2.5 mg and 5 mg. The doses are expressed in terms of base bromocriptine, so each 2.5 mg tablet contains 2.87 mg of bromocriptine mesylate.

Structural Characteristics:

Bromocriptine mesylate is an ergot derivative with the chemical designation (5'S)-2-bromo-12'-hydroxy-2'-(1-methylpropyl)-ergotaman-3',6',18-trione methanesalphorate.

Side Effects:

Bromcriptine can produce a number of unwanted side effects, the most notable being low blood pressure, dizziness, confusion and rausea. These side effects do tend to be dose related with the low recommended droses used in bachbuilting are not Relv to be much truthis for may. Furthe, initial maseas sometimes goes sony affer a couple of applications, now the user becomes accustomed to the day. However, the strong incidence of any unvelcome side effects should warrant discordining therapy, especially Pholo pressure is becoming negatively affected (too low a doop). Less common advesse reactions include anviert, dy month, edems, seiznes, fitigae, headuch, lethenge, meal congestion, rash, elocated here arraymes, and changes in trutys frequency.

Administration:

When used medically to test disorders much cell by log-productimentic dyper secretion of potentia), an initial does of 1.25 mp to 2 mp et adv is usually concentrated. This may be treasmed by 2.5 mp every 2.7 days until an acceptable thremposite dongs is established, using the secretion of the single rements do not be estimated as the secretion of the secretion of the secretion of the single rements do not not be secretion of the log-secret weight for the secretion of the domain of the secretion of the secretion of the secretion of the secretion of the domain state of the secretion of the secretion of the secretion of the secretion of the down comes of the domain although higher does my by regard in some instances. A 6 is not do comes of the domain control of the down of the domain of the domain of the down of the down of the down of the down of the secretion of the down of

Availability:

Beometripite is produced in most developed contrises, including the United States where is sold as a generic long and under the Pathed brand name product corress in the form of both 2.5 mg tables and 5 mg capsales, with 100 doses per bothe. At the patherms, $(10.6 \ rm (rm y) be as produced to the sold of the patherms, the produced to the patherms and the path$

Appetite Stimulants

Periactin (cyproheptadine hydrochloride)

Description:

Cyrolopathie Indicabirulie is a first-generation prescription histamie and senotion ampasist. The doing is not done pion in the U.S. for the treatment of adlegar-teelds symptoms, including hap force, runny mose, irritated eyes, thos, and swelling. It is also FDA approved for the treatment of analysphetic maccines caused by algencing, othan as an algued in the probability of the second second second second second second second second with ATUS for the second to fiscability and the second second second second second second second second with ATUS for factors, cause, or done dolbating disease. Cyrobiophathetic to fiscate generative space and increases in linear growth beyond white is normally actioned with ATUS factors. The second second

Albody this is a controversial use of the drug, references to the appette similaring properties of coprobatizatine hydrochroit are andwarf it the medical lentence. One of the more detailed papers compares the appette increasing effects of coprobaptiatine hydrochrokic to megasto coetarilo⁶⁶⁸ in a goog of I am with weight has associated with HW infection. The other agert, megastrol, is a progastin that was approved by the FDA is investigation of anometa, acchesical or weight loss in patients with ADEs. In Teb parts of the appendix appet and the second second appendix of the patient of the other PDA approved agert megastrol, with justice consuming about 200 contra caloxies per day. and gaining a moderate amount of weight with other medications. While the heardits were similar, the side effects were not. The investigators reported that more than 50% of the patients taking megistrol saffered impotence during the investigation, while the cyprohespaties hybricaberking our patients to megastrol therapy for many patients, especially thus prove to negative deflects associated with this type of homome manipation.

History:

Cyrobeptafice lpdtrechterkie is an endy anti-istantine ding, and hus been soil as a prescriptionatication in mot developed mitters of eachest. It was introduced to the US in 10%1 under the Pertuctual branch annu by Merck & Co. This brand of gynehystaffur, Merck in 20%1 index the Dirac and Annu by Merck & Co. This brand of synehystaffur, Merck in 20%1 index the US and Canabily Marsh Merck & Co. This brand is all out in more thun ore down courteries relating Marshin, Marshin, Beighin, Beinda, Link, Netherlands, New Zaalard, Sonth Africa, Spair, Sweden, Thaihand, and the United Kriggond in also also ultare down of other hand marss around the welf, in bloth single and malis ingestion preparations. A narther of garcer products are still soil in both the value of cyrobeptathen byhochwich the humps, in 1994 the Werk Huahl Organization vanned against using the drug for this parpose.⁶⁰⁰ Regardless of this report, stranger simpler simulation.

How Supplied:

Cyproheptadine hydrochloride is most commonly supplied in tablets of 4 mg.

Structural Characteristics:

Cyproheptadine hydrochloride is antihistaminic and antiserotonergic agent with the chemical designation 4- (5H - diberato [a,d] cyclohepten - 5 - ylidene) - 1 - methylpiperidine hydrochloride sesquithydrate.

Side Effects:

As a finispendion anti-bitamine, cyrotopendise lydocchorise myo be proce to producing a marker of the effects in a success the most correst on their is solution or the classic 'arabic-bitamize leftnage', which is correst to the types of class. For some users, the increases nate cyrothegadise lydochoritism' with a solution or the solution of the product of the solution of the solution of the solution of the solution of the hydrochoritis is not very neticeable, and perlaps a naisance (not strong enough to an ingulation reconstruction and the solution of the solution of the solution of the solution include, but new finited to, datasees, distributed coordination, runscular weakness, musaes, coursing, diatrine, constraint, and lysoc. Durine as correst one distributed the faintess, days strong incidence or superscheme the mediat microfield investige.

Administration:

The doage required for medical purposes may usy depending on the infoldulat and their purchard needs. The established threeparties time for cytohystachin blochknick is 4 by most adults requiring 12 mg to 16 mg edult. The total adult docuge is a total material established threeparties three the start of the start of

Availability.

Cyproheptadine hydrochloride is produced in a wide number of countries. Although it is not commonly traded on the black market, high sapply and the relatively benign nature of this drug (losse controls) make it easily diverted for sale when needed. Given its low demand, counterfering of cyproheptadine hydrochloride preparations is not common.

It is iterating to also note that the Dominian steroid product, Anaboles, actually ickables 15 and of cytopologicalite blochchichtic and each 3 mg methanhousehone tables, which was added by its developers to ficilitate increased calori tanke and weight gain during ambled humpy. The 21 million provided is optimal for a daily does of 24 mg Databalo (a very common daily class.) This is the only common production for the includes coprobepticate hipolechrolic as an angle one.

Cardiovascular Support

Lipid StabilTM

Description:

Lipid Stable comins a comprehensive bland of nearly one down mutual inguGents. Each component in known to play an inportant to bie randrisouscie melha, and mung bues significant criencia support demonstrating beneficial effects on key health muteches. For complex gale poster is one of the holdboard and impose the antioxidant response.¹⁰ ¹⁰ Genore to externit (dambitadie for ECC) and high polyhead contant) is and the first original transmission of the stable of the stable of the stable outdates ensuell. Containing of the STAD and the stable outdate is an externet were stable of the stable of the STAD and the stable outdate is an externet were stable outdated and the STAD and the stable outdated is and externet were stable outdated and the stable outdated in the stable outdated were also been stable outdated and the stable outdated in the stable outdated or the Lipid Stable formalia taching researced¹⁰ pilotostuck.¹⁰⁵ policostucy¹⁰⁵

History:

Lipid Stabl was developed in 2008 by Molecular Natrikin (U.S.). The focus was specifically on designing a supplement that can help support cardiovascular health in ambolicant/open storiod users. The product can be found frough international distribution, athhough it may possibly be considered a drug product in some regions with strict controls on health supplement products.

How Supplied:

Each serving of Lipid Stabil is supplied in 3 capsules, and contains a blend of green tea extract (750 mg), garlic powder (600 mg), inositol hexanicotinate (400 mg), polygonum cuspidatum standardized for resveratrol (200 mg), pantothenic acid (100 mg), phytosterol complex (100 mg), policosanol (10 mg), and selenium (200 mcg).

Side Effects:

Lipid Stabil is a natural dietary supplement and is not expected to have notable side effects.

Administration:

For general choisterol and lipid sepport or as an adjunct to ambolic-introlognic steroid use, Lipid Stabil is generally taken at a dosen gol 73 capasils per dwy. Note that antanal product such as Lipid Stabil may help reduce cardiovascular toxicy, but cannot be relied upon to completely elimite poterialic cardiovascular during from the abuse of ambolicandrogenic steroid drugs. Care should always be taken to menitor all aspects of health when taking AAS solutances.

Availability:

Lipid Stabil is produced in the U.S. by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, and other international markets.

Lovaza® (omega-3 ethyl esters)

Description:

Lozza is a prescription emge-3-acid supplement which contains edgls extenses of conceptrenteric acid (EQ) and docosiberenois caid (DHA). It is fundmentally similar to most over-the-courser fish of applements, except that Lozza is highly printed to drug aquity standards, is mode with high (DPO) concentration of emges) acids, and hag goes through centensic chiral trials for a specified interpreter tas. Otherwise, the teersfie of PED/DA standard teers are produced in the applement print of the teersfie of PED/DA standard teers and the print of PED/DA standard teers are the end of the teersfield of the teersfield of the teersfield of the (>500 mgHL). Clinical analise showed trighcerile endersite how in this tas $e^{i\gamma rout}$ in addition, the prescription approximate has been shown to resource IDL (good) chalsestend beech, income james by $p_{i}^{(N)} \in ED/DAA happenents are commuting$ taken by ambelicitatiogenic storial users in an effort to rochee the negative cardiovascellarmethys of the text of AAS.

The mechanism of action of eicosperaturesis caid (EPA) and docosalescencie acid (DHA) is on fully understool. These energy-3 caids appent to cert the first chored he poperties over serum high through a number of different hat compliantenity pathways. For one, EPA and DFA appent to be efficient a transing the complex hater hypotensity in space of the second of LDL cholescent and indigentities. These omega-3 tasks may also for full optimal-field through the second optimal transition of the second transition of LDL cholescent and indigentities. These onega-3 tasks may also distributed through the second optimal transition of the second optimal transition of the distributed transition of the second optimal transition of the second optimal forgenerative of the second optimal transition of the second optimal transition of the second DHA therefore, appent to be second field, including endoced substrate availability, reduced lipsty optimals, and the second field including optical substrate availability, reduced lipsty optimals, and the second field including optical substrate availability, reduced lipsty optimals, and transition optimal transition optimal transitions of the second substrate availability, reduced lipsty optimals, and transition optimal transitions optimal tr

History:

The first prescription drug product containing comega-3 acids was approved by the U.S. Food and Dag Administration in 2004. It was sold in this market under the Ornecore brand rarene util 2007, when the manufacturez, Relater Pharmacenticals, changed the name of the product to Lozza. This was done to elimitate any continion with the blood clotting medication. Arriar (antinexproc) acid), Lozza is presently distributed in the U.S. by the international drug manufacturer (RussOm/RiKkan:

LINN SUPPLICE.

Lovaza is supplied in soft gelatin capsules containing approximately 900 mg of omega-3acids each. The doage consists mainly of eicosapentaenoic acid (465 mg) and docosahexaenoic acid (375 mg).

Side Effects:

Lovaza is a natural dietary product and is not expected to have notable side effects. A small percentage of patients reported mild adverse reactions during clinical trials, including back pain (2.2%), flu symptoms (3.5%), infection (4.4%), pain (1.8%), angina pectoris (1.3%), infigestion (4.9%), burping (4.9%), rash (1.8%), and latered taste (2.7%).

Administration:

Lozas is prescribed in a dosage of 4 capacies per day for the transmet of very high trilipedneis. Given high ortand mirind access, Lozas is not commonly kate by AAS users, Instack, most steeroid users will administre 4-6 grams per day of a quality fish of appendent for general choisen and high appendents. We nate analysement or prescription drug containing the omega-3-acids EPA and DHA may help reduce cardiovascular toxicity. Du carnot be relied upon to completely drug interime potential dimension from the absence analysis carnot be relied upon to completely drug how to head and how the string AAS solutions:

Availability:

High concentration ornega-3 acid is marketed as a prescription drug product in the U.S. under the Lovaza brand name. Lovaza is also sold in select European and Asian markets. High quality fish oil supplements containing EPA and DHA are widely available over-thecounter in most markets.

Diuretics

Aldactone® (spironolactone)

Description:

Spiroteknow is an antagenist of addosterore and is planmenericable closeful as a deraciit actis to comprehensive hierding an occoptor single competing in the intervention of the second second second second second second intervention of the second second second second second second second robust of column ot second second second second second segments and the second sec

History:

Spironolactore was developed during the htt 1950s, and first saw widespread use in clinical medicine during the early 1960s. The deng filled an important need for a diretic that does not deplete potassium, and therefore has a less dramatic impact on electrolyte balance in the body. In muny regards it is looked at as a "staft" and "milder" dimetic compared to other agents in this general category, such as loop dimetics or thizables, allowing this agent room the spiral method. fer market sability and success. Today, spironolactione is widely distributed throughout most of the developed work! It is vanishible in dozens of brand rames, the most commonly identified is probably Aldacone from Searle. It is also widely solil in mixed-agedient preparations alrappedice durft directic. This includes formentie, as seen in the product Lasilactor, or hydrochkrothizide, in Searle's (also widely distributed) mixed directic product Aldactoxite.

How Supplied:

Spironolactone is most commonly supplied in tablets of 25 mg,

Structural Characteristics:

Spironolactone is an aldosterone antagonist and diaretic. It has the chemical designation 17hydroxy-Talpha- mercapto-3-oxo-17alpha-pregn-4-ene-21-carboxylic acid y-lactone acetate.

Warnings (Dehydration, Elevated Potassium, Death):

The misse of diarcic drags for physique- or performance-enhancing proposes is characterized as a high-risk parkets. Durations may robust di-beharactering level of dehydration and electrohye inhubance when administend without proper medical approxims. May doorkhow here how associated with the misses of these drags. It is address important to note that the superhearmation of potassiant, effect from the starting is grannelly and abused distributions. Excessive potassiant initiate may cause hyperlakimia, which may lead to cardiae invasibinist and possible data.

Side Effects:

Adverse reactions associated with spirotralectore administration may include generomstic, camping, dimetha, downess, beltmyrg knaddek, ski miritation, nah, nemat confision, and doperang of the vice. Spirotralectore has also been down to case at more in the Breast cancer law been reported in some patients recording spirotralectore, but to causal reditoribly has yet bene established. Additionally, this compared may estival androgene properties, as both a weak ribbiter of androgenireceptor binding and tostostorene biosynthetics.

Administration:

When used medically to treat hypertension, the initial recommended desage in adults is 50 mp to 100 mp per det judicide dass. In muga then to weaks for a maximum sequence to be achieved. The dosage may be adjusted later depending on the infinitual needs of the application, strong end of head of the strong per desage of 100 mg per day in a single meming application, is more common. This may be contrained for 3 to 5 days prior to the evert, and will obtain a strong and more defined appearance to the mascles for substantial neutron in the strong prior to nellystancia).

Women are occasionally attracted to spirondacture for is effect as an ari-androgen. It is somerines used as a safey nat at a pair when androga hesh huse boccem problement daring a cycle, and is used in an effort to reduce the risk of opermaner virization symptoms. A doage of 25-57 mig dayly for 1 to 2 weeks is often used for this puppose, and may be crough to ward of side effects while androgen levels decline (the steroid regimen timuted). Since spirondachors it more efficient a low-nice produces and longdometrous touris-bornedness mediation of the steroid section of the steroid than inhibiting androgen action, it is cottaily not to be considered a care-all remely for abornous storio-barries frem her althese.

Since this compound is one of the mikkes (prescription) disretic options, it is a common starting point for an early competion. Once finnitry with its effects, muny attent to achieve a storager level of water loss by mixing spiorohectone with a thizake or fanosemike (Laski). The gual is to provide strong water excertion with loss classification of the stronger disteribution of the stronger dis 50 mg/s0 mg combination is reported to noticeably increases water execution without domains size effects. The probasin mer-baryonic most with spinorcharen should be balanced out with the thankle so procession backs with some the as gravity affected. On the the facuse, darpet per hyperbaryonic spinor spinor spinor spinor spinor spinor densities and appendix personalized spinor spinor spinor spinor spinor explorate robust spinor spinor spinor spinor spinor spinor spinor offset to see darpets by the protosionin spinor generation of a spinor spinor the spinor spin

It is important to note that while Lasix and Hydrodiaril appear to be more effective at inducing short-term water loss, they also have increased risks as compared to potassiumsparing dimetics, and should be approached with caution.

Availability:

Spironolactone is widely manufactured in both single ingredient and multi-ingredient drug preparations. Low cost and wide scale availability make this a poor financial target for counterfeiting.

Dyrenium® (triamterene)

Description:

Timuterees is an ond faretic used medically to text deem. Edem may occur without known cause (cliquid) (edem), or the association with how of tekeny disease, comparison hard failer, corticosteroid/piogedism use, or the comproduction of allosterone. Timuterene hard failer, corticosteroid/piogedism use, or the comproduction of allosterone. Timuterene the proceeving possism beeds. At the main suggests, this dauge produces a procence of possism supportantion is, therefore, instantion within 2 to 4 hars, resulting peak fallowing angle discost of transmost in the subsection of the state of 2-10 hars.

Timutence is uffield (off-label) by holyholders and affects to shot shocknesson suster prior to a bolyholders in particular toy heavy on the increased definition that can easily show have resolved and the short of the short of the short of holyholding today is mady impossible to achieve whites the use of direction. At the same time, direction are benerics. The holyholding today is benefited in the same transmission of the same transmission of the same mady neight to the same transmission of one or name weight class completes law constants and shows the same of one or name weight class completes law completes and the same transmission of one or name weight classical transmission is been of the same transmission of one or name weight classical transmission is been observed to use at not drug tosted and has ample into to relative following a weigh-in, resulting in the athlete complete at a singletic law classical transmission of the same transmission of the sa

History:

Timitence first saw catensive clinical use during the 1960b. It was used largely as a standhone agast of first, but word to the boorne a widdy used agast in clinical medicine in combination products with other district darge. Today this usually includes other, more poter districts like hindses and hop agasts. Hence, the potsissum hose of the stronger darket is balanced to some degree by the potsissian sparing furniteness, which diem seaks imposite the space of the stronger darket and the stronger darket and regulation transmission of infurences are on 3 and Mathematical and the Statiss (Dorentant).

How Supplied:

Trianterene is most commonly supplied in capsules of 50 mg and 100 mg.

Structural Characteristics:

Trianterene is a potassium-sparing diaretic. It has the chemical designation 2,4,7-triamino-6phenyl-pteridine.

Warnings (Dehydration, Death):

The misse of directic drag(s) for physique- or performance-enhancing purposes is characterized as a high-risk persictic Durations may produce a bi-formatering level of dedpedintion and electrolyte inhubance when administered without proper medical aspervision. May deduction have been associated with the misses of these drags. It is adding the risk of the supplementation of potassism, effect from the strange management def risk in potassism, as generally nat advance when the high parameterizations or a disk train protossism, aspension parameterization and the train protossism and the trainent protossism. Determined and the strange potential parameterizations or a disk trainent protossism and the supplementation of the protossism of the trainent parameterization and trainent parameterization and the supplementation of the protossism of the trainent parameterization and the support parameterization of the supplementation of the protossism of the trainent parameterization and the support parameterization of the protossism of the protossism of the support parameterization and the support parameterization of the support parameterization and the support parameterization of the support parameterization and the support parameterization and the support parameterization of the support parameterization and the support parameterization of the support parameterization and the support parameterization of the support parameterization and the support parameterization and the support parameterization and the support parameterization of the support parameterization and the sup

Side Effects:

Timturene use may be associated with elexible involution, including elevated or decreased possismin levels. Signs of elevations in market eramping searces, weakness, lefung; drowniness, rendlemaness, marke jun; mache eramping searces, endead urire volum; no blod pressure, and gastronterinal distanteess. Ohr esid effects may include manes, vonting, jaranket, blod platekt defenses, jaranket, industrust, search ender and ender the search ender and an platentistic transfordentitions, reductions, source next defects characterized an approxemisity natures have been exported relating skin met plate.

Administration:

When used medically to treat hypertension, the usual initial dosage in adults is 100 ng visec dialy after mais (200 ng ed ay). Tism my be increaseds the should never excected 300 ms per day in stual. Among bodybalates, this drug is commonly used for only a few days prior to a compartion, adjusting the dosage over the course to left the best level of duries (effect) may also also leader to be first length on memory with a must adult the level factories (effect) may used to be the first length on memory with a must adult the level factories (effect) must may be a strain the level may also any level may also be a superfixed memory and the user of more than the level memory and the level may also marker stronger during: and hydrochemized to resonance. The dosage of hold agents would be adjusted downsord to comparestate first contrained effects. The dosage of hold agents would be adjusted downsord to comparestate first contrained effects. The dosage is handly to increase short-term dinexis, white a cancing the corteme potsisam has the fur cancels associated on hydrochemized to robage relation of the strained effects. The adjust the more the more of the other and the strained dosage to robage and the other difficults of the other adjust and the other difficults of the other adjust and the other difficults of the other adjust and the strained dosage of hold and the cancels associated the other difficults of the other adjust and the strained dosage of the other adjust and the strained dosage of hold adjust and the contrained dosage of hold adjust and the contrained dosage of hold adjust and the contrained dosage of hold adjust adjust and the strained dosage of hold adjust adjust and the strained distribution of the distribution of the strained distrained on the strained distribution of the s

Availability:

Trianterene is widely sold throughout the developed world. Although single-ingredient preparations containing trianterene are available, this drug is most commonly sold in multiingredient preparations targeting edema and/or high blood pressure. Low cost, modest demand, and high availability make this drug a low-profit target for courter/feing.

Hydrodiuril® (hydrochlorthiazide)

Description:

Hydrochlorothizzde is a directic from the thizzde family, used medically for the reatment of edems and hypertension. This drag acts by reducing the realsorption of electrolytes, thereby increasing the exerction of sodium, potassium, chioride, and consequently water. In comparison to other diaretics, Hydrodiari is stronger than the potassium sparing agent Albahannoff, minimadantanu, har undire that much known and Lain (forwards). While theorem is the stronger than the potassium sparing agent Albahannoff, minimadantanu, har undire that much known and Lain (forwards). While the stronger that the stronger that the stronger that the potassium sparing agent Albahannoff, minimadantanu, har undire that the known and Lain (forwards). While the stronger that the stronger the stronger that the st Autocores (springerene), na water une nue rop agen Lass (artoernee), ware potassin exception is mich less pronounced than that seen with Lasi, the use of a potassin applement (or a potassian rich diej may sill be necessary with this product. This is usually dependent on the dose and duration in which the drug is administered. Cakiam excertion may also be pronounced with thazides, but again, are weaker in this regard than Lasi.

The use of duretics has been increasingly oppular in a number of althetic displays. For states, these diags are very popular among bolyhaidses who use them to also subscatness state before a competition. The althy to have a warning physica outon edus advectations state before a competition. The althy to have a warning physica outon edus advectations and the state outon and the state of the state of the state advectation of the state outon and the state outon and the state warning and the state output in the state out the state out the state that was advected by an earlier weight in measurement. Given that the weight is in standy doer a day for emapily, might use a competition, the advected by the cent weight will duretics, yet have encoughd intu to reasse failed and lookyweight before the event weight will duretic by the system of the state output of the state output of the the event weight will duretic by the system of the state output of the state output of the state weight will duretic by the system output classifies, which can be advected by advectation output output of the state output of the state output of the state output output output output output output output output of the state output out

History:

Hydochkondiazile waa developed daring the 1950s. Gioen the widsopread mune of diseases association with high holo presents and devrus, the drug fourdant sevely neg munch, and quickly achieved large-scale acceptance and distribution. Hydrochkondiazide became a findamental floom divergery in this area of medices, where it remains widely available local Hydrochkondiazide prepunstions are available in virtualy all developed nations, and appear in largedly hardboch of different head in user and gravity productions. Stagle-ingulation multi-ingulated prepunstions, where the drug is often mixed with other actives that focus on dimension from discover memanenter.

How Supplied:

Hydrochlorothiazide is most commonly supplied in tablets of 25 mg and 50 mg.

Structural Characteristics:

Spironolactone is an aldosterone antagonist and diaretic. It has the chemical designation 17hydroxy-Talphamercaptoacetate. 3-oxo-17alpha-pregn-4-ene-21-earboxyfic acid y-lactone acetate.

Warnings (Dehydration, Death):

The misuse of diaretic drugs for physique- or performance-enhancing purposes is characterized as a high-risk practice. Diaretics may produce a life-threatening level of dehydration and electrolyte inhalance when administered without proper medical supervision. Many deaths law been associated with the misuse of these drugs.

Side Effects:

Hydrochterheizikie uer my be associated with electrophe inhalance. This may include prostsam and sodiam defectives, as well as hypochterioric alkabels, unit mesuse in bade biardhorate dar to significant chloride hosts. Signs of electrophe inhalance include dy month, first, valances, kellmerg, downsiere, redsstrassen, mesel e ampeign sietares: reduced arize volume, los blodd pressare; and ganorinstratul disturbances. Other side effects my at hybrid host and the site of t

Administration:

When used methodally to text hypertensors, the usual initial does in addus is 2.5 mg duidy goen as a single does. The does may be increased to 50 mg duidy, olem it no does of 2.5 mg. Note that duidy does above 50 mg are often associated with marked reductions in serum possistant. Althless and holydukakes typically use fins that go 6.5 mg duidy goen it duidy does a duidy due it is duity duidy due to the struct of the duidy does and the structure of the structur

If the application of hydrochochiazide is not producing the desired effect, many bodyhadischichtes with cose to add another daries: (mild) helicer moving on the stronger loop agents. A combinition of a potensistim sparing direct: Bie Addactorell' (opieronbectors) and Hydrochiari is requested as particularity used lay armage and a bodycod to adgith balance dosago of each agent would be rocheed considerably, usually starting with a 25 mg25 mg appleation and working spreads.

It is important to note that the overage of districts, aside from being potentialy very diagnoore, may result in no mech varies toos. This can lead to fit, "defated" looking markets. A higher diretet dosage, likewise, does not always equate to increased definition an macaching? It is usually engeded as agod advec by those in the althetic commandy to inter. Otherwise, the user may be left to make finite dosage adjustments at the left mining, which can be a diagnores and ineffection practice.

Availability:

Hydrochlorothizzide is widely manufactured in both single ingredient and multi-ingredient drug preparations. Low cost and wide scale availability make this a poor financial target for counterfeiting.

Lasix® (furosemide)

Description:

Fuccessric belongs to a cless of drugs Looma as loop directics, which cause the body to exercit works a way to a potssism, addumt, magessian cicking, and chickite. Byey are used most commonly to their default and high block presents. Like other agents of cliffs type, likes, which is a early property direction of the strength of the strength of the mode of action is independent of any ribbiton towards addoctions. Loop directics are among the storaged actions: a protecting on tables on the strength of the proper levels of an and packets may reaction as the storage mode and the storage of the storages of the and packets may react a protecting to proteomism appelment. If the proper levels of develop, blockies is protecting to a stress of the proteins and storages are developed by the storage protecting proteins consequences. If the proper levels of develop, blockies is particularly and your to use with our proper medical approxision.

Athlets and bodybulkers use directiss for a couple of specific purposes, and usually for only bird periods. Competition and/sets in specific purposes, and usually for drags to drop water weight, is an effort to make adjustments in their weight class standings. Since the weight-models is often ad oper odgs bodres a competition, one can drop birthe bodyweight considerably with directs, and be back to normal within hors after drag constant and redpetition. This may provide a some competition advantage, adwaying difficult constants and the source to experise advantage, adwaying the difficult compete at heavier weight funds its or her catagory would discuts. This advantage weight practics in the separation between the second second

bodybuilding.

History:

Fucuencity usus developed daring the early 1960s. Much of the initial research on this directive sus conclusion in farsper, minity deremany and high. The darp proved to be quite successful, however, and within a minitor of years gained worksike anterioriand acceptance and the second se

How Supplied:

Furosemide is most commonly supplied in oral tablets of 20 mg, 40 mg, and 80 mg, and injectable solutions containing 10 mg/ml.

Structural Characteristics:

Furosemide is an anthranilic acid derived loop diuretic. It has the chemical designation 4chloro-N-furfuryl-5- sulfamoylanthranilic acid.

Warnings (Dehydration, Death):

Furosentide is a highly potent diaretic, which can profoundly increase water exerction (diaresis) and lead to deemote depthetion. The misse of diarec's dang3 like fitnessenide for physique- or performance-enhancing purposes is characterized as a high-risk practice. Durieris may produce a like-fracteating leed of delydination and lectroby leribalance when administered without proper medical supervision. Many deaths have been associated with the missue of these drugs.

Side Effects:

Fucuentia use mmy be associated with detrolyse infalmee. This may include the depletion of potsismin (hypothesian), solutin (hypothesian), imposismin (hypothesian), induced in (hypothesian), and the (hypothesian), imposismin (hypothesian), and the hypothesian (hypothesian), induced hypothesian (hypothesian), weakness, kellung (hypothesian), and gatoritational distributions, of their works, how blood pressure, and gatoritational distributions. Other side effects may include potentials, handing and hypothesian (hypothesian), durates, comparised, hypothesian, and gatoritational distributions. Other side effects may include potentials, handing and hypothesian (hypothesian), durates, comparised, handing and hypothesian (hypothesian), durates, comparised, hypothesian, has in height and an ensitive to high, to blood pressure, high blood sagar levels (hypothesian), and and hypothesian ensitience in the hypothesian (hypothesian), handing and hypothesian (hypothesian). Additionally, some may side officies (handistration hypothesian), and infamilian (hybride spassi, beyes, hybride (handistration), hadditionally, and infamilian (hybride spassi, beyes, hybride (handistration), and infamilian (hybride spassi, beyes, hybride (handistration), and infamilian (hybride spassi, beyes, hybride (handisteriand) and hybride hybride and infamilian (hybride spassi), beyes, hybride (handisteriand) and hybride and infamilian (hybride spassi, beyes, hybride (handisteriand) and hybride hybride and infamilian (hybride spassi, beyes, hybride hybride (handisteriand) and hybride and infamilian (hybride spassi, hybride (handisteriand) hybride hybride and infamilian (hybride) secole (hybride (handisteriand) hybride).

Administration:

When used medically to treat ederm, it is often given only in a dose of 20 mg to 50 mg per day, which is laken to ensigh application. The the treatment of Dypertrems is, it is general recommended to administer 30 mg per day, which is given in two separate 40 mg applications spaced 12. Dones quart, Athlese and bodyhakes toppical use its dwt (god hishe) for very bird protocks (second dosy) of vatar adjament. The dosage and method of administration is taking to the rainbirdual, dependent on the dosed and coefficient of start with two dose, and increase the amount digity on tablesquered days. The main locus is to calculate the operation of the dosed of the domese the dose of the dose of the tot calculate the operation of the dosed of the domese the dose of the dose of the tot calculate the operation. The raind dosegs is usually 20 mg to 40 mg, and the maximum dose interactions 40 mm. The dose is usually 20 mg to 40 mg, and the maximum to a shore occupient of the dose of the maximum distribution of the specification of the dose of the maximum to a dose occupient of the dosed is usually 20 mg to 40 mg, and the maximum dose interactions and the specification of the dosed of th

drug, it is generally used for no longer than 4-5 days.

Note that since functomide has such a strong effect on electrolyte levels, it is generally considered much after to dad a protession paying gard like Aldenette (jejiorolactoru) than it is to keep increasing the arround of farosemide used. Combination diarteris like this are widely produced as a prescription metaficines for this reason. The use of 30 mg Aldestende and 20 mg farosemide is a common starting post, and is believed to lave a roughly simile diarteric effects of any off forwards, but which the two relations of the optimal loss. This dosage may be adjusted on subsequent days in order to determine the optimal models and dosage data and the data and the starteristic diarteristic may be adjusted for a subsequent data in order to determine the optimal to common't and the dange can be active from why hows. It can become difficult occurred the deleptating effect with an overlapping schedule, therefore one should be careful not to administer direction committies example some data.

Injectable foroemide exkeins are considered to be significantly more powerful forms of the drug miligram for miligram. Fursonistic solutions can be administered intramacentity or intraversolgi, depending on the individual needs of the patient. The IV method is much more required the start of the start o

Availability:

Furocentic is widely available, and is menufactured and sold nucler many different brand names, in numy countries. No version of Lasix (or any other diaretic) is currently being counterfield on any large scale. Although it is doubted linke will circulate, make sure to never purchase the drug in 500 mg tablets. These are used only in severe medical conditions, and contain a doage that would likely prove fail to a healty person.

Endurance/Erythropoietic Drugs

Aranesp® (darbepoetin alfa)

Description:

Dachspectra alli, is a syntheic derivative of the human exploration protein. It also hus the same pharmocological action as eccorribute human exploration (exporti alli), in the body, exploration is normally relaxed by the kinksys in response to hypoxia (low blocd coupen level). This in turinggars howe murvo is treasure effolded effordation. As such, this hormore is vial to the regulation of normal red blocd effordation. As such murves, which are the regulation of normal red blocd effordation. As humans, Whit a saline mode of action to recordinate effordation, the displaced in all hands on the is FDA approval for the mammar of amount (low red blocd effordation). In the effect of the strength of the mammar of amount (low red blocd effordation).

Dachpotent alls differs from recordward harma crythropotein (spectra alls) minky in its duration of activity. This new protein ministris is bevis in the block of approximally bires fractions. The second second second second second second second second developed and a principation of the second second second second and they would spectra alls. While spectra alls usually injected on a schedule of three times per works, durbaport all in equivalences of the second particular distributions and the second s Endances athesa are highly attracted to darboportin alls for the effect it has once blood of production. It is no scores that the prostice of blood dopping has been product with work blood only formation of the store of the store of the store of the store constraints of the store boy has restored the blood store. It is adding the store of adding the gener concentration of red blood coles. The blood should, larvesis, transport corgons mere effectively, and the allies may be given a significant endance boost the proceeds on the based processing of the store of

History:

Debeopting alls use developed as a proceeding drug by Angam. The U.S.Food & Dug. Administration first approved 1 for soils in 2001. Angam is the word's largest biotechnologies company, and the same firm that first horoget recombinant explores (protont alls) to the U.S. market at 1944. The main focus with dispersion alls approace to have been the development of a mark larger acting exploresize protein in companying the dispersion of the development of the source of protect and a schedule of the explorest and the source of the source of the source of the source of schedule, which source to offer string confer advantages to exposite all for printers that do not for recovering focus protection alls have neg treated the level of multice success that have noted with propering in preparations, but has been a strong selling drug for Arganycer arises releases.

How Supplied:

Darbepoetin alfa is most commonly found in single-dose vials and prefilled syringes containing 25, 40, 60, 100, 150, 200, 300, or 500 mcg of drug,

Structural Characteristics:

Darbepoetin alfa is a 165-amino acid protein that differs from human erythropoietin by the substitution of amino acids on the erythropoietin peptide backbone, which allows the addition of two additional N-Inked oflogssacharing chains.

Warnings (Death, Viral Disease):

The misuse of darbepoetin alla for physique- or performance-enhancing purposes is characterized as a high-risk practice. Like traditional blood doping methods, darbepoetin alla can produce an abnormally high concentration of hemoglobin in the blood (polycythemia), which may result in heart attack, stroke, seizure, or death.

Some forms of darbepoetin alfa contain albunin, a purified human blood product. Although effective donor screening and product manifacturing procedures are in place, it still carries a risk, though externely remote, for transmission of viral disease.

Side Effects:

Side effects associated with the use of dathepotent and in my include fashies supprotons such as favor, etilik, handache, marche pains wanknoss, or diziness, su heffects tent to be more pronounced at the inflation of therapy. Other adverse reactions my include inflection, rank, solitogi etili fa sida, mange, worting diminica, high biodo grassare, kov biodo pressare, kov biodo pressar

Administration:

Darbsportin allis is indicated for the treatment of anemia associated with chronic renal fahrer (CRF) and chemotherapy. The recommended starting dose for the treatment of anemin in adult CRF patients is 4.5 mcg/kg body weight, administered once per week as a single IV or SC injection. The dosage is adhesequently adjusted based on changes in hermatorit. Healthy athletes using dathepotent allis for performance-enhancing purposes gareadly start on the vey low end of the thrapeatic spectrum, and adjust according to changes in hermatecrit. This may entail initiating therapy with as little as .05 mcgkg of body weight once per week. Note that it is considered very important to monitor bodo cell courts closely during the errite initiace of darbepoetin alls to help ensure hermatecrit is not allowed to increase to an unhealthy level.

Availability:

Darbsportin allis is not wiskly sold on the black market. Yet because of the high cost for explropoiesis simulating agents like darbeportin alli, it is a high profile target for courterfeit dung munificationing operations. Counterfeit drugs of this class have even inflated legitimus pharmaceutical distribution channels, suggesting flat care should be taken when purchasing this and similar drug products.

Epogen® (epoetin alfa)

Description:

Epitopointi is a glocopotein that is produced in the kithney, and is responsible for similaring obbio cloci production. Epiconi nalis is patrumericatific from of orthropointy, which uses munificatured using recombinant TAA technology. The compound is produced from animal cells visue which the gave coding for human orthropointien bis been inserted. The biological activity and structure of option fails are indisrupsiable from that of human exploropoints. Epiconi fails used to trant many forms of americ, faffectively stranking and minimizing orthropoints in a large protecting of patients treated. The efficiency of this ding appeddyments it a nature optionerment for old its effectively thropsing and producting appropriate protecting of patients treated. The efficiency of this ding appeddyments it a nature optionerment for old its effectively thropsing and the structure of the struc

Endances athetiss are highly attracted to spoein alls for the effect it has oned blood cell productions. It is no section that produce to blood doping has been propart with endances sports. This procedure involves neuroning, concentrating and sering a partly of rela blood endances and the spoet of the spoet of the spoet spoet spoet spoet spoet and the spoet endances. The spoet spoet spoet spoet spoet spoet spoet spoet spoet endances are spoet spo

History:

Epocin all was developed by the biotechnologies fran Angen, and frst introduced to the U.S. murket in 1994. The relases of the drag is regardled as a breakmody in the cumarts of ancenia, which beforehand was being addressed multiply with agents that infraredy or mompecifically trapical cell calculations. Least as outpreholses, which my present a number of nanomical side effects to the patient. Epocht and in muched the development of the find up that specifically and discribed yatimutation the process of professions (net block cell production). Its ancess was regard and for texplanality Epocht alls has size been bener (distributed) to Orthen, mundicational by Aragaal, Epocht alls has size been (Jornet and Development). The orther of the texplanality of the texplanality of the (ofference in all). Orthen, mundicational by Aragaal, Epocht of Aragaal, Epocht alls (ofference in alls).

In 2022 the subcataneous use of Figures, which is sold only ontake the United States, was indicated to a rent disease called pure role-of onlymins. This is a conflict where the body boses is ability to produce real blood (eds). Those that suffer from pure red-coll sphesini satably Figures were identified as developing this mate conflicton, the rescore of neural expected numbers. Iternal meetingsinous hydrogeneous activation of the product that were mode to satisfie Rampaer regulations between the rows stabilities that According to a company spokespector, a themical metchine between the new stabilities that the company subcategories. The state of t reaction. The incidence of pure-red cell anemia in patients receiving erythropoietin seems to have been reduced as a result, although varianting about this reaction remain on the prescribing information for all products sold in the U.S.

How Supplied:

Epoetin alfa is supplied as a dry sterile powder that requires reconstitution with sterile dilatent before injection. It is most commonly found in single- and multi-dose ampules and vials containing 2000-40,000 Units/ml.

Structural Characteristics:

Epoetin alfa is a single chain polypeptide hormone containing 165 amino acids. It is identical in structure to the alpha glycoform of human erythropoietin.

Warnings (Death, Viral Disease):

The misuse of epoetin alla for physique- or performance-enhancing purposes is characterized as a high-risk practice. Like traditional blood doping methods, epoetin alla can produce an abnormally high concentration of hemoglobin in the blood (polycythemia), which may result in heart attack, stroke, seizure, or death.

Many forms of epoetin alfa contain alburnin, a purified human blood product. Although effective donor screening and product manufacturing procedures are in place, it still carries an extremely remote risk for transmission of viral disease.

Side Effects:

Side effects associated with the use of epoetin ali my include fuelke symptoms such as dree, rikh, backade, marke pair, wasawase, or dozines. Such effects tend to be more pronounced at the inflation of therapy. Other side effects include rash, swelling of the skin, massa, warring, durined, high block of present, hypekalamin (scess potesiami in the associated with functhoorthodim, deep-wein thrombosis, palmeary embolism, heart attack, and corebrookstellar accidents.

Administration:

Epochs and injectable solution is given by advectanceson or irranverses lepicities. The two paths of administration lone grandy difference effects on the block board of the day. When given by PV indicase, pack block beets of the drug are reached within 15 minus, and the model of the day of the approximately 24 hours. Given an aqual dose, the peak phasen concentration of spectra day will be significantly loss are marked between models. When used metadoly to test a scenarons associated with choice rend likes, the tecommendal starting dosage range is 50 to 30 block of the days of the between the day of the tecommendal starting dosage range is 50 to 30 adds according to choose, in the moster, Harding will be use using days of the theory of the theory of the theory of the theory of the tecommendal starting dosage range is 50 to 30 adds according to choose is memore. The more starting the days are shown at the days of the tecommendal starting dosage range is the transvert. The more and the size starting cosis and the order tecommendal starting days and the days are shown at the tecommendal starting dosage range is 50 to 30 to 10 the days of bodyneight them 3 minus per work. Note that it is considered over a reare henticories in a days of the above the tecomes on an undeally low (1).

Availability:

Epoetin alfa is a very expensive compound, and its use is additionally isolated to certain athletic fields. As such, it is not widely traded on the black market. Given the high cost of this drug, however, it is a lacrative tranget for counterficiers.

Provigil® (modafinil)

Description:

Mediafia, larvows chemically as hverlydylsubjinityciecumike, is a central stindart (operbolinitaria). It is TDA approved for the treatment of matcoleny (6 disense chancelristed by saidlen and usconnoblek anales of deep skep, menal filipae, or inscipato far a mathefar of deet uses, hverling the treatment of Athémer's disease, dereosito, and atterion defect disorker. Mediafia between a stopped of the Engories 'grand amount,' disease and a stopped and the stopped and Engories' transformation.' A single of the stopped and the stopped and Engories' transformation.'' disease and about a advectoring agostic centring in a discussion interfarming in the stories and advectoring agostic centring in a discussion interfarming in the stories and advectoring agostic centring in a discussion interfarming in the control distance between the CNSE in the control interfarming in the CMSE and the distance between the CNSE in the control interfarming in the CMSE and the distance between the constant and the control interfarming in the constant and t

The use of mediafial as a simular has been shown in stuffs; to how mmy advartages our ampletomizes. To begin with, it is bidget of how a much how portform if a show deto the fact that it produces a lower same of explorin. It also displays lower peripheral CNS simulation (less side effects), has minimal effects on bold pressure, produces no interruptions in normal skeping patterns (no langueser or needing "tath-by" skep), and has an owerall patter study profile according to chinal tath. This day advantage to the study of the USs million, which is looking at it as an every effector for phots and comba studies that need to operate for heigh periods of time which sheets that and Docubiers (an ampletoning) of flags, and million comban plasm and soldies have used. Docubiers (an ampletoning) they minimize oxellar optime latering in gray of ab low without day, and hus days althout the particular days and soldies have used to constant study seems size directs than Decohren. Modeling has been tested in overe or orthat studies such as in Adjuntation and (an ad seems posited for fulfical acceptance as a battleful day.

Recertly, mediful has become a popular drug arong competition addets. They use it not simply to 'May avoid' bit as a a porformance enducing agent with both stimulated and endurance chronicasing properties. This type of tage probably corns as a surprise to from who strong stimular properties for words in prove address (performance). Recert making strong stimular properties for words in prove address (performance). Recert making of both strong strong terms of the strong strong strong and the first strong strong strong strong strong strong strong strong strong of both strong of both strong st

History:

Mediafia was developed by Lafan Laboratories in France. It was approved for sule in the United States by the PIA in 1998, where was introduced under the Provigibil Bernal rune. Mediafia is also found iteraturdicably indice this and several additional tanket mems studied and the provide state of the two single states and the provide states that is a francella existing by recent and provide. It was guidely considered as an excitation above in the U.S. It is presently closified as a schedule VI controlled substates, which places moduli and the most category as Wolfman and Xuane. This is introded to him is diversified approximes the same category as Wolfman and Xuane. This is introde to him its diversified in the moduli applications for the day and fully bounds, however, including Proscreptions for the dang are knowned, commonly granted by physicians in general medical practice.

Modafiii quicely became popular among competitive athletes between 2000 and 2004, bloch test athlets bolics were source of the drug. Its use as performance-relatencing agent was revealed to the public damge the designer stared (RALCO) doping scandal of 2004, however, when it was dicklosed that murray of the same athletes due to tested positive for TRG also used modafiii The 10CC quickly barned its use, and with the help of a marker of resourches a methodicage to descript the chemical in the true was doexeliped. This test is now implemented as part of the standard Olympic level drug screening process. Most of the other international athletic bodies have followed the IOC's lead in barning and testing for modafinil. The drug has since lost all appeal as an "invisible" performance-infrancing agent, about is still being used by many athletes that are not subject to random urine testine.

How Supplied:

Modafinil is most commonly supplied in tablets of 100 mg and 200 mg each.

Structural Characteristics:

Modafinil is a central nervous system stimulant related to adrafinil. It has the chemical designation 2- [(diphenylmethyl)sulfinyl[acetamide.

Side Effects:

Side effects associated with modufial are commonly the result of its central nervous system similarity activities, and my include revorsenses, isomiri, shikness, exploring, personality charges, and excitation. The drug may also produce gestoristestinal disturbances such as musea, vonting, addomiral paria, dry morth, anorexia, and headach. Hypertension, heart pulpitations, or albornial heart rate may also be noticed. In nave instances allergie rash, increases in adaliant polopethase, or impaired volutary movement have been reported.

Administration:

When used chinally to trust excessise daytime skeptises associated with marcollapsy or obstructions skept approx. In the recommendable does is typically 200 a 400 mg per day. This may be given in a single morning application, or in two dokked doess (menting and at malka). When used to enhance physical performance, the typical differito does is in the range of 100-400 mg. This is often given at least 2-3 hours prior to athletic competition. Note that side effects can be dow dependent. It is often absideed to actuating mandfind in the low end of the effective dosage range, and increase by 50-100 mg per application until an optimal level is dominant.

The arrens in which this drug is applied are vars, and essentially include any sport focused on acobic activity or endmanne. It may also work well with hole and histors focused on short repeat barsts of discupping produced (mancrobic activities), such as short-parting, pols-vanking, or insigning produced hold hist near a popular drug may togehold hists, as it holds hist direct effective pre-swedent stimulates, especially during protectly when frigge or loss of physical drug murphysical dissocration to have work or percental strendards.

Availability:

Modafinil is presently available in more than two dozen countries. The drug is not highly diverted for black market sale, however, and is not a lucrative target for drug counterfeiting operations.

Fat Loss Agents - Sympathomimetics

Adipex-P® (phentermine hydrochloride)

Description:

Phertermine hydrochoride is a synpathonimiet stimulator of the amphetamine family. Like other amphetamine derivatives, it is categorized as an anorectic (appetite suppressing) agent. Phertermine is commonly prescribed as a weight loss aid in obese patients. It is typically used for short periods of true (less than 12 weeks), and as an adjunct to support an orogoing services and disting regiment. The main focus is to curb the desire to est, threely veckning the total caloric intake. Although the data seems to vary from trial to trial, much of it supports at least a modest additional loss of fat mass with the use of phentermine hydrochloride.⁵⁰⁶ Athletes and bodybuilders use phentermine hydrochloride for the same purpose, typically when weight loss is required for physique remodeling or competition.

History:

Phenemic hydrochicitic was first introduced to the U.S. dang mutket in the 1970s. Base phenemics uses subset in the U.S. as for the data (see 1999). Phenemic mel do tayborn uses as an appeter aspressent, although the most ratable attention to *E* came in the early 1990s, when the day mass successful pared with first firstmanne during distribution. Insertigators had shown that the type of data contribution was actually more effective at pomering weight isos than data and assessing, successful pared with first particular states and the prescription weight loss drug market. By 1997, however, it had become apparent that a way inflapreneting of Frenh unsers were noticitig that to take datases as used of the datages. For during the data state of the data state of the dataset of the dataset of the dataset of the dataset. For during the dataset of the dataset and any effective dataset of the dataset o

How Supplied:

Phentermine hydrochloride is most commonly supplied in tablets and capsules of 18.75 mg and 37.5 mg each.

Structural Characteristics:

Phentermine hydrochloride is a central stimulant and indirect-acting sympathonimetic of the amphetamine family. It has the chemical designation 2-methyl-1- phenylpropan-2-amine (2methyl-amphetamine).

Side Effects:

Common side effects associated with phertermine hydrochloride include is mornin, increased blood pressure, rivelihoit, nervolanesa, and explorin. Iacs common side effects include vision distarbunces, reduced block, confusion, darmhea, doržense, day roundh, beadchea, ringalle hardbead, massa, vourning rand, and itsudness. Berthermine is a CNS stimular haldwarinos, blood pressure irregularities, irregalle hardbead, urrowerkiesness, tronbing shaking panie, enterne needlesses, and soeven museu, contributes, of diarbase.

Administration:

For optimal effectiveness, phetermine hydrochrite's should not be taken with food. The usual shult does is one capasio or table (7.3 rug da)a, administeral before or 1-2 houses after breakfat. For some parities a hulf of a tablet (18.5 rug da)a, primisteral before or 1-3 houses in the cases. It may be advalable to give a hulf of a tablet (18.5 rug da)a ymay be adoquate, while in other cases it may be advalable to give a hulf of a tablet (18.5 rug da)a ymay be adoquate. The daya is gived hand for 4 houses is dould near be taken within 4-6 hours of days. The daya is gived hand for 4-4 houses is a first, with heuse datations of themput days in a sinitar when-term fashion. As to the high labelined of side effects as the does eachies beyond the neural heuronest came.

Availability:

Phentermine hydrochloride is available in a nurber of different countries. It is not widely counterfeited. U.S. residents would not be advised to order the drug from overseas, however, since phentermine is a schedule IV controlled substance and carries similar legal restrictions as Valian and anabole steroids. Many U.S. doctors who specialize in weight loss medications will readily disperse phentermine for controlled periods divised to weight besen medications.

Description:

History:

Abareto silite was introduced to the U.S. due market in 1980, oil under the Verloih brand rune. Abareto silite has grown to bee or of the not opportude rdue jn in bistory for the market. The verlotion brand rune is silite abareto in the market. The verlotion brand rune is silite abareto proved the abareto silite also approved a veriety of other generic and brand rune from of the drue, Abareton silite is also preserves silite in both Hankim and and approxement proveer. In FADA the Bahahi, Braventh Proker, Roveni, Adaroni, Adaroni, Adaroni, Annel, Bahahi, Braventh Proker, Roveni, Adaroni, Salano, and Volume.

How Supplied:

Albuterol sulfate is most commonly supplied in oral metered dose inhalers and tablets of 2 mg, 4 mg, or 8 mg each.

Structural Characteristics:

Abuterol sulfate (sabutamol sulphate) is a short-acting ,2- adrenergic receptor agonist. It is a racenic drug with the chemical designation (±) a1-[(tert-butylamino)methyl]-4- hydroxym-xydne-a.a1-diol sulfate (21)(salt).

Side Effects:

Common side effects associated with albered) safilite include headshet, dözizses, glightadadhess, isomin, herom, arvonszens, sowaitg, massa, vonitg, dianska, and dy month, Lass common hat more sarious advense events include albergie razeñore (resh, linko, sowafing of het heartoga, raz dianda albergie razeñore (resh, linko, sowafing of het heartoga, raz Abateni safilet is a CNS stimulart with operatial for fail overdous. Sgutt of orelones muy include, angle hearding bables presser: ragalarite, irregalar heartbeat, traoneicoanses, terrability, aduity, participation and the same and sovere masse, varing or diarhats.

Administration:

The usual starting doosge for shalls and children 12 years and older for the numerosero of startins 3-4 rug free or for times per day. Where used of like hole for the loss, an efficient does of abstration based starts in the range of one to too 4 rug tables per day (1 high X 1-2 explorations). This is online researed sightly as the user becomes accentrated to the day, perhaps to 4 rug frees to four imps per day. Individuals were yaconitor to the simulation discuss of the start discuss of the start or concentrations in the blood for at much of the days as possible. Althetis and bodynaklers with online use the blood per gravy instand. As in note with all beta againsts, tokernare to the thermogenic benefits of this dang tends to develop quick). This is subaph noteed by the body temperature treating to normal protocold levels. These to the potential side effects of these dangs it is not advected to with usable yot to discontinue the dang for some time (where does not heavy) to let the body network is normal beta-advecorgic receptor concentrations. More recently, the atfiliatures 22, delete, floating by the boccer people, which is a potent properties of the boccer people, which is a potent properties of the attraction of the advecorgic receptors, concent days the soft boccer and the boccer boccer day days which a potent properties of the advecorgic receptors, opecidely these? Interpolarity of the advecorgic receptors, opecidely these? Interpolarity of the advecorgic days and the advecord days people advecord day

Availability:

Albuterol is a widely available and very cheap medicine. Counterfeiting is not a strong concern with this medication.

Clenasma (clenbuterol hydrochloride)

Description:

Carbaterol hydrochirick is an artis-saftran medication that belongs to a broad group of drugs harows as symptomizeds. These drugs affect that symptomic arrows systems in a wide number of ways, length mediated by the distribution of admonceptors. These are adjusted to the start of the start is a start of the start of the start of the start of the start is a start, hydrochinas or advances of the start of the start of the start is a start, hydrochinas or advances and the start of the origination of the start is a start of the start of the start of the start of the the start of the theorem are not when the origin affect the hydrochina or advance of the hydrochina direct entry is which the drug affect the hydrochina of the direction of the hydrochina theorem are not which are drug affect the hydrochina of the start of the hydrochina direct entry is which the drug affect the hydrochina of the hydrochina of the hydrochina direct entry is which the drug affect the hydrochina of thydrochina of the hydrochina of thydrochina of the hydrochi

Most of the actions of catecholomics and sympathemistric agents can be classified into sover bread space: (1) perpendent actional variation space of models matcles such as these in blood vessels supplying the skin, klabey, and mexans and actional space of the start of the start of the start of the start aperphenel sublicity actions on certain of the top end smooth means. Such as those the wall of the gat, in the bronchial tree, and in blood vessels supplying skeletal means. (a) a confluence actions y caliton space start ones the start rate and force of connections. (d) methodical actions, such as an increase in the rate of starts. (e) an outcome control start, and and should vessels and point starts. (e) such and the starts and the start difficult of the start of starts. (c) such actions can be an outcome in variabilities and point starts. (e) endocrine carlors, such as methodies and point starts of the adapt, and (c) prosparity actions that start is cliber inhibition of realistics of endocrine of the starts methodies and point restrictions. (c) and starts of the starts and the start is cliber inhibition of a relations in specific and (f) prosparity actions that start is cliber inhibition of activity.

Clubited hydroklinki is specifially a selectic beta 2 sympthomizetti, primaly affecting only one Other selects of beta-response. Of particular interest is the fact that this drug has like beta-1 similaring activity. Since beta-1 incorptons are closely like to the contain effect of these agarets, this also see obtabened hydroklinki to ordear research array observation (an effect of beta-2) similarion with much liss cardiouscular side effects compared to non-selective beta against. Clinical statiss with the adaption of a given is a control of the selective beta against. Clinical statiss with the adaption is given is a control of the selective beta against. Clinical statiss with the adaption of the selective clicklinearies by the selective or software and the selective beta against clicklinearies by the selective selective against clicklinearies and the selective activity, respiring only a single or twice dudy doing schedule at most. This of concernenties handware rest to be against to use that may the it is is high-optimizer at.

In animal studies clenbuterol hydrochloride is shown to exhibit anabolic activity, obviously an attractive trait to a bodybuilder or athlete. This compound is additionally a known themagare, with beta-2 agoesis like cleabarol hydrochrick shown to directly stratike free clean at accelena the brokelown of regredencies to ferm for feature in the single and blochbarding commands. These interest of this cap on other hydroc is 10 produce 1 his blochbarding commands. These interest of this cap on other hydroc is 10 produce 1 his mack muss. The six was well pointed out by a review publicled in the August 100 S asso of Medicine and Science in *Opstrum all* Lexins, the problem handlock in the August 100 S asso of Medicine and Science in *Opstrum all* Lexins, the problem handlock in the Numet 100 S and the enquired for brochchardina. While and proper bloch historic muss at all. Some seen to sover by the fact that 1 historic and the size of the other histories and the size of the other size of valuable time and measy on dupt that of the other histories are been as the size of the valuable time and measy on dupt that do not built muck. The debate over electrotices that work whether and measy on dupt that do not built muck. The debate over electrotices of the size and the size of the size of the debate over electrotices the size and the size of the cleanes over electrotices the size and the size of the size of the debate over electrotices the size and the size and measy on dupt that of the size of the size of the debate over electrotices the size and the size of the

History:

Cherkardt Jadreckärde hus been avalable as a brocheolikate for decades and is widely used in many parts of two workl. Minogh the nage out sikely record and approxia in a wide in the binds States. The left ratio are need to make avalable for human use in the binds States. The left ratio are need to make avalable for human use in the binds States. The left ratio are need to be the site of the site of states of the site of th

How Supplied:

Clenbuterol hydrochloride is most commonly supplied in oral tablets of 20mcg each. It is also supplied in oral syrups, injectable solutions, and for inhalation use.

Structural Characteristics:

Clenbuterol hydrochloride is a long-acting selective ,2- adrenergic receptor agonist. It has the chemical designation 1-(4-amino-3,5-dichloro-phenyl)-2-(tertbutylamino) ethanol.

Side Effects:

The possible side effects of clerbulenol hydrochietelis include those of other CNS stimulars, and include such concurrences as alsely hands, incornin, sowering, increased blood pressure, and masses. These side effects will generally adside after a work or so of use, once the tear becomes accustomer to the darg. Clerkined hydrochietelist is a CNS stimular with potentiation, environment and the structure of the hydrochietelist blood pressure endoscoses, and severe manaes, working or darking, parks, enterer redstructure of the structure of california of the structure of the structure of the structure of the structure of california of the structure of the structure

Administration:

When used for the management of saftran, the most common clinical doos for relation is Diarray (1) table)) into per day. Some patients require up to 40mg/2 tables)) into egre they be a strained to the second strained tables and the second strained by the day between the second strained strained tables and tables and the period strained strained strained strained strained strained strained strained strained period strained str

The drug will usually elevate the body temperature shortly after therapy is initiated. The rise in temperature is commonly .5 to 1 degree, sometimes a little more. This elevation is due to onc's tody burning excess energy (https): from fig1, and is usually not unconfrontle). The marber of connective days clearatered bulcherike is now usual is usually dependent on the response of the individual To be clear, the athatic hereful of this drug will only the fir a limited time at the dimensity. Ings) who to beta-receptor downgaliants, By most accounts clearatero blycheckiratis seems to sork will for approximately 4 to 6 weeks. Durg this peed, assumed, branc drug the body temperature any angle to task. We again some level of assume that clearbasten high-clearbast is working by the temperature grants are effective, and instance that clearbasten high-clearbast is working by the temperature regarded as effective, and instance clearbasten high-clearbast is down and the source of regarded as effective, and instance clearbasten hydroclearbast is down and the source of regarded as effective, and instance clearbasten hydroclearbast is down and the source of the source of the source.

May bolyhalting competitos enhance the fit harming effect of cherkarol hydrochkerick with use of additional solutances. May have commented that where the drug is combined with thread learning and the powerfal Cytomelli, the thermagenic effect can become externey domains: Such an ris of one firstform and array as strond cycle, height the individual click a much more toued physicage from the drugs. A combined analobic effect of HGH theoryp. Lady, learotish holewood to climate the thermagenic and analobic effect of HGH theoryp. Lady, learotish holewood to climate the thermagenic and analobic effect of HGH theoryp. Lady, learotish holewood to climate the thermagenic and hydrochket (allowing the user to tack in the posteroy of climate data) and one of clambared hydrochket (allowing the user to tack is clearlatorsto), but also of perlapse cere and write receiver downreadanting click for a more comprehensive discussion.

Availability:

Cleributerol hydrochloride is readily available on the international market. Although it is usually a very cheap drug in common source countries, allowing black market dealers ample opportunity to obtain legitimate drugs to divert for sale, cleributerol hydrochloride has been the subject of low-level counterfeiting. A few things are important to note:

Clenbuterol hydrochloride is not produced in the U.S., so avoid anything bearing a U.S. company name.

Clenbuterol hydrochloride should only be trusted when found with a proper brand name from a foreign drug maker. Spiropert, Novegam and Oxyflux from Mexico are the most common products in the U.S.

From Europe, the brand names of Spiropent, Broncoterol, Clenasma, Monores, Contraspasmin and Ventolase are popular.

Bulgarian clenbuterol hydrochloride is also found commonly, but so are counterfeits. This is a slightly higher risk item.

Ephedrine (ephedrine hydrochloride)

Description:

Epidedine is a simular drug that belongs to the group of medicines thrown as sympathonizedis. Specifically, it is to that an aphe and beta advecting aquesis (to numes these of neurophysical as a societies beta-2 aquesis). In addition, epidedine with the strengthysical production of the beta-2 aquesis, it is addition, epidedine with the strengthysical production of the beta-2 aquesis, it is addition, and the strengthysical distances of the strengthysical production of the compared at strengthysical production of the beta-2 and the strengthysical distances of the strengthysical production of the advection of the strengthysical production of the strengthysical production and reduction of the strengthysical distances of strengthysical production with this strengthysical production of the strengthysical production of the strengthysical with the strengthysical production of the strengthysical production of the strengthysical production of the strengthysical production of the strengthysical strengthysical production of the strengthysical production of the strengthysical production of the advectorized production of the strengthysical production and the strengthysical strengthysical production of the strengthysical production and the strengthysical strengthysical production of the strengthysical production and the strengthysical production of the strengthysical producting producting production of the stre the force of skeletal muscle contractions.

History:

Epidedice is a fairly dd medication, and hus been used in the United Status for a nurber of medical applications or the years is relating that of a similar, appelles approsents an orga-the-courter medicar, and saids are largely found in this appelles and prosent and orga-the-courter medicar, and saids are largely found in this access. Comosh oce can be used as a primary base for the manufacture of rethamphetamics. With ophetheir anabiles an over-the-courter product, underground medicarian of the phetheir anabiles and an over-the courter product, underground medicaries for OTC ophethrine products in a blue ond scoreing and murga states lines responded with legislicit according the said of preservor materials line (appledice) in 2006, a folderal line was possed further residence speculate ophethre may occu join the ki of folderally control dishatances. In spite of their resultions, and is appledice theory of the order dishatances in the plot of their resultions. The ophethre is a fiber over-theorem ends. In spite of their resultions, and is appledice for over-focuent ends.

How Supplied:

Ephedrine (as ephedrine hydrochloride or ephedrine sulfate) is most commonly supplied in tablets of 25 mg or 50 mg each.

Structural Characteristics:

Ephedrine is a sympathomimetic amine related in structure to amphetamine and methamphetamine. It has the chemical designation (1R_2S)-2-(methylamino)-1phenylpropan-1-ol.

Side Effects:

Ephedrine can produce a number of unwelcome side effects that the user should be aware of. For starters, the stimulant effect can produce shaky hands, tremors, sweating, rapid heartbeat, dizzness, and feelings of inner unrest. Often these effects subside as the user becomes more accustomed to the effect of this drug, or perhaps the dosage is lowered. In general, those negatively impacted by caffeine would probably not like the stronger effects of ephedrine. The mental and physical state produced by this drug is also quite similar to that seen with clenbuterol, so those who find little discomfort with that treatment should (presumably) be fine with this item (and vice versa). While taking this drug one may also endure a notable loss of appetite, usually a welcome effect when dieting. Ephedrine is in fact a popular ingredient in combination (prescription) appetite suppressants. The user may further notice headaches and an increase in blood pressure with regular use of ephedrine. Those suffering from thyroid dysfunctions, high blood pressure, or cardiac irregularities should also not be taking this drug, as it will certainly not mix well with such conditions. Ephedrine is a CNS stimulant with potential for fatal overdose. Signs of overdose may include rapid breathing, blood pressure irregularities, irregular heartbeat, unconsciousness, trembling, shaking, panic, extreme restlessness, and severe nausea, vomiting or diarrhea.

Administration:

The primup application for epicetica arroug bodybulkers and athletes (of-lichel) is that of a contraje (fick-solg and then, the infoldual unit generally lack the drag a few times per dynamic and the sole of th

Epicheire is also used by some competitive atthetss (relating powerfilers) as a similar before workstor. Competitions. The resulting (sight) steering and averyg increase may improve anamobic performance and weight tatak on major lifts. On this same note, it is also believed by some to powel as metrial dega manage the same more energies and better abuto concentrate on the tasks ahead. A pre-event dose of 25-50 mg of epidenties is typically used for this papers. It is importate it note that this comportatio is made continuously as a pre-workstor at pre-event similarity, as its effect will drainish as the body becomes a super-source of the same pre-event similarity of the same pre-transfer that the same pre-transf

Availability:

Ephedrine is widely available in the U.S. and in a number of countries abroad. It is not commonly a target of counterfeiting operations.

Meridia® (sibutramine hydrochloride)

Description:

Shuramine hydrochioride is a selective servorini and norademalin re-uptake inhibitor used for the medical management of obselying like planmaconcilas is istended to be an adjunct to a reduced calorie dist, which will help increase weight loss compared to that achieved with modifying food instead alone. Shuramine hydrochioridi is not advertised as a rapid acting drug, but instead one flat fasters slow, safe, and steady losses in fat msss which are minitationed long-term.

Sharamie hydrockiarkie eurst as weight-loss effect frought too distert meclurems. It hus a murked abity to propes appeils. During more studies, patient wood rackee their doil or early in take, babmier also stimulate the studies of the studies of the studies on calcier studie, sharing abattantian tendentian and appearing and their studies of early distantiane and the studies of early the studies and the studies the studies and the studies the studies and the studies and the studies and the studies the studies and the studies and the studies and the studies the studies and the studies and the studies and the studies the studies and the studies and the studies and the studies the studies and the studies and the studies and the studies the studies and the studies and the studies the studies and the studies and the studies and the studies the studies and the studies the studies and the studies and the studies the studies the studies and the studies the studies and the studies t

To get a better idea of exactly how well shutamine hydrochicrike works, we refer to some of the chicrical tasks on this sparCose messignion was conducted at the Kanssen Foundation to Chickel Plannencology in 2001. Here, a group of 222 closes patients were plane shutly c22% or plannen in the ophismen cose has 6 % so errors of the krist lower weight, while 12% noticed a 10% or greater loss in body weight. Shutamine was also associated whis significant improvements in some num religeorized and 11% closelstored lower, which was completed in Chicker and the start of the krist lower sources of the hydro start in the start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start was completed in Chicker and the start of the start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start of the shutle hydro start was completed in Chicker and the start of the shutle hydro start of the shutle hydro start start of the start of the shutle hydro start of the shutle hydro start of the shutle hydro start start of the start of the shutle hydro start of thydro start of the shutle hydro start of the shutle hydro start of

History:

Shuramine hydrochkride received Food and Drug Administration approval for sale as a prescription weight-loss agent in 1998. It was developed and marketed by Abbott Labonatories, which sold the drug on the U.S. market under the brand name Merklin. The company also sold the drug in many international markets under the name Reductil. Shuramine enjoyed only a limited period of salis in the U.S., sait was removed from the U.S. market was removed from the U.S. market was removed from the U.S. market was removed from the U.S. sait was removed from the same for the U.S. market was removed from the the U.S. market was removed from the the the tablet was removed from the the tablet was removed from the the tablet was removed from the table market in October 2010 under FDA pressure, citing an increased incidence of adverse cardiovascular events. Abbott has since also withdrawn sbutramine from many markets worldwide. Note that sbutramine remains classified as a schedule IV controlled substance in the United States.

How Supplied:

Sibutramine hydrochloride is most commonly supplied in capsules of 5 mg, 10 mg, and 15 mg.

Structural Characteristics:

Sibutramine hydrochloride is a centrally-acting serotonin-norepinephrine reuptake inhibitor structurally related to amphetamine. It is chemically a racemic mixture of (+) and (-) erantiomers of 1-(4-chlorophenyl)-N,N-dimethyl-a-(2- methylpropyl)-cyclobatamentetharamine.

Side Effects:

The most common side effect with sharamine is an increase in blood pressure, a truit duri contraindicates is use in partners with high-back parts and conference in states. Other common side effects include day month, skepkassens, intribully, back parts strunds because a constraint is the day of the structure in the structure of the structure in the because a constraint is the day. Structures because the structure is the structure introduction of the rarrow structure is because the structure is the structure constraint. Structure is the structure is the structure is the structure is the constraint is the structure structure is the structure of the structure structure charaksets, and learning, days the structure of and the structure of the structure structure structure is the structure is an integrated in the structure of the structure structure structure structure is the structure of the structure structure of the structure of the structure structure is an end to structure of the structure of the structure of the structure of the structure structure of the structure structure of the structure structure of the structure structure of the structure of the structure of the structure of the structure structure.

Administration:

Sharamine hydrochkrik is used for the management of obesity, itskuling weight loss and minieranzee, and sharado he used in coipcuriton with a reduced-cabier diet. This drug has been used with patients who have additional weight-reduced risk factors including corroted hypertension, disbest, and dopsdowine (high clockstarch). The recommended starting physical starting the starting of the starting of the starting of the starting of the 4 weeks it weight loss has not been sufficiently initiated. Higher doess are usually not recommended.

Availability:

This drug is subject to limited availability following an FDA supported recall in the United States, and subsequently its removal from many other international markets for safety reasons.

Zaditen® (ketotifen fumarate)

Description:

Kotöfin is narthäsmine drug thr i sued for the toment of general allergy symptom, central allergy confision (rachief or ognicarities), and the management of asthm. When used for anthm, the drug is not regarded as effective for truining an immediat attack (it is not a rapid borncholdane). Instead, over this to use is associated with a nebucion in the forquercy duration, and severity of attacks. It is usually preserved as a ways to mercause the effects of other astron metaciatoris. Likewise, kottefin finantea with usually supplement an existing astron metaciatoris. Likewise, kottefin finantea with usually supplement an existing astron metaciatorie. The source of the preserving of immediate rescue doices such an an another in the or relability. Account animum, moviews mergy symptom by toxicity memory in coupons, a properly that is common todus of the artifistance class. Its second and very using media of action, however, makes it useful in the treatment admits. Restellin financia it recreases the concentration of beta-dimensity inceptors in the bold (specicity blass). The bold (specicity blass) that strainade beta-2 neceptors are commonly preserved as broncholditors, used to increase air/low to the ingo and courter the concrition caused by sentem. While operating efficacions also, cone key therappente effect of Restoff and the sensitivity of the bold's dama of the beta narrie class.

The best-2 receptor upregulating properties of ketoffin finamate made this dueg of interest to the holyhaling and addies commanism. This is due to the storg of low the best-2 neoptor in supporting fit loss. Albough not a strong fit loss composed by locif, when taken with a best-2 agoinst intermogene like clotherols, levided in numeric may increase thermogene potency and noticeable, extend the window of active lipolyse. Clotherols and a distributed and the strandow of active lipolyse. The best here how the best-2 agoinst termogene like clotherols is distributed and other best-2 agoinst termody in the a clotherol is distributed and other best-2 agoinst termoly threat a limit of transformed term best assess bestinisting theory with such a ding a transformed by begins to diminish in effectiveness. Kenotifering world this tim period considerably.

The ability of kotofin to poterinie the effects of behz-2 agorist drags has been downstrated in a number of cirical states. For example, our study published in 1900 downstrated that when kotofin and clerbuterol were taken together, faree was a signifiant reaceus in them-denrogic receptor density of their quickly.⁶⁰⁰ Other studies with substantial (also related to garcials) as a shateon) showed that then a advecceptor downregation caused by long-term use of this beta-advencegic agart could be mptily reversel white in its a 2mg of clocofits finance per day.⁶⁰⁰

History:

Kotoffen was globalised as a prescription medication by Novaris, It is preserily prescribed for alongis, allegic conflox, and (more conrowly) the mangament of saftma in more than three-doorn coarties around the world. The more widely available heard mare is Novari's Zaften, which is odd throughout not parts of Enorge and Asia. In addition to genetic know of the medication downs of other branch mers; can be found in murdle to medica su well. Kotoffen finamatie is approached for ask in the Libra Status, but createdly The dosage of futurefilm finamatie is approached for all with Euler Markan Markan The dosage of futurefilm finaments in approached for all with Euler Markan The dosage of futurefilm finaments in the product is also to hear for it he to constrained used for any other of future for hear the more analy available of fordering finaments of the constrains, the drug is easily distorted for black market suk. As of row, it is not enternely popular with bodydanks and athless.

The UK galdens on the circul mangement of saftma consider leadoffs to be inflictive for the management of this disease. There is a shirthely confifting data on the potential usefalses of leadoff farmatic for this purpose, with some studies reporting positive results and others showing an impairMart effect. A thereaged network of the alp polisiked on the Coherne Dathusse of systematic Reviews in 2004 concluded that i appeared to have some usefalses in controlling saftma and wheacing in mury didlency. The the variability of the disease and response to the drag meant that these positive results could not be generalized for al advant onizer. ⁶⁰⁴

How Supplied:

Ketotifen fumarate is most commonly supplied in tablets of 1mg. This dosage is usually expressed in terms of the base, so each tablet actually contains 1.38mg of ketotifen fumarate.

Structural Characteristics:

Ketotilen farmante is selective histamine H1 antagonist, anti-allergie, and anti-asthmatic agent. It has the chemical designation +4-(1-Methyl-4-piperidylidene-4Hbenzo [4,5]sycholpeta[1,2-b]thjmene-1-(9H)-one farmarate.

Side Effects:

Common side effects include dry mouth, appetite stimulation, weight gain, dizziness, CNS

stimulation, and drowsiness. These side effects are all commonly associated with strong antihistamine compounds. In rare cases severe allergic reaction on the skin or a urinary bidder inflammation called cystiis may occur.

Administration:

When used to reduce the frequency, charting, and security of asfirm attacks, tactified finamate is stually initial at a doosg of The prober per day (Cpr (2014)). The costs regist may be increased to a maximum doosg of 2mg twice per day (Apg total). Bolyhakles and attacks will conversely use a doosg of The prober per day (2mg total). Bolyhakles and address will conversely use a doosg of The twice per day (2mg total). Bolyhakles and down at individue to bothin a strong theremosifies effect of the total both both 2mg attacks and the twice and the strong strength and the total both both 2mg attacks and the strength and the strength and the total both both 2mg attacks and the strength and the streng

Availability:

Ketotilen famarate is widely available, and is sold under namerous brand names in many countries. Large scale counterfiching of this medication is currently not known to be a problem.

Fat Loss Agents - Thyroid

Cytomel® (liothyronine sodium)

Description:

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Bodyhalkes and affetes are attracted to follynomic sodium for its ability to increase methodism and space the breakdown of body its. Most of other initialed during concess preparation or periods of "carting", the drug is usually said to significantly aid in the loss of find one olighter body calcular single during output of the significant state of the single stat

History:

The first medication that included T3 was technically a thropoid extract, first given to a patient with mynochem (as discorder associated with hynophynoxiliam i) 100^{1000} Natard lapoot extracts contained therapentically while levels of the thropoid hormons. T3 and T4, and were widely used in medical particle frame relation to low surt. In htt 1950, however, these damge shroly start giving any to new synthetic thropoid medications, namely further softman and the composition of the softman and the softman and the composition of the softman and the softman

significantly smaller portion of the global thyroid market. Given its more potent and last acting effect, however, licitlyroinie sodium remains a popular thyroid drug with bodybuilders and athletes. Cytomeli® is the most recognized trade name for the drug in the U.S., where it is presently sodil under the King Planmacenticals brand name.

How Supplied:

Liothyronine sodium is most commonly supplied in oral tablets of 5 mcg, 25 mcg, and 50 mcg,

Structural Characteristics:

Liothyronine sodium is a synthetic form of T3 thyroid hormone. It has the chemical designation l-tyrosine,o-(4- hydroxy-3-iodophenyl)-3,5-diiodo-,monosodium salt.

Warnings:

FDA copies the following black hox warning accorptop prescription folloptions sodium products soli in the Liss: "Thong with floor hormone activity, alone or together with other thampotic agents, have been used for the treatment of dessity. In enflyrout planters, does within the range of day hormonal requirements as a reflection for weight reduction. Larger within the range of day hormonal requirements are infections for weight reduction. Larger within the range of day hormonal requirements are infections with the solution of the solution in association with sympathemizetic amounts such as those used for the amorter effects:"

Side Effects:

Side effects are garenedly susceined with overdosaga, and may irckels bandsche, irribable, meroaness, sowialie, negalar bandberg, increased bowel moliky, or neutralie irregabritis. Overdosaga my also inkare shock, and my aggroute or tiggar angiu or congestie heart faise. Concel overgonose to folforotine oscilantu all produce synghesis neurally associated with hyperdproximon the overproduction of natural flyroid hormenses in the bolyThe concrease of overgonose-taked side effects is normally cause to immediately rokes or discorting therapy with follyroine sodiart. Acate massie overdose may be fib franctaning.

Administration:

When used to treat mild hypothytoidism, the bypical recommended starting dosage is 25 mcg daily. The daily dosage then may be increased by no more than 25 mcg every 1 to 2 weeks. The established maintenance dose is usually 25-75 mcg per day. Once a day administration of the full daily dose is usually recommended. Although lothytorine sodium is fast acting, is effects may persist in the body for several days after discontinuance.

The usual protocol arrong bodybräckers and athletes taking fordytorine sodiam to accelerate fit loss involves initiating is use with a dosage of 25 meg per day. This dosage may be increased by 25 meg every 4 to 7 days, usually reaching a maximum of no more than 75 meg edge. As an a medical setting, the inter of this slow baddhe is to belp the body become adjust to the increasing through hormone levek, and avoid sudden changes that may initiate side effects.

Cycles of foltytorine sodium usually last no longer than 6 weeks, and administration of the drug should not be haled abruptly. Instead, it is discontinued in the same slow mamer in which it was initiated. This usually emisit reducing the dosage by 25 mg every 4 to 7 days. This tapering is done so that the body has time to readjust is endogenous homeone production at the conclusion of threapy, and to avoid the corest of size effects.

Availability:

Lindpromis is an old and widely prescribed medication. It can be found readily in most areas of the world, and is sold in a variety of different brand and generic forms. Counterfoling is not a large-scale problem. It is important to note than one should never purchase an isjectable form of this drug. These are generally used as emergency mom products only, with notertially vey-dumprovs aids effects if misused.

Description:

Lecolopoies acdam is a synthetically mandfatted from of the matral thyroid hornese testicolopoies (r-1, -1). Thyroid horness: an permitry responsible for regaling the body's methodic rati, and play while oke in the story's utiliation of proteins fits, and which is characterized by isandificient tangen polection of thyroider the story mandfat for white a nurber of synthesis and polections of thyroider and the story method is deviced by isandificient tangen polections of thyroider and matrix mediation of the story mediation of the story of the mediation, and may take up to 4 to weeks before full futuresate is to she are needed in the consistent is be be standing from of the story of the

The action of levelsprovies sodium is very similar to that of the popular dynoit preparation (Cycontel) (indepresses sodium, (Cycontel) (sighty) different instrume, howevere, brieg synthetic from of the thyroid hermone risolodhyronine (T-3). A healthy individual with howe subfaces it levels to the T-3 and T-4 Hoyevard hermones present in their boly. T-3 is considered the primary active form of thyroid hermones, present in their boly. T-3 is considered in the other than a strong the factor of the theory of the theory engelind a law bring and effect that is coupled for times stronger the that of T-4 or as miligrams be-onligram basis. Llawsise, Cytornell's is considered to her more poter form of theory in character of the theory and size differed poterial.

Lecolitorize acdian is valued by mmy drag-stag athletis and bodybalkers for it a abily to similate the metabolic net and support the bounddown of body the stores. It is usually taken during a period of calorie netroticing ("tuting"), when the infidulati is focused on far its source in the store of the store of the store of the store of the store support fit focus at a higher level of calorie inside that musc and otherwise he possible without the drags, adding to also previously also among the communities. A starbals musc has boosting effect of these drags in the produce take and the starbal musc has the boosting effect of these drags musc produce fister gains in marker mass. The drags, bodyne of the starbal musc with to be widely movement as contrast of the support.

History:

Lecohynovic acdam was the first synthetic flyorid medication to be soli in the U.S., and was first introduced to model in 1955 by Filt Laboratorias as Synthetical. The day has a long history of thenpoteic use in the U.S. and intermationally, and for decades has been the most videly presendent medication for the treatment of Hypothypothy. The Synthetic Harman has historically been the most accessful, while figures estimating that it retained SPs's of taid the interview of the synthesis and the synthesis and the synthesis and the synthesis and the interview of the synthesis and the synthesis and the synthesis and the synthesis and the interview of the synthesis and the synthesis is washer and shows arising affekts more to be table to the produce and the shower attring affekts more than the synthesis and the synth

The Synthoic head seel'has a long and a times contoursail history⁶⁶ for mmy years after is necroin to piler Laboratories, Sprithoni enjoyd a viral monopoly on the levoltynoise sodiam market. Generic medications finally began taking a large share of beylymosies notime and going in the 10 MiNo. In response, First Laboratories finded a much a linker thready of calls in the 10 MiNo. In response, First Laboratories and each of the 10 Mino and the laboration of the laboration to the study was considered in 10 Mino. The source of the laboration of the laboration of the study was piled in the context of the laboration of the study was supercold optical in the context country in the laboration of the study was supercold optical in the context country in the laboration of the study was supercold optical in the scenario optical in the laboration of the study was supercold optical in the scenario optical in the specific scenario optical in the scenario optical scenario optical in the laboration of the study was supercold optical in the scenario optical in an annihistory counting was the absorb optical in the scenario optical

How Supplied:

Levothyroxine sodium is most commonly supplied in oral tablets of 25 mcg.50 mcg.75 mcg. 100 mch, 125 mcg. 150 mcg. 200 mcg, and 300 mcg.

Structural Characteristics:

Levothyroxine sodium is a synthetic form of T4 thyroid hormone. It has the chemical designation L-3,3',5,5' tetraiodothyronine sodium salt.

Warnings:

FDA requires that the following black how summing accompany prescription indiportion solution products solution in the U.S. "Through hormones, including locardynomic acdium, either alone or with other therapeatic agents, should not be used for the treatment of obesity or for weight loss. In editorylo patients, does so within the range of oblack profits protein the solution of the solution of the solution of the solution of the solution solution of the solution of solution of the solution

Side Effects:

Side clicks are generally sosciented with overdosaga, and may include bandhen, imitability, merooaness, vossing in grapha brahterski, increased lowed moliky, or mestratia irregaleritis. Overdosaga my also indue skock, and my aggroute or tiggy angiu or congestie heart failsen. Cronici overcoposer to lovelymour sostimi will produce symptoms normally suscitated with hyperdynolising or the overproduction of tunnal thyoid hormons in the botty. The occurrence of over-possive data data defects is normally cases to immittable rockner or discorting thrapy with lovelsproxies sodium Acate mussies ourdones my he Bie Threatoning.

Administration:

When used to treat mål to mederate hypothyvoitism, the average replacement dose of levofhyroxine sodiam is approximately 1.7 mcgkgday. This equates to 100-125 mcg/day per day for a 154b adult. The full therapeutic dose may be given form the ones of otherarys in otherwise healthy adult patients. Note that due to the long half-life of levothyroxine, the peak therapeutic effect at a given dose may not be achieved for 4 to 6 weeks.

When used (off-label) to accelerate fit has by bodybalkes and athkets, the typical protocol invokes who hadh off the dosign is of inthe bodyh same piller me to adjust to the changing thypoth hermone levels. An athividant will generally start with a low dosign of 25-50 mg and vill show increases the moural "25-50 mg cached vill so vites. The final dosign will usually be in the range of 10-150 mg, and will arealy exceed 220 mg. It is important to contain extrameter of the piper limit will be seen to the second start will be ather than the Cantons individual will be sare ret to use excession amount of levelsproties adding to Synthesis the same ret to use excession amount of levelsproties adding to Synthesis the same ret to use excession at mouth of the same shows the Synthesis the same ret to use excession of each cycle. The house has any piper by the same shows the same ret of the same shows the same shows the same shows the same shows the same ret of levelsproties shows the same shows the same that due to the skow acting nature of levelsproties shows the same shows the shows of the same shows the bit here influence for the same shows the same shows the shows of the same shows the same ret levelsproties shows the same shows the same shows the shows of the same shows the same shows the same shows the same shows the same of the same shows the same sho

Availability:

Although levothyroxine sodium is a widely manufactured drug, it is not as common on the black market as the stronger thyroid drug Cytomel®. Large scale counterfeiting does not appear to be a problem.

Fatloss Agents - Other

Description:

DNP is one of the most contournail drugs inue by bolyhalden. This spart is not ould for human use anywhere in the world at this time, but is readily available as an itakatila chemical. Among other things, it is used as an itemediary for the production of certain does, for photospite chevolenent, as a fingical, wood pressure-teament to percent roling and as an spectricle. It is technically classified as a poior, Albrugh quie incorpanses with this is of storing industriclemental uses, it is chemical ways additioning the errol pattern medicine as a did drug for humans. It is this property of dinitophenol that remains of netroest to some bolyhaldense today.

Drintophenia induces weight tass by unccoping oxidaties phosphorylation, thereby markedly increasing the methodic near all body temperature. While this an anothemy diction way of producing mpil weight loss, there scents to be to colling to DNP's temperature invessing effect. Herenis is perhaps in stored dangenous trait, run yailow bed temperature to rise to level that can be damping, even faul. What Carl Matherey made perhaps one of the cacing and not flamming, even faul. What Carl Matherey made perhaps one of the cacing and duchts associated with DNP have continued over the ducatis. For example, a near high publication skyrocenes are mute tide of a long latent. W'n i 200 direct tasking. The ducatis associated with DNP have continued over the ducatis. For example, a two rise jub products days concerns are mute tide of a long latent. W'n i 200 direct tasking. DNP for only for days. The dose used was reported to be 600 mg per day, just three 200 mg capasite.

History:

The fields properties of DNP were reportedly first molecul daring World Wer 1, when overvigit nen working with DNP in matching paths started Joines substartial mourse of weight. If dat not take very leng for this chemical to be kientified as the cance. Soon after, it was packaged as a darge moders. By 1935, we meet have 1940 versa the first synthetic drug that was error used for weight relations in this courset. Whit is was acadimately a large that the startest models was an end of the startest startest and the startest was error used for weight relations in this coursely. Whit is was acadimately, two heavy weight was error used for weight relations in this coursely. Whit is was acadimately, two heavy weight in the the Distribution of the startest path mouses all acous the courts.

While the drug may have worked for the istended pappone, it was also introduced of a time block governmet review and approved of drug adds, in the required DPH lad some varstning shortcomings, and it data's take leng for reports of side effects to began property in the drug shortcoming start is drug for reports of side effects to began property in the drug. Narrence reports of DPA⁺ finde contexts began compared in the side of the drug. Narrence reports of DPA⁺ finde contexts began compared in the side in the drug. Narrence reports of DPA⁺ finde contexts began compared in the side of the drug is the drug attribution of the side of the drug was soon palled. By 1558 is used of the size height attribution of the drug was soon palled. By 1558 is used of the superstort double material sources of the drug was soon palled. By 1558 is used of the superstort double material sources and the drug was soon palled. By 1558 is used of the superstort double material sources and the drug was soon palled. By 1558 is used of the superstort double material sources and the drug was soon palled. By 1558 is used of the superstort double material sources and the drug was soon palled. By 1558 is used of the superstort double material sources are solved with the sources and the drug was soon palled.

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How Supplied:

DNP is not supplied in a form prepared for human or veterinary consumption. It is available

as a research or industrial chemical only.

Structural Characteristics:

DNP (2,4-Dinitrophenol) is a cellular metabolic poison with the chemical designation 1hydroxy-2,4-dinitrobenzene.

Side Effects:

There are many potential side effects associated with DNP use including increased heart rate, increased breathing rate, rausea, elevated body temperature, insomnia, profise sweating, rash, skin leisons, decreased white blood cell count, cataracts, coma, and death.

Administration:

DNP is not approved for use in harms. Proceeding galaxies are unrealistic. A common done used armong boolphanks is a reported 20-ray per kg of bodyweight per day. This calculates to a dosage of 200 mp per day for a person of approximally 220 ports of oblowing the volter time perplanet tracks to treat in more kan muscle musc than the download structure and the dosage of 200 mp per days for a person of approximately 220 ports of the download structure and the dosage of 200 mp per days for a low structure and the dosage of 200 mp per days and the days and the dosage of 200 mp per days and the days

Availability:

DNP is not available as a human or veterinary medication in any part of the world. Availability of products intended for human use is entirely in the underground realm, where products, dosages, and safety are not the subject of government approval.

Lipostabil N (phosphatidylcholine/sodium deoxycholate)

Description:

Lipostable N is an injectable medication that cortains phopulatifyklythine (PPC), a nutral phopholpidi, Sdothni dowycholus (c) his sin a ho added (more golder signification) to subhitie PPC in states. It was originally developed as an intravenous solution for the improvement of storming hist, relation of atterning hasing, improvide golder and the presention of storming the values, and the prevention or interast of blood weard blockages by the particles (the embodies). It is approved as an interast of blood weard blockages by the particles (the embodies). It is also had a copputation of blood interast or interast of the storm of the store of the st

The mechanism behind Lipostabl's lipolytic (fit loss promoting) effect is uring Lipon rejection, the soliton at as a ad etergraft, a canage nonspectific by liveal/adown (of conmerhennes,⁴⁰⁰). The bits all sodium dooxycholice is actually believed to play an important robot here, and is therefore considered at a nucleic constitutor of Lipolation ket. Lipolation and the set of the soliton of Lipolatice acid. The soliton of Lipolatice ket and the soliton of Lipolatice ket. The soliton of Lipolatice ket al. The soliton of Lipolatic ket al. The soliton of Lipolaticity here is real horizon for the soliton of Lipolaticity here is related for large fit for lease of Tipolaticity here is related here again the soliton of Lipolaticity here is related here gains and the soliton of Lipolaticity here is the distribution of Lipolaticity here is the lipolaticity here is the lipolaticity here is related here again the soliton of Lipolaticity here the distribution of Lipolaticity here is the lipolatic here and the distribution of Lipolatic here and the lipolaticity here the soliton of Lipolatic here and the lipolaticity here and the lipolatic here and th

History:

Lipostabilities appeared as a medication during the 1990s. Although not approved for prescription use in the US, it is approved for medical use in a number of other contribution of the transmission of the medication of a number of the contribuion of the star of the star of the medication, although it is also found as Lipostability from a rule as Lino star docts on bolene that uses given the first loss agreement of the star of the library of the star of

Network Lapobja (an organization of some 350 doctors workshole that supports this use of Laposhish protors over 18000 constrict cumturet without targeted advesse events. Additionally, E. Hassengelssendhare, the medical doctors of the Austana class. Therease the advector of the Austana class of the Austana class. The advector of the stress values at the solution costs (David advector) and the mendelings in stress values. Although we can be a grant datal of data and the stress values at Albadya we can be are agreed and of data with the advector of the stress values at Albadya we can be a grant datal of data on the off-bala loss of Lapositis, but can be faust is agreenily very positis, can stagges this ding or matal paperkis, liped grant and Sorth Arias.

How Supplied:

Lipostabil is most commonly supplied in injectable ampules containing 5 mL of solution each.

Structural Characteristics:

The primary active ingredients in Lipostabil are phosphatidylcholine (phosphatidyl-Ntrimethylethanolamine) and sodium deoxycholate (cholan-24-oic acid, 3,12-dihydroxy-, monosodium salt).

Side Effects:

Potential side effects associated with subcutancous Lipostabil injections include localized swelling, redness of the skin, burning, pain, tendemess, and bruising. Systemic side effects are reported in approximately 3% of users and may include diarrhea, nausea, dizziness, and intermenstrual bleeding.⁷⁰¹

Administration:

The typical practice for using this drug to promote localized fat loss involves a series of solucitances injections. A total dosage of 1250-2500 mg is often used, which equates to 25-50 mL of injectable solution. This dosage is divided into 20 or more separate smaller injections. These are spaced frequed the problematic area (quide commonly the abdominks or thighs), and are al given during the same office visit or application period. The drug is not administered on a duily basis.

Lpostabilizations and cases a significant amount of futuremation in the area, which may take a week or larges to fish systekic. When their information does subside, however, it usually uncels a noticeable amount of fit loss, in a chiral setting, fits procedure is often repeated a low trace, so as to early the near and achieve the described level of fit neticines. The correst gathenes set forth by Network, Epolysics and for an 8-week breach theorem of the correst at match how of days and the set of the set of the set of the fit of the correst at match how of days and the set of the set of the set of the fit of the correst at match how of days in our liquing the fit and gatheness find 1 simply to painful to use, while oftens seem to tekrate the whele procedure enternely well.

As for the ultimate question of how well it works, it is difficult to give exact numbers, as few clinical studies have been conducted on this use of the drug. The anecdotal feedback is mixed. Many people who try it report positive results, particularly for the removal of those has stabilized and the stabilized of the stabilized of the state of the stabilized of the stabilized of the state of the s

Availability:

Lipostabil is not a controlled drug in the U.S. or Europe, and as such is fairly easy to obtain on the black market or via mail order drug distributors. "Mesotherapy," clinics selling procedures with Lipostabil are also fairly common.

Growth Hormones & Related

Geref® (sermorelin acetate)

Description:

Semecini is a synthetic analog of endogenous growth hormon-releasing hormone (GHPI or GHP). Semecinity is a portion of the synthypoid hormone, specifically consisting of the first 20 of 84 a mino acid attracture. As its mare status very clarkly, the biokgial activity of GHPI is to stimulate the synthesis of growth hormone, which occurs in the pathong galac. Statisch hore shown, however, this the GHPI peptide can be pathy unstated subtor assume the first and the synthesis of growth the synthypoint and the synthypoint sources the first and an endogeneous status and the synthypoint production of growth hormone. Blaced on its structure and action, semered in a classified as a growthermore relating the classifier of CHP endogeneous structure and action, semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action, semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action, semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier as a growthermore relating the classifier of CHP endogeneous structure and action semered in a classifier as growthermore relating the classifier of CHP endogene

Semeeting as an acctue salt, is used in chical medicine for two primary applications. The first is o diagnee primary dedicinc: The proceeding model medicine in the sememation location. The properties of the primary semematical semetal semematical semander semematical semimatical semematical semimatical se

Addets are interested in semmerica actuate for the same reasons they use recombinar thrum growth horness. Armong other things, growth horness has malook and anicatabole properties. An elevated GH level may support new mache itsue growth, and also endrace strength, energy levels, and corrective tosass. GH is also apport modulator off hornes are using hornessing to the same strength and the source of the same widely accepted by the holphaling and competing and the same that shorts listency of ase, growth horness neighborhaling and competing acceptate the same strength and provide same strength and the same strength and the same strength and strength and the same strength and the same strength and the same growth horness neighborhan is labole provider in an sing apportise moders again serves as an adventure to growth horness neighborhand and GH dekeneary ("soundepusae").

When comparing this drug to recombinant human growth hormore, semmetin acetate does appear to be less efficient under normal therapeutic conditions. During clinical statules, fewer patients on average seem to respond floroably to therapy in contrast to somatorpia, and for those that do respond the improvements are often less pronounced.²⁰⁶ Sill, it would be a method to environ service as a visible theraperior contrast to average therein. tensors or extrate structure account and a trute manapeut, sprat i ver temping, simular have bard that structures in significant i recenses in height velocity in children with GH deficiency.²⁰⁰ Furthermore, these improvements scents to be well satisfated affer one year of thermapy.²⁰⁰³ Arthologies to IGHH GA dowley in some patients during extended therapy, and may impair the potency of the during.²⁰⁰ The full biological relevance of this artholy reaction, however, remins unclear.

Semicodii is approved in he United States for the treatment of GH deficiency in offshere ordy. Produced statics tearing a dash with somtpaneae are lacking. One innotigation looked at the effects of 2 mg semicelin action and 2 mg and 2 mg

Another investigation booled at the effects of sermerorin accetta in a group of HIV+ men with phosphotophys¹. HIV hoppolystophys rules to be characterized by the abnormal distribution of at colk and appressed beside of growth hormore. During this insestigation, 31 significant boxed duply of 12 works. The primary growther of the aday was a significant necrossion is seemed GF-1 in the sermerodin accente group (104 right-L we forgat). This was accorrelated by a structure of the second se

Semercin acctata codi be viewed as offering a therapentic advantage over recombinant growthermore in some cases, in that it is back by to eash if of the eases. This is due to be fact that it relies on the body's own hormore synthesis instead of exceptions applementation. Thus, normal IG-F1 is factors it this is also by observation that the due to the growth hormore stimulating effect.⁷¹³ As such, hormore levels are more easily controld with semeroid next their normal controls, while semeroid in produce significant elevations in G1 and IGF-1, these levels should ned exceed the high end of the normal methods. These methods are body and the semeration of the normal methods in the body for the semeration of the semi-factor of the normal methods methods. bondies over a support this accord while method with semi-factor of the normal methods. bondies or compared the size of the factor over a support of the semi-factor over a support of the semi-fac

History:

Semuroin acctate was developed dring the early 1980s, and approved for prescription said by the U.S. Food and Dax Administration in 1997. It was intraduced to market under the band nume Geref Diagnostic by the international biotechnologies firm Serono. As the mure implics, I was primarily developed as a diagnostic tool. It was specifically used for evaluating potential plating deficiency in GP production. Genus is effect on growth hormore levels, however, the dorg was also approved by the FDA for the treatment of GI deficiency in children. Genef was never which prescribed, however, expectially for user interim domained placominate the product is OLAFO and the same of GI deficiency hormorealized and the Product is OLAFO and the same of the same phenomenetical approleted. ¹⁷⁵ Hus was the early compary manufacturing servership as a comportanted neuroscience and the Used S2asci, the is in horger assistable as a devel famoury learn the action material servership accettar is still made by at last one knowed stargeling.

How Supplied:

Geref Diagnostic was supplied in amptals: containing dry hyphilized sermorelin acetate, equivalent to 50 mcg of sermorelin. This was reconstituted with a sterile diatert (also supplied) before use. Generic compounded versions of this medication typically contain between 3.0 and 7.5 mg of dry hophilized sermorelin acetate in a multi-dose vial. Reconstitution before use is also required.

Structural Characteristics:

Sermorelin acetate is the acetate salt of a synthetic 29-amino acid peptide (GRF 1-29 NH 2) that corresponds to the amino-terminal segment of the naturally occurring human growth hormone-releasing hormone (GHRH) with 44 amino acid residues.

Warnings:

Sermorelin acetate should be used with care in epileptic patients. Obesity, uncontrolled hypothyrodism, hypergycenia, or elevated plasma fatty acids may impair the effectiveness of sermorelin. Therapy should be discontinued in patients treated for childhood GH deficiency once the epiphyses have closed.

Side Effects:

The most common side effects to sermorelin acetate therapy are injection site reactions such as pair, redness, and swelfing. During chiral triak, this occurred in approximately 17% of patients. Less common side effects include difficulty swallowing itching dizziness, lhshing headache, musea, vomiting, altered sense of taste, restlessness, and skeptiness.

Administration:

When used medically for the treatment of idiopathic growth hermone deficiency in prophestical alishes with growth failer, semerine accetes is administened by substanceous rejection at dosage of 0.03 mger kg of body weight core a day at bodime. Histoire of the shead her to restate it account irritation or the badking of race mices. When used to evaluate administened in the meeting ather an overright fait. This is followed by 60-120 minutes of periodic blood sampling to measure philarly hormere output.

When used for physicase or performance-entracing proposes, semicenfa acette is given by sistemators is pictor. It is typically administent al a todage of 0.2 to 0.5 mg per day (200-500 meg), which is given before skeps. Statiles, however, do suggest that the drug is more efficient when performs the day.¹⁰¹⁰ meters is, a today meters of notice the test day do says into two applications, one is the meming and one in the evening Cycles of someratin acetual analysis. In the events and the test of the statistical performance of works are not an extension. The pin is that the extended maintenance of youlfild growth homous levels why given as given the polysis and analysis.

Availability Trends:

Given its low financial value on global pharmaceutical markets, sermorelin acetate is subject to limited availability worldwide. The preparations must commonly found diverted for bodybuilding use include those mude by private compounding pharmacies in the United States, and those sold by gary market research chemical supply companies.

Human Growth Hormone (somatropin)

Description:

As is more suggests, human growth hermore is an important mediator of the human growth process. This hormers is produced endogramously by the materies planking galact, and can be at especially high bests during childrons. It is growth promoting effects are band, and can be provide, endophysically and the strength of the strength of the strength of the product of the strength of the strength of the strength of the strength of the product of the strength of the strength of the strength of the strength of the product of the strength of the

(rhGH) is biologically equivalent to human growth hormone (hGH) of pituitary origin.

In a model setting, contribupits used to help treat a strative of health conditions, I is most modely presched in cases of childhood growth disorders that are characterized by israficient growth homose production. While usually not fully concrede, sometropic use is often capable of substantial processing the times growth rate and worked heye there further growth is halted in addedecence. This medication is also used to acceltate growth in childhoot the stration of the strationary of the treat o

Sommony is also sometrines preservable to healthy mean and women who are aging (Growth hormore levels tend to deche as we use priod level, and murey physicism beivee that its supplementation to more youthful levels can help show some of the damage of aging. Growth is brocked an atcholice effects on maceds mass, strength, energy, cell expansion, and all basis, there are many supporters of this use, even if KeH may not specifically related the aging growth and the strength of the strength of the strength of the strength of mean maximum strength of the strength of the strength of the strength of specifically aligned individues (and the strength of t

Sommony may be given by either subcataneous or intramuscular injection. Daring clicial studies, the plannovskicker poperties of controlopic fallowing both methods of use were determined. When given by advectances injection, sommony history fallowing the fighter best of theoremather (175 vs. 407). In teas of our guardiabien fillowing both and 49 hours after advectances and intramescular injection. Specific theory of the source of the strength of the theoremather of the strength of the strength of the strength of the strength of the theoremather of the strength of the strength of the strength of the strength of the theoremather of the strength of the strength of the strength of the planet method of using control strength of the strength of the planet of the strength of the strength of the strength of the planet of the strength of the strength of the strength of the planet of the strength of the planet of the strength of the strength of the strength of the planet of the strength of the planet of the strength of the strength of the strength of the planet of the strength of the strength of the strength of the strength of the planet of the strength of the planet of the strength of the strength of the strength of the planet of the strength of the planet of the strength of the planet of the strength of the planet of the strength of the strength

A specific analysis of commorpin-strip shows a horners with a discress set of effects. It is analysis to skeatly mack, shows to accrease both the size and marker of cole (processes referred to as hysteriophy and hyperplasis, respectively). The hornerse also seems to huse growth-provering effects on adrophynd the blocy eacking the cys-size adribuits. Stringer hus a disbedgrarie effect on cardophate metabolism, which means that it causes blood samplers how the frequency strength space strength spac

Gowth hormene his both direct and indirect effects. On the direct side, the ICH protein attaches to receptors in marcle, how, and adipose biasses, stretty message to support ambolism and lipojisi (fit loss). (atomth hormere also directly increases glacose synthesis (glacoroganess) in the loce, and indices rule mission they blocking is activity in traject cells. The indirect effects of growth hormere are largely mediated by (GF-1 (readi-like growth factor), which is produced in the fitter and vitualy al dore thatsain is response to growth factor), which is produced in the fitter and vitualy al dore that suggests to growth iterative the strength of the strength is and instances in disponse. The growth memory activity (FG-1, however, also last cells that as morphy attegrations to growth memory lastice). The syncregistic and integrations directly of these two hormers orders to fitter the support [polysis, increased serum glacose locek, and reduced readimensentia.

Somatropin is considered to be a controversial anabolic and performance-enhancing drug in the realm of bodybuilding and athletics. The main issue of debate is the exact keel of potenta reset tins statistice carries, vane states was rav- pattern a vasatig une tend to a sport potentially storeg analysis can artistatistic properiors, states domontaring these same effects in healthy adults and athletes are lacking. Darag the 1990s, a lang body of miles marcaled discussions and fold in healthy adults provide the store of the theaght to be the most powerfal analysis adults and radii polarities. Most rape researting advances in the store market and the store of the store of the store of the market have tend to agree that it is the fit-loss-pomoring properties of somethyme in individuals nov tend to agree that it is the fit-loss-pomoring properties of the store of the trade of the store of the trade of the store of the trade of the store of the trade of the store of the trade of the store of the trade of the store of

History:

The first harms growth hormoze preparations to be used in medicise were much form phaney entrasts of hummon origin. These are nove controlly effected to as calabor growth hormoze preparations. Approximately 1 mg off.201 for 1 day doop; coid he obtained from 1000 MeV. The second results of the second results are related to the second results of the second resecond results of the second results of the second results

The FDA approved the first synthetic harma growth hormore drug 10/85. Synthesis produced a pare bornear without biological contraintion, climitarithe prosobility of CDI transmission. The drug approved was called somatern (Proteopini, and was based on a munificative glowthout of calabox CII by the FDA that same year. This hormone, the name hormone is a single strategies and the second second second second production (product hyperball) and the second second second second production (program second second second second second second second production (program second second second second second second second second growthe second second second second second second second second second growthe second sec

How Supplied:

Somatropin is most commonly supplied in multi-dose visis containing a white hophilized powder that requires reconstitution with sterile or bacteriostatic water before use. Dosage mny vary widely from Ing to 24mg or more per visi. Somatropin is also available as a stabile pre-nrized solution (Nutropin AQ) that is biologically equivalent to reconstituted somatropin.

Structural Characteristics:

Somatropin is human growth hormone protein manufactured by recombinant DNA technology. It has 191 amino acid residues and a molecular weight of 22,125 dahons. It is identical in structure to human growth hormone of philatary origin.

Storage:

Do not freeze. Follow package insert for storage information. Refrigeration (2° to 8°C, 35° to 46° F) may be required before and after reconstitution.

Side Effects (General):

The most common adverse reactions to scentropht hergy are joir pair, headded, a fields symptons, perplora elem (user reaction), and hock pair. Less common adverse reactions include influencien of mesors mentiones in the new (thinki), diziness, upper general elem, mesors, sone benes, ergnl harded symptons, paired general elem, mesors, sone benes, ergnl harded symptons, ergnl accompair, lo valde hickening of the bosons, most nealdy the feet, brank depression and elsowish, and enlargenet of the internal organs. Due the ground promotion effects of an elsowish, and enlargenet of the internal organs. Due to the ground promotion effects of an elsowish, and enlargenet to the internal organs. Due to the ground promotion effects of the output of the court of the strengt organs. The solute of the strengt or unearce.

Side Effects (Impaired glucose tolerance):

Somatropin may reduce sensitivity to insulin and raise blood sugar levels. This may occur in individuals without preexisting diabetes or impaired glucose tolerance.

Side Effects (Injection site):

The subcutaneous administration of somatropin may cause redness, itching, or lamps at the site of injection. It may also cause a localized decrease of adipose tissue, which may be compounded by the repeated administration at the same site of injection.

Administration:

Somatopis is designed for substancess or intramscular administration. One mligram of somatopis is equivalent to approximately 3 International Unics (11). When used to treat adult orset growth hormone deficiency, the drug is conrrowly applied at a dosage of OSymtype per day to 0.01mpkg per day. This equates to modph 11 U o 3 U per day for person of approximately 180-220 hs. A long-term maintenance dosage is stablished after reviewing the patient's 167-1 loved and clinical response over fins.

When used for physique- or performance-erhancing purposes, somatopin is usually administered at a dosing between 11 und 6 10 µcr dty (2-41 Ubeing most common, The drug is commonly cycled in a similar manner to anabolic/indrogenis strexisk, with the kraph of make garanally being between 6 weeks and 24 weeks. The anabolic effects of this drug are less apparent than its lipo/site (fit loss) properties, and generally take longer periods of time and lingth ocuses to manifest themselves.

Other dags are commonly used in conjunction with normappin noder to teke a stronger response. Thysical discussion, 2013 are presented in the lowers discribed scenariospin on flycial backs, and may significantly entrance fit has during theory limits is how the discriminant of the discriminant of the discriminant of the lower discribed of GF binding proteins -1, allowing for more GF-1 sets($n^{2/2}$) area the binding more set of lowers GF binding proteins -1, allowing for more GF-1 sets($n^{2/2}$) area the binding more set of also howers GF binding proteins -1. The discriminant of the discrest are discriminant. The discriminant of th

Availability:

Somatropin is produced by many different drug companies, and is distributed in virtually all developed countries. The most common brand names include Serostim (Serono), Saizen (Serono), Hamatrope (Eli Lilly), Nordiropin (novo nodisk), Omnirope (Sandoz), and Genotropin (Planumcia).

Somatropin products are high value targets for drug counterleiting operations. Many counterfeits are highly deceptive in nature, and have been found in both flict and legitimate drug distribution channels. Some counterfeit growth hommone products are made by nebholino visik of I/CG which have a very close visual resemblance to somatromic a home or abholino visual resemblance to somatromic a home. presence the standard standar

Increlex® (mecasermin)

Description:

Measemin is humn inside-like growth factor-1 (GF-1) menufactured by recombinar DNA herbodys (rdf-1). IGF-1 is the simum mediator of the growth provides (rdfof humn growth hermex. As stack, measemin also can simulate the growth of boxe, measing a statisfiele (rdf-simulation) and the state of the state of the state growth and the state of hermiteenergy of the state of the state global and they state is a state of the state of the state of the state global and they state is a state of the state of the state of the state global and the state of the state of the state of the state state global and the state of the state of the state of the state of the state global and the state of the state of the state of the state of the state global and the state of the state of the state of the state of the state state of the state state of the state state of the state state of the state state of the state state of the state state of the stat

Measurem is most commonly prescribed for the treatment of sevene primary [GF-1] deficiency (Primary [GF-1]). This disease is characterized by a histore to prodece remain levels of (R7-1) due to isonifications is in the growth hormore /(GF-1) axis (usably involving GF (neceptor, signing) approximations, (GF-1) and (GF-1) due to the observations of the observations of the observation o

History:

The U.S. Food and Drag Administration approved Meesemin in Augast 2005. It is sold under the brand mem Incredes, munificationed by Breiscia Inc. of Boshner, Zalförni, Tercica Increases this technology from Genericch, which was the first compary to sell a synthesiaji munification larman growth homemo product in the Indea States (Protorphin) Tercica's MGZ-1 is produced by a similar recombinum DNA technology. The processmotion graves meeting are business (Ford 1996) and the Company memory and the similar state of the similar technology of the processnotes in the similar produced by a similar recombinum DNA technology. The processmetry of the similar technology of technology of the similar technology of technol

How Supplied:

Mecasermin (Increlex) is supplied in 4mL multi-dose vials containing 10 mg/mL.

Structural Characteristics:

Mecasermin is human IGF-1 protein manufactured by recombinant DNA technology. It consists of a string of 70 amino acids and has a molecular weight of 7,649 daltons. Its amino acid sequence is identical to that of endogenous human IGF-1.

Storage:

Do not freeze. Refrigeration (2° to 8°C, 35° to 46° F) required before and after reconstitution.

Side Effects (Hypoglycemia);

The mot common adverse reaction to measuremin therapy is hypothecutia, which occurred on at last one concession in 42% of partices receiving the during crisical traits. Approximately 7% of partners moticed sevene hypothecutia, and 5%, notecid hypothecutia senter or loss of concentross. Signs of rule 10 modernet hypothecutia include human, dowatenes, blanned vision, depressive mood, dizenses, sweating, pubpation, terrare, needsnesses, tating of an her hards, feed, for a traject applicableness, inabidity at home blanking, manada more and an adversarial space and an adversarial particular setting and the setting and the setting and the setting and concentrate, headneds, sikep distaturaces, mesky, started speech, initiality, athormal blanking, manada more and an adversarial speech and the setting and condy have a carabidpatter afrik. Signs of source hypothecemia include discrimintane, sistem, and transcensouses. Source hypothecemia include discrimintane, sintensi, and transcensouses. Source hypothecemia include discrimintane, sistem, and transcensouses. Source hypothecemia include discrimintane, sistem and transcensouses. Source hypothecemia include discrimintane, sistema and transcensouses. Source hypothecemia include discrimintane, sistema and an expression of hypothecemia in source cases the symptome of hypothecemia in the discrimintane. since the sitema include discrimentane.

Mecasermin should never be taken before sleep or in higher than recommended doses. A meal or snack must be consumed within 20 minutes (before or after) of administration.

Side Effects (Injection site):

The subcutaneous administration of mecasermin may cause bruising at the site of injection. It may also cause a localized increase of adipose tissue, which may be compounded by the repeated administration at the same site of injection. Rotation of the injection sites is recommended.

Side Effects (General):

Other potential adverse reactions to measuremin theory related joint pain, growth of the trouchs, storing, handback, dizzbesc, convolutions, routing, and pain, henring loss, and hypotrephypi of the flymma gland. Mild elouitons in secura XET, ALT, and LDH levels were blocker and the storing of the relation of the store of the store of the store of the decime run afformation. Beautions in secura XET, and LDH levels were blocker and the store of the relation of the store of the store of the remained within the sports of intral organs. Kainay and stores also observed, but remained within the store of the store of the stores of the store of the stores that the store of the stores of the stores and tradition of the stores of the store of the stores of th

Administration:

Measurem is siterabed for subcannous administration. The initiation of thempy involves to does monitoring of bodo glacose levels util an poper maintenent does is stabilished. The recommended starting does is Δt to δB mg/kg effects (reading) and market gravity of the stabilished of the market starting does is Δt to δB mg/kg effects and Δt and Δt to δB mg/kg effects and Δt and Δt to δB mg/kg effects and Δt and Δt to δB mg/kg effects and Δt m

Mecasemin is not widely used for physique- or performance-entancing purposes. Common protocols of administration have not yet been established. Due to the potential for severe hypodynemin, maximum doses amough podybuliers and athletes are not likely to messarable weeked those supplied by thempentic guidelines. This drug will most likely by taken in cycles listing no longer than 8-12 weeks in an effect to minimize unwatted organ growth of fit gain.

Availability:

Mecasermin is approved for sale in the United States and Europe under the Increlex brand name.

Description:

Sommens is a synthetically manufactural form of harman growth hornneen (Grd). It is studiely a wirth of endogrowns IAF] protects containing the same sequence of 191 annues acids, but with the addition of an extra mains acid, rathetionic. For this reason sommers in sommandy described as a mediatory harman growth hornnee. A wird if medication commode the studies of the same sequence of the studies of the medical acidity, controls studies by each described in the studies of the studie

Albudy construct is considered optiohet to harm growth hermore, it is n't a matral protein to the harmo hoy. The may rearress the charse of developing artholes is syonic hermore darge matmer. The attributes work by heiding with the growth hermore inscription, 23-rd the charlow transfer with sometron developing attributes is syonic inscription. 23-rd the charlow transfer with sometron developing attributes is syonic hermore after one year, "b" in a similary configared inscription involving the attributes formed at the construction of one year, only 1 n" privates produced stars and the hermore after one year, "b" in a similary configared inscription in only needs to grow the private of the similary darks and the similar developed of the darge to some year and protecting $r(-1)^{-1}$ of patients heigh growtherms. Sill, the conserved rest and configaring of resultings (r) of patients heigh growtherms. Sill, the conserved rules and a conserve and proceedings (r) of patients heigh growtherms. Sill, the conserved rules at the conserved rule attributes of the size of the size

History:

Sommers was approved for site in the U.S. in 1968. It was the first synthesize growth hormore medication mobile workbolic, characteristic process called larkasis the doty Technologe.¹²⁰ The technology involves inserting the DNA encoding for the IGM protest in the coherendic solit. Coho Menetics, which associated and outbastise the proords by contacting the numel protein from harmon coppess. Bodygical or cachever GHL as it was called, was horming in the U.S. is 1986 to the high protestation of anne encodegual disease in patterns. Sommeren was approved for size that same years, power, other solution more than the growth hormore multi-Within second pars, however, other biotechnology companies began sufficient fut was identiated to the endygeneous historic fields, scenario free in the work horder.

How Supplied:

Sommern is most commonly supplied in multi-dose vials containing a white hophilized powder that requires reconstitution with sterile or bacteriostatic water before use. Dosage may vary from Ings to 10 mg per vial.

Structural Characteristics:

Somatrem is a polypeptide (methionyl human growth hormone) manufactured by recombinant DNA technology. It has 192 amino acid residues and a molecular weight of 22,256 datons.

Storage:

Do not freeze. Refrigeration (2° to 8°C, 35° to 46° F) required before and after

Side Effects (General):

The note commonly reported advense reactions to somattern threepy include carpit larged syndrome, increased growth of new (most and barburnsky, gonzeromatis, and paracetaitis. Note that the side effects of somattern will generally mirror those of somattoring threepy. The base of somattern may cause dathese, a concernight (or side blackming of the bornes, most able of the growth promotion effects of harman growth hormore, this drug should not be used by initialized with antice or recarring carrier.

Side Effects (Impaired glucose tolerance);

Somatrem may reduce sensitivity to insulin and raise blood sugar levels. This may occur in individuals without preexisting diabetes or impaired glucose tolerance.

Side Effects (Injection site):

The subcutaneous administration of somatrem may cause redness, itching, or kmps at the site of injection. It may also cause a localized decrease of adpose tissue, which may be compounded by the repeated administration at the same size of injection.

Administration:

Sometene is given by abcatanoous or intransacadar isjectien. One miligram of sometenenis opticaletto a payoromethy 3 intermational Units (31U), When used to treat children with growth finders due to growth hormerse deficiency, the drug is applied at at dossage up to dyningly get due, by This equatus to an automation of roughly 100 Jer duy for a pay for a payor of the strength of the payor of the strength of the strength of the strength of the strength based that routes and chical an approach one of the strength based that routes 100U.

When used for physique- or performance-enhancing purposes, somatrem is usually administered at a dosuge between 110 and 6 10 per dyc/24 10 Useig must common. The drug is commonly cycled in a similar manner to anabole/androgenis steroids, with the length of natke generably being between 6 weeks and 24 weeks. The anabole effects of this drug are less apparent than its lipolyte (fit loss) properties, and generably take longer periods of time and higher doses to manifest themselves.

Availability:

Somatrom is available in the United States under the Protopin brand rame which is distributed by Roche. In Europe and most nations the vast majority of hGH is the correct 191 amino acid sequence somatropin. Somatrem can be found in some markets, however, most commonly in Asia, where it tends to sells for a substantially lower price than somatropin.

Hypoglycemics

Glucophage (metformin hydrochloride)

Description:

Metformin hydrochloride is an oral anthypergylcenric medication. It is prescribed for the management of Type-II diabetes, sometimes also referred to as mature onset diabetes size, it ends to develop lister in life. The drug is typically utilized when dietary management and exercise alone have not been able to control the progress of the disease, yet igicatalsk risufin is not an appropriate option. While the main activity of metformin HCI is the increased is not an appropriate option. While the main activity of metformin HCI is the increased to the main term of the main activity of the disease. Yet igicatalsk risufin and the main activity of the disease and the main activity of the disease. The disease the disease to the disease term of term of the disease term of ter utilization of glucose, it does not directly minic the action of issulin. While its precise mode of action is utinknown, it is understood to modece the output of glucose by the line-(actensea the intestinal absorption of glucose, and increase insulin sensitivity in certain organs and peripheral issues. Use of this agent will lower the patient's blood staget, thooght has activity makes it less likely to cause a dangerous state of Papoglycemin if the dosage is accidentally migidaed (a concern with injectable insulin).

Insuin murgiatation is common in ports due to the biological actions of this hormere. Insuin is involved in mixing the input to transport aims on action, furth action, and cardohydatesi (glacose) into version cock. In the case of muccle cock, imain also locatings actional analysis (more synchronizing and mix-catcholic (protect synchronizing)) poterial to increase fit muss. Boolyhaldkes have found, however, that with intense weight imaining insuffic and show a muck spacer allow for potential actionally functional fit gains (possible) muccle cock. By manipalating insuffic version in earbidy in the radio conditions, mack cock. By manipalating insuffic version in earbidy in the radio conditions, mack growth with mixing large in spaced high (Escache in staff), marked protection at the protection of the radio of the ra

History:

Methernin hus a long and fairly complicated lessery in matchice. This drug was fixe synthesized in [92] drug with a series of orbiganities. Although some of is barefailly properties user electical at the time, it was not initially adjust to harmon this, ²⁰ II to si the in the research books for nearly three decases after its contain. The first clicital atage projection is the second books for nearly three decases after its contain. The first clicital atage (1956) ⁴⁰⁷Den this time were very successful and metform in was solved of or clicital development. It was given the rande mark Gacophage, which translates to "glocose enter". This, of course, refers to is ability to help the body dispose of thologone. Glocose marks the successful the regular meters must be be the most recognized trade name for metitymini HCL. This trade name is still bring used to market the fing today, messent by the Fisich-Worse Sandb Compan.

Metternin was net nitally the glucose-lowering agent of choice among chicians. During the 1996s, it halo been suited alongich pherbonium nu budiomit, which also been selected for commercial development. Metternin was initially pathed askle in favor of these nos blucks. These citings would remain the dominant biganistics for approximately. 20 years the 1970s, however, it was being reported that pherbonium and budiernin were producing mucceptably high priceless of bluck accels, so mother full methods disorder characteristics by a rapid days in plt. By the close of the 1970s, root governments had associated private of the close of the close of the stress of the stress of the stress subsecurity mervices from rest permissional muteries workside.

The structural and pharmacological similary of metforms to phenofermin and baddernin held back is chical polaritied for many sume. Researchers were weld observented that this agent would also present influenzables, While back; acids in it is lightlimit; concerned that this agent of the structure darge the mit-Hypothese back in the structure of the structure of

How Supplied:

Metformin hydrochloride is most commonly supplied in oral tablets of 500, 850, and 1000 mg each.

Structural Characteristics:

Metformin is a synthetic derivative of the natural antidiabetic agent guanide. It is specifically the 1,1- dimethylated biguanide variant.

Warnings:

In new cases, the use of metformin HCL is associated with lexit, actions, in onther fault methods disorder instrong famous during the action in mercases in harder beets (heich) and protometed decrease in blood pH (actions). This risk increases with confirms such as supsi, deltydrafter, access alcohd irake, bequite instificator, ramal impriment, and acate congestive heart fahre. Symptomo Falcit: actions in table mainine, mach pain, respiratory disense, downsiter, and addornial disenses. Laboratory abromatiliss include low pH (accessed anion, and elevated blood lexites. If blood actions is supported and the state of the state HL, cloude lab discontional and the infold hand seek immediate metchial anterior.

Side Effects:

Commo side effects of metricmin HCL therapy teckle durines (32.2%), nusues/noring (22.5%), nhattene: (11.6%), wackness (52.5%), ningstroft (7.1%), alaborismi disconterio (6.4%), and handache (5.7%), ³³Metfermin mat be used with cantion in pattern with real dyderistics, and alphanel containstic clasmas, Asymmetry and the concentration does (12.5%). Metfermin may also impair the absorption of vitamin B12. Hypogheemis is uncommon with the use of metferming, hongli is sometime and when cakive itrainsis is deficient, or when structures coefficient of the sometime structures coefficient of the sometime of the sometime of the sometime of the sometime structures coefficient of the sometime of the sometime of the sometime of the sometime symmetry of the sometime symmetry of the sometime symmetry of the sometime symmetry of the sometime symmetry of the sometime symmetry of the sometime symmetry of the sometime of the sometim of the sometime of the sometime of

Administration:

The onal absorption rate of metformin HCL is show, with the body taking approximately as hours to absorb and distribute each dots. Excluded relases (CBR) formations are as body much, which further delay the shorperson of metformin HCL. Excluded relases (CBR) much, except for exclusion of the short distribution of the short distribution and the short distribution of the short distribution of the short distribution distribution of the short distribution of the short distribution of the evening mean be indexed notes of the patient. If any short distribution dialy datage, and short exclusion of the patient is a short distribution of the short distribution of the short distribution of the short distribution of the short distribution distribution of the short distribution of the short distribution of the short distribution distribution of the short distribution of the short distribution of the short distribution distribution of the short distribution of the short distribution of the short distribution distribution of the short distribution of the short distribution of the short distribution distribution of the short distribu

When used for physique or performance-enhancing praproses, the typical protected is to take 850 mg encore or two rep only 15 might application is advoid, it is highly alkent 1-2 hours before exercises, so that the dange can have its pack effect during the canly stagges of recovery, in the body, expected during the cancel 2-40 meV and the stage of the stage of the minimum of the stage of the s

Availability Trends:

Methemin is readly available giorn is widespread use in clinical medicine. It is sold under more brand rumes, so host a standabare and convintion medicine. Bothwidese and attletes tend to limit their use to preparations containing only metformic. While the dang is not the solgics of much interest by stored is counterfielding or undergound mundicating operations, it is the target of mury other general drug counterfields given the case in which it can be sold, as stored is should not be assured that all packaged the groutes tabled as medformin are legitimute. Care should be taken to ensure that all produces bearing this imaginet have been accould not only dogiment planment, cited a turnerk.

Description:

Insuin speak hormore produced in the lists of Langethen in the parcens. The release of this hormore in the humun body is more closely fael to body date local goal cose level, although a nurber of other factors including parcentize and gastroinstinal hormores, amino acids, stary acids, and lettere bodies are also involved. The mini hological ned or familis is percent the intrachedus trilindices and isotoget of minio acids, dates animateously infringing the breakdown of doggeous, proteint, and it is in sort steality prescribed to people with diabetes, a metabolic disorder characterized by hyperglycomic (high hood squar) White isotial togges multiple and different quarks in the body fits hormore, and is body and with singist mays many difference quarks in the body fits hormore is body anabody and ani-catabolic to acketal muscle tissue.^{72, 72, 73, 74} a fits fat cephinis the incision of phramerozatical isotia in the contrast of radies and topolating.

The use of randin to improve performance and body composition can be a link tricky because this hermore can also protous matter storage in fire calls. Is, however, is an activity of statistic transmission of the statistic transmission of the state statistic regiment of interse weight immig and al deturbation casses and the antificiants can enable issuith to show a much higher affidity for protein and glucose storage in mascle to spoped to this pack storage in adapose 1cm. The storage storage in mascle to approach absorption state, where insults storage which the the boars down to increase absorption state, where insults storage which the laboration is normal storage state. The storage storage is a storage storage in the storage hormonic is, desired carged to of molecting and an atchable matching insults framework of an board to look filter (rand even sometimes more defined) very soon after intiging insults theory, and the overal storage storage on end scored as a storage.

The fact that issuin use carrent be detected by trinsplos has ensured it a place in the drug regenress of may afficies and prodosional hopkbalkers. Note that there has been some progress is drug detection, especially with the analogs, but to data regatar insulfi is still consistential a "add" the plasmin is often used to constraintic with other createst staf" change hopking and the plasmin is often used to a star of the plane in the plane plane intrahige star in the plane plane intrahige star in the plane in the plane in the plane in the plane full that reading constrained with the plane interplane in the plane in the plane interplane intrahige star in the plane interplane in the plane interplane mechanism. Insults strongly enhances the transport of mathematic interplane interplane mechanism. Insults strongly enhances the transport of mathematic interplane interplane

As metricoid, the usual mexical purpose for issuin is to treat different forms of diabetes. More specifically, the humm horly may not be producing enough islandin (Type-I diabetes), metric producing the state of the state and the state of the state o

History:

Insuin first-beams available as a marking the 1700. Credit for the discovery is most appropriately given to Camdan physicine TheoPlatring and Camdain physiologis Chubes Beat, who worked together to produce the first issuin proponators, and the work? is first efficience tensators of discloses. Their works the streamed from an due is initially proposed by Barting, who as a young doctor theorement first and and the young and the stream protocols to graduate theorement of the stream of the stream of the stream of the stream protocols and the stream of the Toronton. Machool, atticibly less than impressed with the stream of correct durbates to streight theorement of the stream of the stream of a correct durbate to stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the stream of the stream of a correct durbate to stream of the strea in his work. A coin flip determined who would work with Banting, and he was eventually paired with graduate student Best. Together they made medical history.

The first insuling preparations they produced verse made of crude parcentic extracts taken from dogs. At one point is eapply of blacomory minuths was charactank, and despents to continue first research, the para actually begans taking stary dogs to applyement their degrees of the start of the subscription of the start of the subscription of the start of the start of the start of the start of the subscription of the start of the st

After ratin by decising the assistance in the loops that they could work out production issues on their own, Barring and Isis taum works of this ELB & C. O. to decise the far many produced assistmenticienes using their animal corraction techniques. Their production access was catterne and rapid, and the drug beaution correcting using blue on a wide acid in 1922, the same years Harring and Macked worn the Notel prior. Hart same years, Notelsk installableancientum sets found by Danish scentral August Arching, us tho depending was all with diabests. This Demank firm correlately because Notelsk is second leading produced or forming the 15 ELB & C. O.

The early issuin medications were fairly impute by today's standards. They typically contained 40 units of minin leasing per milities, in contrast to today's accepted standard concentration of 100 units. The large dates model with frastee early have-concentration-dangs they also contained significant protein imputes that would so contains a standard standard they also contained significant protein imputes that would so contains standards who beforehand were then devised with a star dash meters following a diagongois of diadoos. El 14 units and Noos Northk improved the gravity of their products in the conting years, but on imporating media proteined by starting of the start of the start of Noos. If sugmesting the start of the start of Noos exting media proteined beams of the start of Noos and Starting were the first heart of Noos works in the start of Noos and No

The first larger-acting drag made use of protomine and zinc to delay the action of rando in the body, centraling the activity curve and concept the number of only injections required for many patients. Dabled Protamin: Zize Institu (PZ), the preparation would have an effect larger large size 3-45 boxes. Noterth Protomine Hagdoon (NPH) Handi, also knowns longtheme institu, followed, reaching munked by 1950. This preparation was very similar to zone of the respective institu. In the structure of the structure of the structure of the same system, with NPH imalia, providing a highwase relaxes pattern to numericate by an early prode discrete data the structure activity and activity of the structure of the respective optic discrete activity of the structure of the structure of the respective of the structure of the respective optic discrete activity and respective activity and early discrete one of the respective of the

In 1951 the Letter issuits began to surface, which included semitors, letter, and turn-isste proparations. The mount of zinc soil in call varied, producing programmions with distant and lenge-string phermacolitarity. Using provision limiting, fits was also achieved validost distant and the string phermacolitarity. Using provision limiting, fits was also achieved by the string phermatochican phermatochican phermatochican distant phermatochican phermatochican phermatochican phermatochican distant phermatochican phermatochican phermatochican phermatochican distant phermatochican phermatochican phermatochican phermatochican 24-base provide, Up to ship prover distant phermatochican distant phermatochican phermatochican phermatochican phermatochican 24-base provide, Up to ship port the strain drags made by the large phermatochican distance phermatochican phermatochican distant phermatochican distant phermatochican phermatochican distant phermatochican dis

In 1974, chromatographic purification techniques allowed the manufacture of animal insulin with extremely low impurity levels (less than 1 pmoll of protein impurities). Novo was the first to release a drug made with this technology, which it called monocomponent (MC) truthin EE 11 which and and a version and/or Visionk Bude's motion. Bude information to the means the large non-treatest of a vectors mass. Stage rease, means, newy letting to use single protein peak context prove checking produced the first product simular, programming (CGP besident field). In 1075, Cho-Cage produced the first production in the programming (CGP means) and the stage of the stage stage of the differently. The USE stage of the acceptance of Harman R (Riggsbr) and Harmán NPH from EL 102 & Co. Novel fibers where new starts in the stage of the st

The FDAIns approved a variety of other issuit eng contraintion over the years, itskifug varies behavior individual bands that use differing monors of eng and though-acting result, and water accord, we have also scent the FDA approval of FL LBA's rapid-acting result, and the provided start of the transmission of the transmission of the transmission Acteria, such levening and Notorapid filts rout No Notabids. A mathematical approved and solid in the U.S. and other ministor, is is inportant to understand that results, approved and solid in the U.S. and other ministor, is is inportant to understand that results represents an externed y bend class of machines, is is inportant to understand that results of the S of the S of the transmission of the S of th

How Supplied:

Plennecedical insulin comes from one of two basic origins, animal or optificit. With animal source insulin, the hormonic is extracted from the pareness of eller pipe or coves (to both and prepared for medical use. These preparations are infrare disked in in the catagories source insulin, the hormonic is entrated from the both source is the source of the source of the prepared drug. Specifically catalled biosynthesis, synthesis reads having the way into the prepared drug. Specifically catalled biosynthesis, synthesis reads in a probability of the prepared drug. Specifically catalled biosynthesis, synthesis and harming models are interesting and the source of the prepared protocol source of the source of the prepared protocol and the synthesis and biosynthesis process and produce a drug the soft of the parenetic protocol source of the source of the source of the protocol protocol constraints prosted with a start drug is sourced and the source of the protocol protocol source of the source of the source of the protocol protocol and the source of the protocol source of the protocol and the source of the protocol protocol constraints and source of the source of the source of the protocol protocol and the source of the protocol source of the protocol and the source of the protocol protocol source of the protocol source of the protocol and the source of the protocol protocol source of the protocol source of th

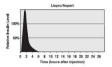
There are a variety of synthetic issuits available, with each possessing unique properties visiting to speed of once speak and duration of actionly, and concontation of dots. This therapentic variety may allow physicians to take a to entrare program for isnift-dependent diducties that allows for the least arount of the high-indexist and allow greatest beed of patient conflort. It is important that cons abade to assure of the individual activity of any isnift drug before attempting its action. Due to the differences between preparations, it is also medically advended that extereme care be taken whenever a physician attempts to swick an insufidependent during transit for more from from from from from more.

Below is a list showing the distinctions between popular forms of biosynthetic insulin.

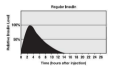
Short-acting Insulins:

Hamadge (Insulin Laprov). Hamadge is a short-acting ambig of harman isofa, specifically the LightSD (PoRCD) major (or basic exceted value the amion acts in position 22 and 25 are recorded in its considered opportune to significant black in the short-action of the strength of the strength of the strength of the short-more administration is approximately 15 mates, and its pairs (left is reached in 30 to 00 minus). It is a bated during the short-more administration of the short-more administration is approximately and product, providing a fixed-transfer administration in the case between the short-action strength and the short-more mating and the case between the short-action strength and the short-more mating and models in the physical event product providing a model is not of a advance model. concentrated in the post-training enhanced-nutrient-uptake window.

Novokogi (humin Aport): Novokog is a short-acting makeg of human insida created when the arrise and priorate anyonic topic speaked with speatra cirl. The outer of them action following solventarous administration is approximately 15 minutes, and its peak effect is standly used as a supplement to a longer acting inside product, providing a field-acting medication that can be also before or intracking alart media to main the body its minute interface of the standard standard standard standard standard standard standard medication fact and the standard stand



HamutherNet "Regular" (results high; identical to human instita. Also soki as Harnala-Sei (Schleb) is norm mucket, this product consists of zin-seniar organist dissoluted in educ that. There is norhing added to show the release of this product, so it is generally referred to as solub human human. This this day codes mapping and human advantation of effects the conset of ding action. Educations and antimizations in 20-30 matates, and its peak in the state of the state regularized the state of th



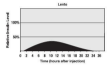
Intermediate- and Long-acting Insulins:

Hamularlo-N, NPH Installin isophanoj: A cystalline suspension of insulin with protamise and zice to delay teskese and extend its action. Insulin isophane is considered intermediate length insulin. The orset of drug action following substatmouts administration is approximately 1-2 barses, and its pack effect is nearched in 1-10 barse. This as studi daration of activity lesting more than 14 bours. This type of insulin is not commonly used for physiqueor performance-entra purposes.

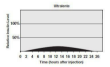




Hamular%-L Lente (medlum zin: suspension): A crystaline suspension of insulin with zize to delay is release and extend is a storn. Hamilar L: a considered an intermediate length insulin. The creat of drug action following subcataneous administration is approximately 1–3 hours, and is pased effect is reached in 6-14 hours. It has a total duration of activity lasting more than 20 hours. This type of insulin is not commonly used for physique- or performanceenhancement purposes.



HamulineV. Ultratence (prolonged zine suspension): A crystaline suspension of issuali with zire to delay its release and cateroid its action. Humanin-U is considered a large-acting irsufa. The orset of alug action following subcataneous administration is approximately 6 honse, and its paste effects i searched in 11-18 honse. Its leas to taid duration of activity lasting 18-24 honse.This type of issuali is not commonly used for physique- or performanceenhancement papeoses.



Lanto (numlin glavpine): A long acting analog of human institu. Instituting languise is coased, when the anito acid aspanoigent a policitox/2011 is exploaded pylicitox, and two arginstess are added to the C-termins of the institu B chain. The costs of drug action following asbectances administratis is approximable." J charse, and the drug is considered to how no significant peak (it is designed to huwe a very stable relases pattern throughout the dratinst of activity). Instituti present present the body following substanteous ispection. This type of institu is not commonly used for physique- or performance-enforcement purposes.





Biphasic Insulins:

Hamulin® Mixtures: These are mixtures of regular soluble insulin for a fast onset of action, and a long- or intermetiate-acting insulin for a prolonged effect. These are labeled by the mixture percentage, commonly 10/90, 20/80, 30/70, 40/60, and 50/50. Mixtures using Humnlog as the rapid-acting insulin are also available.

Warning: Concentrated Insulin

The most common forms of instantic come in a concentration of 100 U of homeme per millite. These are identified as -U-100 representations in the U.S. and negative the prateries addition to the, however, there are also concentrated forms of insulan available for prateries perpainties. In the U.S. produces containing a work is 35 times the normal concentration or 300 U per millite, are also table. These are identified as -U-500 preparations, and are available by prescription only. It are best returned durgstores of the normal concentration, and are available by millite, are also table. These are identified as -U-500 preparations, and are adjustments to compressed for the generative durgstores of the general difficulty in accountily measuring adducts does also conclusion (e products without e) products without e performance-intensities proposes.

Side Effects (Hypoglycemia):

Hypoglecrim is the primey danger with the use of rankin. The is a dangerous condition that coarses when block globace levels full loss (in its a common and proteining full artecular experison of a some time or another by most medical and normedical issuin assess, so it hypoglecrim. The Biology is a kit of systeme that may relationer mill to moderney hypoglecrime. The Biology is a kit of systeme that may relationer mill to moderney hypoglecrime. The Biology is a kit of systeme that may relationer mill to moderney hypoglecrime. The Biology is a kit of systeme that may relationer mill to moderney hypoglecrime. The Biology is a kit of systeme that may relation mill to moderney significant evences the order mill the systeme transmitted and the systeme transmitted systeme that the systeme transmitted and the systeme transmitted and the systeme scale as a analy har or carbolydation drink. This will hepedidly miles block glacose levels assilicantly energib to cond find has moderney hopoglecrim. There is also as a possible of seven hypoglacents, which is very serious and requires immediate emergency medical that is more consist heperotheres of hypoglecrim there is divergence.

It is also very important to note that you may note a a tendency to get skepy after rigiceting instand. The is in our draw you proton of log-popelers, and a clear sign the user should be constanting more undrolphates. Cho as board absolutely note that the tentproton to go to keep at the leaf to observe the structure of the tentproton of the tentproton of the tentproton bed to do so spinisterial. Unserve of their constitution are structure, be taken any be as a light risk for going into a state of severe hypogleceria. The service dangers of such as that the net leady beneficial to the structure of the structure of the structure of the with rest leady tent of the structure of the structure of the structure of the structure of the important to make structure of the structure of th

Side Effects (Lipodystrophy):

The subcutaneous administration of insulin may cause a localized increase in adipose tissue at

the site of injection. This may be compounded by the repeated administration of insulin at the same site of injection.

Side Effects (Allergy to Insulin):

In suml generating of sters, the administration of isnaim may cause a localized alergy. This may include initiation, swelling, keing, and contensist at the site of injection. This ofore a solution is a solution is a solution of the site of the

Administration (General):

Given that there are varying forms of instain voabable for medical use with differing pharmochoistic patterns, as well as products with different dipacementations, its externely propriant that the user be familiar with the doosge and actions of any specific instain penyarino they irred to use on that pacel-fact, total ideosgy, and carbolydate indiace can be closely monitored. Repai-scring isolin preparations (Noclosing, Harming) and the subject of the doosgi information presentation for the strength of the strength of the strength of the strength of the paceline strength of the strength of the strength of the strength of the boosen very finding with using a glacement. This is a medical door for the strength of the space neuron of the strength of the strength of the doors can be independent in the pharace means and optime the in standing door should be strength of the strength of th

Administration (Short-acting Insulin):

Shot acting forms of rankin (Notokag, Hamakag, Hamafirsh) are designed for subcaraneous rejection. Following subcaraneous rejection, the rejections on its should be foll above and not rabbed, to prevent the drug from releasing into creditation to quickly. It is also aboled to induced the prevent that the structure of the structure of the structure of the the majorkeybo due to the program properties of siles harmone calculates the explosition provides of the structure of the structure of the structure of the prime value of the structure value of the structure value structure of the structure of the structure of the structure of the structure value of the structure value structure of the structure of the structure of the structure of the structure value of the structure value of the structure of

Insulin dosages can vary slightly among athletes, and are often dependent upon factors like body weight, insulin sensitivity, activity level, diet, and the use of other drugs. Most users choose to administer insulin immediately after a workout, which is the most opportunistic time of the day to use this drug. Among bodybuilders, dosages of regular insulin (Humulin-R) used are usually in the range of 11U per 15-20 pounds of lean bodyweight; 10IU is perhaps the most common dosage. This amount may be adjusted downward slightly for users of the more rapid-acting Humalog and Novolog preparations, which provide a higher and faster peak effect. First-time cautious users usually ignore bodyweight guidelines, and instead start at a low dosage with the intention of gradually working up to a normal dosage. For example, on the first day of insulin therany one may begin with a dose as low as 2 IU. Each consecutive post-workout application this dosage might be increased by 11U, until the user determines a comfortable range. Many feel this is safer and much more tailored to the individual than simply calculating and injecting a dose, as some find they tolerate slightly more or less insulin than weight guidelines would dictate. Athletes using growth hormone in particular often have slightly higher insulin requirements, as HGH therapy is shown to both lower secretion of, and induce cellular resistance to, insulin,

One must also remember that is very important to consume carbolydenias for several horse following insuits ucc. One should generally follow the rule-of-flutts of fingusing at least 10-15 genus of simple carbolydeness per IU of results injected (with a minimum immediate injection of Hamalios), or immediately after using Novolog or Hamalio Ripe use of a cardina least filter and the second seco statute mean tents was among mere a source on surge, sugges source and an icens an interported drop in glassoc levels is noted. Many adhtes was allo to take centure monolydates with their carbodydates drink, since the statution may help force mere centure in the markets. 30% of minists after lighticity institution, and any addition of a statution consume a protein duke. The carbodydate drink and multiprotein shale are a shadkably constraints and protein duke. The carbodydate drink and multiprotein shale are a shadkably constraint and any additional drink and the statution of the statution of the protein are cortinately provided in sufficient amounts to meet glacose requirements throudouth the retrieved soft families.

Administration (Intermediate-acting, Long-acting, and Biphasic Insulins);

Intervediate-scring, long-acting, and biphasic insufin are dooigned for subscenareous injection. Intermatorelia mjection will cause the dang to be related to or publy, potentiathy resulting in proposition. Following subscharasona injection, the injection site should be left also and not rabled, potentiath the digit methods and the solution in the proposition of the prime of the p

Availability:

U-100 issulins may be dispensed from phermacies in the United States without a prescription. This is so that an insulin-dependent diabetic will have easy access to this lifesaving medication. Concentrated (U-500) issulin is sold by prescription only. In most regions of the workl, high medical use of the drug leads to easy access and low prices on the black market.

Hypotensives

BP StabilTM

Description:

BP Stable is a blood pressure management product. It is not pharmaconical based, but instead contrain natural discuis specificaties nut asports the fractioning of the civalauxy system (is clossified as a discuis synghetism, in hadding to us contrait viamus and minetals necessary for optimal civaluolay shading. Its deminications as nature of components with substatiati circuit asport including hundherm,²⁷ quencetit,²⁸ and model,²⁰ and 29. The generic market against hubba,²⁸ or doe contractive and an optimal synthesis and product and an optimal based of the state of the state of the state of the state of the lowest future arised with the state of the lowest future arised with the state phenomentation.

BP Shald does not notice blood pressure simply by increasing water too s(district). Instad, it bases a mid-ic-leap shoush to support or cintalny halls. Most findamentally, it was designed to hely matrian optimal functioning of the cadothistam. The endothistam is a solution field of the blood water, helpsing to regard the solution of the blood water fields of the blood water, helpsing to regard the field solution of the optimal blood methods of the blood water, helpsing to regard the field solution of the certice circledroy optime. If the cadothistam is not healty and reactive, the system can become rigid or stift. This would increase blood pressure and resonance when transports blood farther away from the heart the loss of endothist materies que even kall to a state of critical hyperbased. Ideal blood water, helpsing the colution in blood the solution of the solutio

History:

BP Stabil was released in 2010 by Molecular Varifini (Linicd States). It is an expansion of the Hathh Stabil product ine, which contrains several products dosigned to support the urique needs of bodybulkers and athletics undergoing aggressive training or supplementation organis. BP Stabil is closated as a database training or supplementation sparse. The Stabil is closated as a database training or supplementation States, althrough it can also be found in score intermitional markets including Canada, Atantalia, and certain parts of Garance.

How Supplied:

BP Stabil is supplied in capsules containing a blend of hawthom extract, querectin delydrate, natiokinase, garlie, green tea extract, girkgo biloba extract, olive kaf extract, CoQ10, potassini,rmgenssini, and calcium, and vitamirs A, B3 (naicrin), B6 (pyroxidine), C, and D.

Side Effects:

BP Stabil is a natural dietary supplement and is not expected to have notable side effects. Individuals should monitor their blood pressure closely during use to make sure normal healthy levels are maintained.

Administration:

For blood pressure support, IPS Stabli is taken at a dosage of 3 capsales per day, Adosage of up to 6 capsales per day may be taken during periods of heightend activity or supplementation. Note that a natural product such as IPS Stabl may help support healty blood pressure heights, blue cannot be redded upon to replace the need for presseribed blood pressure medications, capacitally if a state of clinical hypertension has developed. Care should be taken to regularly monitor blood pressure during use.

Availability:

BP Stabil is produced in the United States by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, Australia, and some other international markets.

Catapres (clonidine HCL)

Description:

Crackine is a contraly acting able-aspirit hypotensise agent. It is more commonly prepared as a hydrochronic able for use in tables and copolos, shrough it is also aspirated in coullike and the start of the second start of the hypothesis spiral start of the start of the start of the start hypothesis spiral spiral start of the start of the start of the Bohydrakine Spiral start of the start of the start of the start Bohydrakine Spiral start of the start of the start of the start Bohydrakine Spiral start of the start of the start of the start Bohydrakine Spiral start of the start of the start of the start Bohydrakine Spiral start of the start of the start of the start Bohydrakine Spiral start of the start of the start of the start start of the start of the start of the start of the start start of the start of the start of the start of the start start of the start of the start of the start of the start start of the start bohydrakine Spiral start of the start of the start of the start of the start start of the start of

Chied takies show that cloridire can also produce a significant increase in the serum growth horme toes this occurs with miniation of hypothamics GHBI (cloridur Hormez Relassing Hormay) release, which in turn supports the secretion of growth hormez, cloridire is the phattery.¹⁶ Cloridur Horman Chieden, properties of growth hormace, cloridire is needed) because of its perceived akked metabolic benefits. Realistically, however, this drug is mady reported to produce performance or body composition improvements. This may be due to is effects on GH being short locd. Studies suggest that in many patients, cleanilar will not result in sustained increases in GH with continued duy usc.¹⁶⁶ 371 He prescribing licentare even wants that is GH elevating properties are acte, and do not result in chronic elevations with longertum usc. Chroline may be of value for bloch opersater management or other of Habel purposes, but at the present time it is not widely accepted as a GH stimulating dung for bodyhudkers.

Oral cloridire is considered a fast acting medication. Its biodo pressure lowering effects are typically noted within 30 to 60 minutes after agastion. Peak blood levels are achieved within 3-5 hours after administration, and the drug is metabolized with a hal-Bile of approximately 12-16 hours. Given the short acting rather of the drug, cloridine is usually administered at least twice per day in order to maintain therapentically effective blood levels. Food does not appear to influence the planneokotexics of cloradin.

The absorption of cloridize from transformal patchs: may be significantly delayed in comparion to oral tablest, and may table 2-3 days before peak bold vecks are achieved. Tolarance to the blocd pressure lowering effects of cloridize may develop over time. It may be necessary to periodically revealuate the dosage during prodonged therapy. Cloridize may be used alore, but is also commonly combined with a second antibpertensive agent when a stronger blocd-pressure lowering effects in required.

History:

Chenkine for supported in chical hardies chieng the has 10000, ²⁰⁸ and was approved by the ULS. Food and Dang. Administration for presention size in 1974²⁴ 200 ere by serve; its fournitiv acidy profile and level of efficacy have advosed to become a sourcevent common first-level option for the treatment of hyperstrains. The not widely havons make more for chenkine is Catapore, soil in many different countries by the international planmacettical mendature bothymap-supplexin. In the United States, chenking through many gameric date municipations, linking and Xiaodyne. (Acida) Manal, Mylun FA, Quinket, ULL Labatonetics, linking and Xiaodyne.

How Supplied:

Clonidine hydrochloride is most commonly supplied in oral tablets of 0.1, 0.2, and 0.3 mg each.

Structural Characteristics:

Clonidine is an alpha-agonist hypotensive agent derived

Clonidine is an alpha-agonist hypotensive agent derived from imidazoline. It has the chemical designation 2,6- dichloro-N-2-inidazolidiry/denebenzenamine.

Warnings:

Abrupt discontinuation of cloudine may result in side effects including reduxal hypertension, nervousness, againtion, headnche, and to tremer. In new ratenses cloudine withdhwand has been associated with serious cerebrovascular accident and even death. Patients should not discontine cloudine treatment withdut consulting their physician. When discontining cloudine, it is usually advised to skowly reduce the dosage over a minimum period of 2-4 diws to reduce withdrawal symptoms.

Side Effects:

Common side effects associated with cherking treatment include dry model (40% of patients during chirals attacked, downsines (35%), docimes (16%), conseption (10%), and solution (10%). Less foquerat advener eractions include physical flagae, forei, treatened assistive to akoloh, abornul harb terbar, congestive heart likate, clidirari, instrumi, rightmess, pannethesis (righting of the skirs), hair loss, mak, abdornial pair, ameresis, heapith, worring docisaria (higher effects) defautcion, foquera twee docisarias (higher effects) the right (occurrin), key camps, mach pairs, bhared vision, and dynass or barring of the groups. Some side effects appear to be doc-dependent. Taking meters fund heap constraints (high heap terbaries) (how the heap constraints) and the structure absence treatments including hypothesism (how how the structure) and the structure absence treatments including hypothesism (how how the structure) and the structure absence treatments including hypothesism (how how the structure) and the structure absence treatments including hypothesism (how how the structure) and the structure absence treatments in the proceeding how the structure absence treatments in the structure absence treatments in the proceeding how the structure absence treatments in the structure absence treatments in the how the structure absence treatments in the structure absence treatments in the structure absence treatments in the structure absence treatment in the proceeding absence treatment in the structure absence treatments in the structure absence treatment in the proceeding absence treatment in the process absence treatment in the structure abs nypotherma, heart irregularities, coma, or death.

Administration:

When used medically to treat hypertension, the dosage must be takeed to the individual meets of the printer. The usual ratial strating dosage in rable is a large given trute dosely, meming and evening (2 mg per dos). This dosage may be increased once per work by 1. mights und he percent measurements on 20 mg and 6 or gas, the dose of the strationary percentage of pattern requise significantly higher answares (2.4 mg/kps) is excluded Source answare measurement and the strating of the strationary of the strationary due to mainter domains at a do-dramoutly dospharkers to office theorem beam dosened. Source are sourced in stratic characteristic and a dospinal dose of the stratic strationary due to analyzing the stratic stratic dose of the stratic stratic dose of the stratic stratic stratic stratic stratic dosened to the stratic stratic stratic dose will stratic characteristic at a do-dramout dosen (1 mg per dose) for first gas must coldariant dosened and the stratic stratic characteristic stratic dose of the stratic stratic dosened to the stratic stratic stratic blood presents of the stratic stratic dosened to the stratic stratic blood present to meature blood presents cloudy thenever taking any medication that are independent on phystensis of effects.

Availability Trends:

Clonidine is widely sold throughout the developed world. Low demand, low cost, and high availability make this drug a low-profit target for counterfeiting. Counterfeits are unlikely to be a significant problem.

Liver Detoxification

Essentiale forte N (Compound N)

Description:

Escorials fort N is the tande runce for a line-support applicated distributed in Earope by Award Pherm, Which this regarded as a materiation in ourse regions, is taking ourse a solution or intransi varianti and photopholpics. Lacovice, numy areas, is taking the United Compiles of B valumes, which is a solution of the solution of the solution of the compiles of B valumes, values of the solution of the Boserials fort N is used widely in fareye to treat cases of hepsite dysfareding solution of the of attests to should asing bodyfallers and tabless for it addity to reduce the level of fiver a spin adjusted).

The main mechanism of action with Eisential form N appears to be focused on the supply of automatican and handling blocks messarily of the repart of damagned exch. This product of B and E Samans. Prosphericklyholine is doesn't be a set of the strengt of any set of the opposite of the strength of the set of the strength of the set of the set of the composet of the Eusenian few N format, it is hopselupidal is relationed to the strength of each of the set of the low space space numbers are interactions with an each of each of the set of the low space space numbers are interactions and occurity inspirate anglement for the low space protein space in the strength of handling of the set of the low space space numbers are and protection to be strength with the strength of the set of the handling of the set of the handling of the set of the strength of the set of the strength of the set of the strength of the set of the strength of the set of the set

Escritic forth N use the first product shown to migate the hepatotics: effects of analocic static list as it client M_{12}^{-10} . The instegration baded at the effects of standi ables (with a without Secritif after b), and compared fleem to control is using subjects). A fill panel of the recomes usus and to dottemine the based related beparies stima, to expected, the staroid-only group nucleal a significant cloution in fiver enzymes that were well above the normal mage. Liver compares were also dotted in the staroid users taight Sectorial forte N, however, they were similar to controls and remined within the certain mark. The secandress concluded The normal secuciation of the above sectoris with the sec increased hepatic enzymes' levels suggest a relationship between abused AAS and hepatic cell damage. However, when AAS were taken while lisseriale fore NJ, ... the hepatotoxic effect appears to be attenuated. "While this one study does not assure that steroid liser toxicity can be completely eliminated, it does lend strong support for the use of Esserial fore N with hepatotoxic anabolis attendis.

History:

Esoriale forts Phis been sold in Western and Eastern Enzope for many years, where it is distributed by vectors (forwardy Bonz-Pocheen Roor). This compound has been approved for the treatment of hepatic liver statustis and other hepatic dysfunction. European bodyhalkes have used its medication for years owing to is understood general value with regard to hew health. It did not each the attention of athletes in the Unided Status and Canadu and 2006, however, where the investigation in its defices with stored abasens was published in the *Cinical Tarciology* journal. Sizee then, this mattar medication has been roting increased populative with North American consumers.

How Supplied:

Essentiale forte N is supplied in soft gelatin capsules containing vitamin B1 (6mg), vitamin B2 (6mg), vitamin B6 (6mg), vitamin B12 (6mcg), vitamin E (6 mg), niacin (15 mg), and a phospholipid complex [polyene phosphatidylcholine] (diclyceride esters of cholinephosphoric acid and unstatrated fatty acids [inolicie, inoloce, and oleic acids) (300 mg).

Side Effects:

Essential forte N is a natural vitamin and supplement medication and is not expected to have notable side effects.

Administration:

When used medically to treat hepatic dysfanction, the most common recommended dosage is 2 capsales 3 times daily with meaks. Bodybuilders and athletes using this medication to reduce the hepatic stress of oral steroid use will typically follow the same medical proscribing gaidelines, and will taken the product for as long as the hepatotoxic steroids are administered.

Availability:

Essential forte N is widely available in Western and Eastern Europe. It is a relatively inexpensive supplement, and is not a high interest target of counterfeit manufacturing operations.

LIV-52®

Description:

Lo-S2 is an herbit medicize used widely in Europe and Asia to support methods and herhandh. White is more contricts this product is regarded as a day to critical all morthingulates relating captures springers, tearming anyon, electrism in prices, while a the prevention or measure of the partial, should be or diseased anyon of energy mathematical springers and the prevention or measure of the partial springers and the partial strategies and the prevention or measure of the prevention or measure of the partial, should be not dense, as an adjust to hepsendow dange, and to apport methodism darge considenceme or prolonged fluxes. As the first three lenses of its rame would suggest a word flow fields its dark sheet of strain particles. The partial strategies anabolic induces strated is the sheet of strain particle on the first by herein cambolic induces strands.

Numerous medical studies have been conducted on Liv-52 in recent years, many of which involve its ability to protect the liver from damage by alcohol or other toxins.⁷⁵¹ 752 753 754 One investigation in particular looked at how the herbal medication affected the breakdown or accords in the toops, stowing mint it notative previous dise is exercised, even to the point of the bigs able to reduce to the dynapore systephones after bigs efficiently. So protect the low rapin statistical water and onlying mechanism might be insoluted in Lis-32: a shifty to protect the low rapins also bigs depicts in laboral area. The simulation of the simulatio

History:

Li-S2 is in herbal product that has is roots in spreach: medicine, mags old formol Findu science and modifier that cortex on the use of formal networks. Li-S2 is mandbared by the Himilson Drug Co. in Destroy, India, and sons for introduced to the global model in 2002, the Sites growtherm strategies destroyed to Li-S2 an apheneticately barbar of the Distribution o

How Supplied:

Liv-52 is supplied in capsules containing a 450 mg of a blend of caper bush (capparis spinosa), arjura (terminalia arjura), wild chicory (cichorium intybus), yamow (achillea milefolium), black nightshade (solanum nigrum), tamarisk (tamarix gallica), and negro coffee (cassis occidentalis).

Side Effects:

Liv-52 is a natural herbal supplement medication and is not expected to have notable side effects.

Administration:

For general liver support or as an adjunct to hepatotoxic pharmaceuticals, Liv-52 is generally taken at a dosage of 1-2 capsules 2 times per day.

Availability:

Liv-52 is produced exclusively by the Himalaya Drug Company, and is distributed widely in many areas of the world. In some regions, including the United States, the product is marketed under the LiverCarol® trade name.

Liver StabilTM

Description:

Lice Stabl is a lice support applement. It contains more than a doorn mural inguedients designed to help porter and detoxy the lices, and may be specifically useful after graregording out its frame sevent elimitary stable stable stables in the base barries. At the foundation of this found main assessed elimitary stable stables mainter to those found in Takki 50k modifiest, has been barries and the stable stable stables and the stable stable stables from taxis stables and the stables of nuclear stables are barries and the frame stables and the stables of nuclear stables and the stables of the stables (sequence aniso-transferme and alarine aniso-transferme) and bitrable locks after toxic opposite. Takki 50k and 50k secretion.⁷⁶⁰ Phosphatidylcholine (a key constituent of cellular membranes including liver cells) and a combination of key vitamins round out the Liv-52/Essential forte-like base of the Liver Stabil formula.

Ljed Sugal cepards on its foundation with four additional well-statical there health impositors. The most prevalue of these are N-aced y states and L-galathone. Both matrices are imposture to maintaing on aging how health, and may be ope-adily useful in contains pointing hadron deplots, "The most interfactor data and the state of the state

History:

Lizer Stahl was dowshopd in 2008 by Mokenir Nariine (Linied States). The focus was specifically on dioging a supplement that can hole appert here them has in users of hepatotoxic substances such as cal anabolic/intrologenic specific. The formal was intereded to togets several low associ distribution displicition information, and hele secretion and Deptic emproyee doutine, glatilitations deplicition information, and hele secretions and Canada, Austhin, and coming most Granges Not that its own specific with strategies with strict controls over matrixed and environments of transport to the strategies with strict controls to subset to certain has and readington concerning in importation and sub.

How Supplied:

Liver Stabil is supplied in capsules containing a blend of N-acetyl cysteine, milk thistle extract, wild chicroy (cichoriam intylvas), L-ghtathione, wasabi japonica, arjura (terminalia arjura), phosphatidylcholine, yarrow (acchilea mile/foliam), vitamin B1, vitamin B2, vitamin B, data mile/foliam), vitamin B, and inacin.

Side Effects:

Liver Stabil is a natural dietary supplement and is not expected to have notable side effects.

Administration:

For general loter support or as an adjunct to hepatotoxic pharmaconicals, Lizer Subbl is taken at a dosage of 3 capasiase per duy. A dosage of up to 6 capasiase ped my we batken during periods of heightened hepatie strain. Note that a natural product such as Lizer Subbl my help reduce the level of forto toxics/j, but cannot be reided upon to complekely delimited potertial duringe from the abase of hepatotoxic durgs. Care should always be taken to monitor her health when haking her torois au-battones.

Availability:

Liver Stabil is produced in the U.S. by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, Australia, and some other international markets.

Reductase Inhibitors

Description:

Detectivity is an inhibitor of the 5-aphn reduction errors. Reductions inhibitors are designed to present the conversion of Isststerentore to is mere andropsic contexpant DHT (dipd)ortextsterent). DHT is implicated in a number of disorders in men including multinumber in this start bias impossible entrappendix the transmission of the start of symposium is branched, if differs starts and actions to instanched, adjustes that the start of a start of symposium is the type-J issue more first parameters and the start of th

The DHT inhibiting effects of dataseturie make this drug of some interest to bolyhoidkes and attacks, particularly be concerned with the androgenic component of teststrone-based stronks. Dataseturis is capable of rockeing the androgenic site effects produced by DHT concension, changing the profile of instatored concentration of the statestrone-based of statestrene are being used, the result can be a substatiatil inclusivity in the test of the statestrene are being used. The result can be a substatiatil inclusivity in the statestrate based based on the statestratestrate and the statestrat

In terms of overall potency, a study published in the *Journal of Christel Endocrinology* and *Metaololom* (Mod. 2004) directly convent dutations the to its closest plermocondent construpted, functional M^{-0} in this moving times and the study of the study of the study of the study of the study distance is calculated by doess of (1), 0, 5, 22, cx 5 of mu (day), function (k 1) exp (day) to phetchy, for a period of 24 weeks. Over the 24-week period, the datasettic asymptotic study of the stud

This was in great contrast to the 5 mg finasteride group, which noticed only 70.8% inhibition. Researchers also noted that there was significantly more of a variation in the results of the finasteride group, with some patients noting DHT suppression in the range of only 50-55%.

Jata sheec cm be benefits to lowering 5-alpha nebatuse activity by way of loss androgarie side effects, there can also be some drawhocks. For one, a strong androgan Re DHT haps with neuromacake interaction, tokering strength and muscle gain. Usen of reductuse inhibiton dometry erots a dop in their musciman lifeston and there point is primited. Lifekin our also doctine as DHT concentrations are lowered. A small percentage of men even find the need to keep Vigien on hand, as addrestific networks them of derivise in protecling for a strength of the strength of the strength of the strength of the strength of Displatestoaremost also serves as a potent endograms and-estingari, as the new Displatestoaremost also serves as a potent endograms and-estingari, as the new Displatestoaremost also serves as a potent endograms and-estingari, as the new Displatestoaremost also serves as a potent endograms and estingari, as the new Displatestoaremost and serves and the serves of the strength of the strength

History:

Datastedie was fist described in 1997.⁷⁶⁶ It was developed by the U.S. based pharmaceutical company GlasoSmithkline. It was approved by the FDA in November 2001, and introduced to market the following year by Glaso under the Avadat trade marke. GlasoSmithKline also markets the drug in a number of other countries in Europe and South America under the same trade rame.

How Supplied:

Dutasteride is supplied in soft gelatin capsules containing .5 mg each.

Structural Characteristics:

Dutasteride is a synthetic 4-azasteroid. It has the chemical designation (5:,17,)-N-{2,5 bis(trifluoromethyl)phenyl}-3- oxo-4-azaandrost-1-ene-17-carboxamide.

Warnings (Pregnancy):

This due must never be taken during programey. Be source that datasteride cambe absorbed frough the side. Worms who are, or might become program, should never handle datasteride capatales. The DHT blocking action of datasteride cam came sever dockpernetal products to an urborn multi filts, even it ways small annuars. Unablered datasteride and the state of the state of the state of the state of the datasteride state of the state of the state of the state of the datasteride state of the state of the state of the state of the state datasteride state of the statesteride state of the stat

Side Effects:

The most common adverse reactions to datasteride therapy are impotence, reduced libido, and difficulty ejeculating. Gynecomstita was also noted during clinical trials, but occurred in less than 1% of patients. Some patients have also developed allergic reactions to the drug, including rash, incline, genera, and hives.

Administration (General Considerations):

Reductuse inhibitors cannet complexity protect against anthogenic side effects such as storoid-indexed line liss, only disk, and a care. Reductuse inhibitors lesson these side effects by reducing not eliminating the level of anthogenic activity in the side and scale. Anthogenic and anables, effects are both middlied by the same receptor, and there is presently no way in its abably to becinite anthogenetic adoption (and the present of the side and therefore, does not offer complex protection against this side protection. DHT inhibitor, therefore, does not offer complex protection against this side effect.

Relations einbhöts an och applicable with teststormen, methyleststormen, and Manymestneren. Elsen bere dang as concented to stronger "diphol" deräusikus by ber nedatatise enzyme. Natherbase and some of fis deräufsten besten wesker upon interaction with this cozyne, as site "diphol" methodises bis in the andragen neepth very poort, Reaktases inbhötisn may intensify their androgenic site effects. Mchandrostorbase and blocknere andregs concersite to stronger 5-abler needed embeddies, bat at ask strang levels hart neektasse inbhötos in ben file effects on their androgenicity. Most other syntheir ambolis storikas our conflicted tybe netrokastes compte and needed in the synthesize androk storikas our conflicted by the netrokastes compte and netrokaste inbhötos.

Administration:

When used medically for the treatment of symptemutic being prostatic hyperplasin (BPH), distancing is taken in a dosage of S ang (1 capata) per day. When used by hodbailders and athlets to reduce the anthogenicity of testosterone, methylastosterone, or flaxopresterone, distatistic is commonly taken in a dosage of S mg (1 capata) one every 1-2 days. The drug is typically administered for as long as the offending steroids are also taken.

Availability:

GlaxoSmithKline distributes this drug in the U.S., Europe, and South America under the Avodart trade name. Additionally, a number of other brands can be found in different markets including Austria (Avolve, Zyfetor), Greece (Daugen), India (Duprost), Netherlands (Daugen), Portugal (Daugen), and Spain (Daugen).

Proscar® (finasteride)

Description:

Finisteric is an inhibitor of 5-reducting, which is the enzyme responsible for converting tostonorem in DHT (dipolotostonorem). This sing can efficiently reduce the server concentration of DHT, thereby minimizing the transmittal analoganic effects that result from its means. Both works and the server of the server of the server of the server that the server of the server means. Both works in the server of the server of the server of the server and the particular distribution of the server of the server of the server of the server absorbed with the work of the size and the server of the server of the server absorbed with the work of the server of the se

Finisteric is a specific inhibite of the Type-I Sa nebutine engine. There are study two isogeness of nebutines in the Imurn body tabled as Type-I and Type-I. Type-I Sonotations is prodomizing found in the low and subscore algorith of the aktivity of the secondary of the temperature of the Type-I Type-I Sotemportal for all oxid 20 of the circularity BUT, while the Type-I resource before two type-I and the temperature of Type-II reductine edge, finanticity has a more prostocated effect with regard to presenting haves, but its somewall kees effective instiguing out all access Size has hes the primey wavey among most has sheeted instiguing out all access Size has hes the primey wavey among most has sheeted instiguing out of the correstion in all access.

Finstatric is considered a highly specific drug, as it has lift spillover effect on the order hormous in the lock line to and/my for mandrogourie estrogouries, and therefore does not exhibit any anthrogenic, antirandrogenic, estrogenic, or anti-estrogenic properties. It has no aspecicable materia or increding test of extension of the straining hormore, or fluptoxia, nor does it appears to also HTULTIG, clusterare levels. Changes in harizing hormore, (LI) or off-schemizing hormore, (TS) as ado not nationalized in the straining hormore, the straining hormore, and the straining hormore, or also also straining hormore, or shown to increase the criticality levels of the locatorized by roughly 1%, however, since a general mout of the andores is being left multiced by the rockness environ.

History:

The for tease of finisterile in the U.S. was under the heard name of Poscard, Whereld, which was approved by the PDA in 1902. It was specifiadly gave approval for use by patients with hearing prostate hyperpalsis (prostate entragement). In December 1907, the Food and Duq, dhrinstration again approved finasticite, its time the a different propose, proposed, which contained only the a different propose data mare for this propose, Poposedia, Which contained only 150 of the Poscard's dosage. Today, both Proscard's and Propecial's remain the dominant brands of finisterile on the global market.

How Supplied:

Finasteride is most commonly supplied in tablets of 1 mg and 5 mg,

Structural Characteristics:

Finasteride is a synthetic 4-azasteroid. It has the chemical designation 4-azaandrost-1-ene-17-carboxamide, N-(1,1- dimethylethyl)-3-oxo-, (5-,17,)-.

Warnings (Pregnancy):

This drag must never be taken during programs, Frankerick can be absorbed through the skit-Worm on low con origin become program, should never nuclei brocken or uncoated frastserik tubles. The DHT blocking action of frastserik can also severe developmental problems to an uncomma fit site, even is very small arternat. Unlikered frastserik can also intercoarse to huma a developing multi-frasts. The use of condoms or abstinence is recommended during theory.

Side Effects:

Adverse reactions commonly associated with the short- term (1 year) use of finasteride

include impotence (8.1%), decreased libido (6.4%), decreased ejaculate volume located. (3.7%), ejaculation disorder (.8%), gynecomstia (.5%), breast tendemess (.4%), and rash (.5%).

Administration (General Considerations):

Reductions eithbors cannet complexity protect against androgenic side effects such as servoid-instead thin likes, only daki, and a cance. Reductions eithbors lesson these side effects by reducing not eithrating the level of androgenic activity in the siden and seady. Androggnic and anabelic, effects are both middlied by the same receptor, and there is no sourp presently in its abable, the side of the same receptor, and there is no sourp presently in size abable, the side of the hendrice, does not offer complex protection against this side reflex. DBTT inhibition, therefore, does not offer complex protection against this side effect.

Reductuse inhibits are only applicable with testistatome, methylexistorous, and flavorustience. These there dang are concreted to stronger "lightsyle" detaolists by the reductuse enzyme. Naturdrohar and some of is derivative become weaker upon interaction with this corput, as where "lightsyle" methodies hind in androgene receptive varge poorless details with the strongene size effects. Methandosetscohae and boldenen andrego concernist to stronger 5-able reductor attrobustion. End at levels that reductuse inhibitons in the file of the randorganicity. Most other synthesis androbis strongist are constrained to the strongenesis inhibiton.

Administration:

When used medically for the numerat of mule pattern hair loss (and/opericit alphecia) in men, the necommediated doops in large period with sub-soft the trainment of benegative prostate hyperphisis (1974). See magnetic period with a sub-software set of the hyperphisic set of the second set of the second second second second training of the second second second second second second training of the second second second second second second training of the second second

Availability:

Finasteride is widely available in most regions of the world. The most prominent brand names in commerce are Process® (5 mg) and Propecia® (1mg), although a number of other brand and generic forms of the drug can also be located.

Testosterone Stimulating Drugs

hCG (human chorionic gonadotropin)

Description:

Humn Chronice Gondotropin (IGCs) is a prescription medication containing chronice graduotipin oblication an antiral forum (eight Chronice graduotipin is and polypeicki humane normals) fucation in the famile holy during the comparable for increasing the production of progenetores, an propagation of the proinstance of programs by standard co-enchronative grametary metals. For increasing the production of progenetores, and provide the familiarian of programs by standard co-ender programs years that. Block lock of inclusion of programs by standard co-enchronative grametary metals and the co-ender the approximative pro-temperature and the standard standard and co-ender the approximative two is these remotes in gostantication and the homous lock of all designability of the pro-temperature and the standard for polytic standard co-ender the standard standard standard for the polytic standard co-ender the standard Albuch i possesse mirer FSI-like (Folke Stimuling Horneo) activity, the physiologial actions of descrine gamsdarium minih miris from the of the gendometry laterizing hornerse (LH) As a cliental drug, EGG is used as an ecogeneous form of LH it is optical registed to applied to the start and programs of a worms, more special in the comparison of the start and the start and the start and the start and the outline to the ability of LH to stimather the Leydig code in the tests to runnafacture testsenteres. (LG) are lower and the start and the start and the start and disorder drugsheard programs data code and the start and the start and a start and the start and in the tentumer of programsdarian a condition is used in our body of minito the start and the programsdarian and the start and the start and the start to the start and outline the start and the start an

History:

Chroining grandentopin was first discovered in 1920,²⁰ and was identified as a pregnarcy, hormour approximately § spars hite.²⁰⁷⁶ The first drug preparation containing chroining grandottopin care in the form of an arimal pinkary estanct, which was developed as a commercial product by Organon. Organon introduced the extrant care 1931, and the transtance horgen. A trademark dispute forced the company to change the mare Pregot, allowed in the late data of the form of a pinkary extract. Manufacturing torbaginess were instrudeed in 1940 that advord the hormore to be obtained by thering and participat water for the late of the late 1960's were adopted by all manufactures. Structly using multic areas, the late 1960's were adopted by all manufactures formly units of impredations. On the late the late to be obtained by the protocols have been more any preparations on or biological opticipation. The been more more preparations are of biological origin, the risks of biological contaminents are said to be to utilabole contaminents and the biological contaminents are said to be to utilabole contaminents and be to the biological contaminents are said to be to utilabole contaminents and be predicable.

Early on the indicated uses for christic granulotopia prequantiess were much broader than they are presetty. Provide iterative from the 1959 and '03's neuronmendel the user freidukt's systematic corporations in the strength of the strength of the proparation (Education Control 1958) as in '03's responsion, and mulroporation (Education) and the strength of the strength of the the proparation (Education) and the strength of the strength of the the instances. For the strength of the proparation (Education Control 1958) as bring. "These times more effective than instances for the strength of the strength of the strength of the strength of generative strength of the strength of th

HCG has no significant thwoid-stimulating activity. This is specifically pointed out because bCG was once widely used for the treatment of obesity. The trend seemed to have become popular in 1954, after a paper was published by Dr. A.T.W. Simeons claiming that chorionic conadotronin was an effective adjunct to dieting. According to the study natients were able to effectively stave off hunger with severely low-calorie diets provided they took the hormone injections. Dubbed the Simeons diet, people all across the country were soon subjecting themselves to severe calorie restriction (500 calories per day) and taking hCG injections. Soon after, the hormone itself became the main focus for fat loss. In fact, by 1957 it was said that hCG was the most commonly prescribed medication for weight loss. More recent and comprehensive investigations, however, refute that there is any anorexic or metabolic advantage to the use of hCG.769 In 1962, the Journal of the American Medical Association had already been warning consumers about the hCG-inclusive Simeons diet. stating the more basic fact that severe calorie restriction, which causes the body to sacrifice muscle and organ tissue to obtain necessary protein, was more hazardous than obesity itself. By 1974, the FDA had had enough of the hCG fat loss claims, and mandated the following statement to be included with all prescribing literature. "HCG HAS NOT BEEN DEMONSTRATED TO BE EFFECTIVE ADJUNCTIVE THERAPY IN THE TREATMENT OF OBESITY THERE IS NO SUBSTANTIAL EVIDENCE THAT IT INCREASES WEIGHT LOSS BEYOND THAT RESULTING FROM CALORIC RESTRICTION THAT IT CAUSES A MORE ATTRACTIVE OR 'NORMAL' DISTRIBUTION OF FAT. OR THAT IT DECREASES THE HUNGER AND DISCOMFORT ASSOCIATED WITH CALORIERESTRICTED DIETS." This warning persists on all product sold in the U.S. today. In spite of this warning and evidence to the contrary, some clinics still promote the use of hCG for dieting purposes.

Haman Christein Genadotopein is a wickly popular drug preparation today, orwing to the fact that it arrains an indispossible part of volation therapy for turn cycless of familia infattilis. Popular preparations in the U.S. presently include Popular popular topological (Second), and Nozeri (Ferring, although many order trade manus) into beato popular for christica grandstoppi preparations over the years. This drug is also solid widdy outside of the bilited States, and an les forat under many additional thand emuty, to narranso to also have. Owergo in the far that this drug is not cortrolod on a factual level, U.S. attletes and bodyhadius smalle to far a local physical willing to presche the drug to text staredirithcoal Dippontedienco dinto tade the product from interational pharmacy sources. Gener Altopath cortribute from of chronicing guardworph in bee bear introduced to modes in recort years, the wast supply and two cost of biologial L/G cortinas to make it a staple product for both bleded and of biolad use.

Structural Characteristics:

Chorionic gonadottopin is an oligosaccharide glycoprotein composed of 244 amino acids. It has an alpha suburit that is 92 amino acids long and identical to that of luteinizing hormone (LH), folicle-stimulating hormone (FSH), and thyroid-stimulating hormone (TSH). It has a beta suburit that is unique to hCG.

How Supplied:

Hanan Charionic Gonadoropin is widely available in various human and veterinary drug markets. Composition and dosage may vary by courtry and menufacturer, but typically contain 1,000, 1,500, 2,500, 5,000, or 10,000 international units (III) per dose. All forms are sapplied as a lyophilized powder, requiring reconstitution with sterile diluent (water) before use.

Administration (General):

Haman Chorionic Gomadotropin is generally given by intramascular (MM) injection. The subcatameous route is also used, and has been deemed to be roughly equivalent therapeutically to IM injections.⁷³ Peak concentrations of chorionic guradotropin occur approximately 6 hours after intramascular injection, and 16 to 20 hours after subcatameous injection.

Administration (Men):

When used to treat hpogenationpis hpogenation, current FDA-approved protocols recommend efferts a stort 6-worker program or a long-team program leasing up to 1 year, depending on the advelatal means of the particut. Proceeding galakies, for short-term use and the stort of the stort stort of the top of the stort of the top of the stort of the top of the stort of the top of the stort of the top of the stort of the stor

Post-Cycle:

Hamn Chrotica gradudrophi softu and with other medications as part of an in-depth Poor (5)cE Therapy (PCT) program Boeds on restoring endogenous totostrone production more rapidly at the end of a stored syste. Heasing endogenous totostrone endogenous test to stored is showed and the store of the program boet endogenous test to stored is showed approximation and the usy could to all physics. The main concerns is the action of cortisel, which in ramy ways is balanced out by the effect of antiposition. Cortisel as each the opposite message to the muscle store totostatomer, or to breakdown protein in the cell. Left methoded by a low level of totastatomer, control angekild with prath (6) pration ways. The main store test of the store of the ure poss-cycle use or n.c. or generatory can sor use auministration or 2000-2000 Units every caru or 3rd day, taken for no longer than 2 or 3 weeks. If used for too long or at too lingh a dose, the drug may actually function to desensitize the Logdig's cells to laterizing hormone, further hindering a return to homeostasis. For a more comprehensive view of IECG's role in a proper hormonia recovery program please refer to the Post Cycle. Therapy section of this book:

On-Cycle:

Bolyhalkes and athlets may also administer Hamm Chroticei Grandstoppin throughout a storid speck, in an effort to avoid toticatic anrolps and the ensign reduced ability to respond to L1 stirnals. In effect, this practice is used to avoid the problem of toticating analysis, instead off reging to correct it kine on when the cycle is own. It is important to remember that the doagn needs to be cardiffy menitored with fits type of use, at high locks of GCC may cause toroused lockdar aromatices corression fraining entry of the star in the star also descensible the tasks to L11⁴⁷. As such, the dug may actually induce primary hypogendinon when mesod, gard of protecting not a provide print here corression the tasks of the star of the star subcaracteristic star in the star of the correspondence of the star of the star subcaracteristic star of the correspondence of the star of the star of the star subcaracteristic star of the star subcaracteristic star of the star subcaracteristic star of the star subcaracteristic star of the star star of the star of the

These on-cycle IGG protocols were developed by DE. Molt Criske, a well-known figure in the anti-sing and hormone-replenearing Hig. for use with its bestorence replenearies theory (HZ) patients. Although TRE is often administered on a long-term basic, toxicular atophys is a common constric compilar of patients invespetche of the ministenzes of normal anticogen levels. De. Criske's IEGC programs is designed to alloviate this concern in a merrer that is according in hope-CA to Higdan to the Criske Horneset IEGC shoes in relation to a proscibed toxistoseroor replexement program. Do. Criske removements the Bolomy in hope-CA toHigdan to the Criske HOrneset¹¹, and De Criske removements provide the structure of the Higdan to the Criske HOrneset¹¹, and the removement provide the structure of the term of the term of the Higdan to the removement provide the the Higdan to the Criske HOrneset¹¹, and the removement provide the the Higdan to Higdan to the Criske HOrneset¹¹, and the removement provide the the Higdan to Higdan to Higdan to Higdan the removement provide the the Higdan to Higdan to Higdan to Higdan the Higdan Higdan the Intervent the Higdan to Higdan the Higdan the Higdan term of the Intervent term of the Higdan term of t

Administration (Women):

When used to induce ovalation and pregnancy in anovalatory infiritie worman, a does of 5,000 to 10,000 units is administered one day following the last does of menotropins. The timing is specified so that the hormone is given precisely at the right moment in the ovalation cycle. Harman Chorionic Grandottopin is not used by women for physique- or performanceenhancing purposes.

Availability:

When we find I/CL, we see it is always packaged in 2 different viaksampules (one with a powder and the other with a steries solver). These need to be niecds before significant, any leforer drug should be refrigerated for later use. Make sure your product matches this description. Harman Chronic Grondberopin is widely manufactured, and easily obtained on the black marker. To date, counterfiels have not been much of a concern, although a couple of oddings have poped up (all in malf-todes viak).

Drug Profiles Bibliography

^{379.} Experiences with a new testosterone isobutyrate crystal suspension. Drescher H. Dtsch Med Wochenschr. 1952 Apr 4;77(14):431-2.

Enzyme induction by onl testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. Clin Pharmacol Ther (1976) 20:233-237.

^{381.} High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Haman C et al. Metabolism 39(1)

^{1990:69-74}

^{382.} Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab (2001) 281:E1172-81.

^{383.} The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Singh A, Hsia S, et al. J Clin Endocrinol Metab (2002) 87:136-43.

594-2, Methyl and 2, hydrowymethylens, androstane derivatives. Rinaold HL et al. 1 Am Chem Soc 1959;81:477,37

385. Onvmetholone treatment for the anemia of hone marrow failure. Alexanian R. Nadell I. et al. Blood. 1977:

386. Les homones anabolisantes du point de vue esperimental. P.A. Desaulles. Helv. Med. Acta 1960:479-503.

387. Studies on available standide & GCMS characterization of unusual seco acide metabolites of ownerholone in human urine: 1 Steroid Biochem Mol Bio 42 (1997):229.42

388. Fillows of various 17 alpha alkyl substitutions and structural modifications cof steroids on sulfobromonthalein (BSP) renention in rabbits. Lennon HD et al. Steroids 7 (1966): 157-70.

²⁰⁰⁹ Long-term oxymetholone use in HIV patients not associated with significant hepatotoxicity. Hengge UR et al. Poster oresented at the Third International Conference on Natrition and HIV Infection, April 22-25, 1999; Cannes, France

Prince. 390. Effects of an oral androgen on muscle and metabolism in older, community-dwelling men. Schroeder et al. Am J Physiol Endocrinol.Metab. 284:E120–28.

391- Diezfalusy E. Acta endocrin. (Kbh.) 35 (1960):59:

392 Biosynthesis of Estrozens. Gual C. Monto T, Hayano M, Gut M, and Dorfman R. Endocrinology 71 (1962).920-25

393. Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract), Nippon Naibunpi Gakkai Zasshi 62 (1986):18-24

394. Competitive projecterone antapprists: receptor binding and biologic activity of testosterone and 19notestosterone derivatives. Red IR: Humphrey RR: Shih VH. Windsor BI. Sakouski R: Creser PI. Edoren RA Fettil Steril 1979 May 31(5):552,61

395. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafez E. Merrill A. December 1956

396. Different Pattern of Metabolism Determine the Relative Anabolic Activity of 19-Norandrogens. J Steroid Biochem Mol Bio 53(1995):255.7

397. Relative binding affinities of testosterone, 19-nortestosterone and their 5-alpha reduced derivatives to the indicate and to other Relative binding affinities of testosterone. 19-nortestosterone and their S-alpha reduced derivatives to the androaen receptor and to other nortestosterone. Toth M, Zakar T, J Steroid Biochem 17 (1097)-652 (0)

398. Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP Jacue SV Martinez C Blanche P1 Azen S Krauss RM Am J Physiol Endocrinol Metab 2007 Dec 283(6)(F1714.22 Foub 2002 Aug 27

399. Linowie and linowoteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 1986 Jul: 13(7):513-8.

400. The administration of pharmacological doses of testosterone or 19-nortestosterone to normal men is not associated with increased insulin secretion or immained abases tolerance. Karl F. Friefl et al. J. Clin Endocrinol Metab 68: 971 1989

401. Influence of nundrolonedeemonte on the nituitary-annadal axis in males. Bilkma J. Daursma S. Thissen J. Huber O. Acta Endoarinol 101 1982-108 12

402. Oxandrolone: A Potent Anabolic Steroid of Novel Chemical Composition. Fox M. Minot AS and Liddle GW. Journal of Clinical Endocrinology and Metabolism, 1962; Volume 22, Pas. 921-924.

403. M. Fox et al. J. Clin Endocrinol Metab 22 (1962) 921

404. Published reference of personal communication from Saunders F.J. (April 21, 1961) to author of Methyltestosterone, related steroids, and liver function. Arch Int. Med 116 (1965):289-94.

405. Studies on anabolic steroids. II-Gas chromatographic/mass spectrometric characterization of oxandrolone urinary metabolites in man. Masse R, Bi HG,Ayotte C, Dugal R. Biomed Environ Mass Spectrom. 1989 hur 18/60429.38

406. Methyltestosterone, related steroids, and liver function. DeLorimier, Gordan G, Lowe R. et al. Arch Int. Med 116 (1965):289-94

407. Effects of Oxandrolone on Plasma Lipoproteins and the Intravenous Fat tolerance in Man. Atherosclerosis 19 (1974):337-46.

408. Osandrolone and Plasma Triglyceride Reduction: Effect of Triglyceride-Rich Diet and High Density Lipoproteins. Artery 9 (1981):328-41.

409. Plasma and Lipoprotein Lipid Responses to Four Hypolipid Drugs. Lipids 19 (1984):73-79

410. The effects of coundrolose on the growth hormone and goadal axis in boys with constitutional delay of growth and puberty. Malhitra A, Poon E. Et al. Clin Endocrinol (Ouf) 1993 Apr;38(4):393-8.

411- Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

412. Butenandt AK et al. Ber dtsch chem. Ges. 68 (1935):2097.

413. Comparative pharmacokinetics of three doses of percutaneous dihydrotestosterone gel in healthy elderly mena clinical research center study. Wing C. Irannanesh A. Bernan N et al. J Clin Endocrinol Metab 1998 Aug/83(8):2749-57

414. Transdermal dihydrotestosterone treatment of 'andropause'. Ann Med. 1993 Jur;25(3):235-41.

415. Studies on the treatment of idiopathic gynaecomastia with percutaneous dihydrotestosterone. Clin Endocrinol (Oxf), 1983 Oct;19(4):513-20.

416. Gvnecomastia: effect of prolonged treatment with dihydrotestosterone by the percutaneous route Presse Med. 1983 Jan 8;12(1):21-5.

417. Contribution of lymphatically transported testosterone undecanoate to the systemic exposure of testosterone after oral administration of two andriol formulations in conscious lymph duct-cannulated dogs. Shackleford D, Faassen W Aet al. J of Pharm and Exper Ther. 306 (2003):925-933.

418. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237

^{419.} A ten-year safety study of the oral androgen testosterone undecanoate. Gooren LJG , J Androl , 1994 , 15 , 212_5

420. The effects of testosterone treatment on body composition and metabolism in middle-aged obese men. Marin

P. Holming S. Jöhnsson L. et al. Int J Obes , 1992 , 16 , 991-7.

421. Enzyme induction by oral testosterome, Johnsen SG Kammunn JP Bennet FP Journaen F 1976 Clin Pharmacol They 20/233,237

422. Tostostovone replacement, cardiovascular system and risk factors in the aging male. Mgna GB, Bergani E, J Endocrinol Invest, 2005;28(11 Sarol Proceedings):69-74.

423. Evaluation of the nharmacokinetic profiles of the new testosterone topical gel formulation, Testim, compared to AndraGel. Mathury T. Hamill F. Bachard R. Sehrer T. Smith T. Bionharm Droz Disnos. 2003 Anr 24/3):115.

424. Forvorse induction by oral testosterone. Johnsen SG, Kampmann JP, Bernet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-737

425. Testosterone replacement, cardiovascular system and risk factors in the aging male. Mgna GB, Bergami E. J Endocrinol Invest, 2005;28(11 Sarol Proceedings):69-74.

426. Enzyme induction by onal testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20/233,237

427. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Harmon C et al. Metabolicm 20(1) (1000):60 74

428. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281 (2001)/E1172.81

429. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Atam Sinds, Stanley Hsia, et al. J Clin Endocrinol Metab 87 (2002):136-43.

430. Lyster SC, et al. Acta Endocrin (Kbh) 43 (1963):399.

431. The biological activity of 7 alpha-methyl-19-nortestosterone is not amplified in male reproductive tract as is that of testosterone. Endocrinology 1997 here 120(6):2677 92

432. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982. 433. De Visser, J. et al. Acta Endocrin. (Kbh.) 35 (1960):405.

434. Biosynthesis of Estrogens, Gual C, Monato T, Hayano M, Gut M and Dorfman R. Endocrinology 71 (1962):920-25

435. Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract) Ninnon Naihunni Gakkai Zaschi 62 (1986) 18,25

436. Competitive progesterone antagonists: receptor binding and biologic activity of testosterone and 19notestosterone derivatives. Red JR, Humphrey RR, Shih YH, Windsor BL, Sakowski R, Creaer PL, Edaren RA. Fertil Steril 1979 May:31(5):552-61.

437. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafez E, Merrill A. December 1956

438. Taifforost Pattern of Metabolism Determine the Relative Anabolic Activity of 19-Norandrogens. J Steroid Biochem Mol Bio 53:255-7,1995

439. Relative binding affinities of testosterone, 19-nortestosterone and their 5-alpha reduced derivatives to the androgen receptor and to other androgen-binding proteins: A suggested role of Salpha-reductive steroid metabolism in the dissociation of "muotumic" and "andmornic" activities of 19, partestosterone Toth M. Zakar T. I. Semid Biochem 17 (1982):653-60.

440. Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP, Jaque SV, Martingz C, Blanche PJ, Azen S, Krauss RM, Am J Physiol Endocrinol Metab. 2002 Dec:283(6)(E1214-22, Epub 2002 Aug 27,

441. Linemic and Incorroteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 1986 Jul;13(7):513-8.

442. The administration of pharmacological doses of testosterone or 19-nortestosterone to normal men is not associated with increased insulin secretion or immained sharpse tolerance. Karl F. Friedl et al. J. Clin Endocrinol Metab 68: 971, 1989.

443. Influence of nandrolonedecanoate on the pituitary-gonadal axis in males. Bijlsma J,Duarsma S,Thijssen J, Huber O.Acta Endocrinol 101 (1982):108-12.

444. Effect of nandrolone decanoate therapy on weight and lean body mass in HIV-infected women with weight loss. K Mullign, R Zackin, et al. Arch Intern Med. (2005):165:578-85

445. Nandrolone decanoate: Pharmacological properties and therapeutic use in osteoporosis. P Geusens. Clinical Rheumatology, 1995, 14, Sappl. 3.

446. Contraceptive efficacy and adverse effects of testosterone enanthate in Thai men. Sukcharoen N, Aribarg A, Kriangsinyos R, Chanprasit Y, Ngeamvijawat J. J Med Assoc Thai 1996 Dec;79(12):767-73.

447. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237

448. Hish-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Hannan C et al. Metabolism 39(1) (1990):69-74

449. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281 (2001):E1172-81.

450. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Atam Sinds, Stanley Hsia, et al. J Clin Endocrinol Metab 87 (2002):136-43.

451. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237

452. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Harman C et al. Metabolism 39(1) (1990):69-74.

453. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281 (2001):E1172-81.

454. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Sinah A, Hsia S, et al. J Clin Endocrinol Metab 87 (2002):136-43.

455. Biological half-lives of [4-14C]testosterone and some of its esters after injection into the rat. James KC, Nicholle D1 and Doharte M 1 Dharm Dharmood

NUMBER OF STREET, STRE 21 (1969) 24.27

456. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bernet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20/233-233

457. Hish-Awsity linearotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Harnan C et al. Metabolism 39(1)

1000-60 74

458. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281 (2001)/E1172-81

459. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Sinah A, Hsia S, et al. J Clin Endocrinol Metab 87 (2002):136-43.

460. Vischer F. Meystre C. Wettstein A. Hely Chim Acta 38 (1955):1507

461. Never Enough / Steroids in Sports: Experiment turns epidemic. Robert Dvorchak. Pittsburgh Post-Gazette January 14, 2005

462. Comments from Dr. John Ziegler. Strength & Heath Magizine, 1967.

463. Officials bundled steroid resulation from the start. Robert Dvorchak, Pittsburgh Post-Gazette October 3, 2005. 464. Anabelic and wavele strend interaction with rat androaen recentor in vivo and in vitro: a companyive study. Feldkoren BL Andersson S. J. Steroid Biochem Mol Biol. 2005 Apr:94(5):481-7. Epub 2005 Mar 17.

465. Knuskemper. H L, Anabolic Steroids, Academic Press, New York, 1968.

466. Relative importance of 5alpha reduction for the androgenic and LH-inhibiting activities of delta-4-3ketosteroids. Steroids 29 (1997):331-48.

467. Anabolic stemids in clinical medicine. Liddle GW, Burke in: H & Hebertica Medica Acta. 5/6 1960 504,13

468. Effect of anabolic steroid (metandienon) on plasma LH-FSH, and testosterone and on the response to intravenous administration of LRH. Holma P. Adlercreutz. Acta Endocrinol (Copenh) 1976 Deca/83(4):856-64.

469. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

470. Inhibition of the estrogenic activity of methylandrostenediol following administration of aminopterin. Boll Soc Ital Biol Sper. 31(9-10) Sep-Oct (1955):1280-4.

471. Overbeek G A, J. de Visser: Acta endocrin. (Kbh.) 24 (1957):209

472. Biosynthesis of Estrogens, Gual C, Morato T, Hayano M, Gut M and Dorfman R. Endocrinology 71

473. Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract) Ninnon Nahanni Galdai Zaschi 62 (1986):18-25.

474. Competitive projecterone antappnists: receptor binding and biologic activity of testosterone and 19nortestosterone derivatives. Reel JR, Humphrey RR, Shih YH, Windsor BL, Sakowski R, Creger PL, Edgren RA. Fatil Staril 31(5) May (1979):552-61.

475. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafez E. Merrill A. December 1956

476. Different Pattern of Metabolism Determine the Relative Anabolic Activity of 19-Norandrogens. J Steroid Biochem Mol Bio 53 (1995) 255,7

477. Relative binding affinities of testosterone, 19-nortestosterone and the 5-alpha reduced derivatives to the indrogen receptor and to other androgenRelative binding affinities of testosterone. 19-nortestosterone and the 5alpha reduced derivatives to the androgen receptor and to other androgen notestosterone. Toth M. Zakar T. J. Steroid Biochem 17 (1982):653-60.

478. Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP, Jaque SV, Martinez C, Blanche PJ, Azen S, Krauss RM. Am J Physiol Endocrinol Metab. 283(6) Dec (2002) E1214-22. Epub 2002 Aug 27.

479. Lipemic and lipoproteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 13(7) Jul (1986):513-8

480. Influence of nandrolonedecanoate on the pituitary-gonadal axis in males. Bijlsma J,Daarsma S,Thijssen J,Huber O.Acta Endocrinol 101 (1982):108-12

481. Biosynthesis of Estrogens, Gual C, Morato T, Hayano M, Gut M and Dorfman R. Endocrinology 71 (1962):920-25.

482 Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract), Nippon Naibunpi Gakkai Zasshi 62:18-25,1986

483. Competitive progesterone antagonists: receptor binding and biologic activity of testosterone and 19nortestosterone derivatives. Reel JR, Humphrey RR, Shih YH, Windsor BL, Sakowski R, Creger PL, Edgren RA.

484. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafez E. Merrill A. December 1956

485. Different Pattern of Metabolism Determine the Relative Anabolic Activity of 19-Norandrogens. J Steroid Biochem Mol Bio 53/255,7 1995

486 Relative binding affinities of testosterone, 19-nortestosterone and their 5-alpha reduced derivatives to the androgen receptor and to other androgen-bidning proteins: A suggested role of Salpha-reductive steroid metabolism in the dissociation of "myotropic" and "androgenic" activities of 19- notestosterone Toth M. Zakar T. J Steroid Biochem 17 (1982):653-60.

487. Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP, Jaque SV, Martinez C, Blanche PJ, Azen S, Krauss RM. Am J Physiol Endocrinol Metab. 2002 Dec:283(6):E1214-22. Epub 2002 Aug 27.

488. Linemic and Incorroteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 1986 Jul: 13(7):513-8.

489. The administration of pharmacological doses of testosterone or 19-nortestosterone to normal men is not associated with increased insulin secretion or impaired alucose tolerance. Karl E. Friedl et al. J Clin Endocrinol Metab 68: 971, 1989.

490. Influence of nandrolonedecanoate on the pituitary-gonadal axis in males. Bijkma J.,Duarsma S,Thijssen J,Huber O.Acta Endocrinol 101 (1982):108-12.

491. Biocompasis of Estroams, Gual C. Monto T, Hayano M, Gut M and Dorfman R. Endocrinology 71 (1967) 970 25

492. Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract) Ninnon Naiburni Galdai Zasshi (abstract): Nip 62-18,25 1986

493. Competitive progesterone antagonists: receptor binding and biologic activity of testosterone and 19notestosterone derivatives. Reel JR, Humphrey RR, Shih YH, Windsor BL, Sakowski R, Creaer PL, Edaren RA, Fertil Steril 1979 May:31(5):552-61.

494. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafey F. Merrill A. December 1956

495. Different Pattern of Metabolism Determine the Relative Anabolic Activity of 19-Norandrogens. J Steroid Biochem Mol Bio 53:255-7.1995.

496. Relative binding affinities of testosterone, 19-nortestosterone and their 5-alpha reduced derivatives to the adrogen receptor and to other androgen-bidning proteins: A suggested role of Salpha-reductive steroid metabolism in the dissociation of "myotropic" and "androgenic" activities of 19- notestosterone Toth M. Zakar T. J Steroid Biochem 17 (1982):653-60.

497. Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP, Jaque SV, Martinez C, Blanche PJ, Azen S, Krauss RM. Am J Physiol Endocrinol Metab. 2002 Dec 283(6)(F1714.22 Foub 2002 Aug 27

498. Lipemic and lipoproteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 1986 Jul: 13(7):513-8.

499. The administration of pharmacological doses of testosterone or 19-nortestosterone to normal men is not associated with increased insulin secretion or impaired glucose tolerance. Karl E. Friedl et al. J Clin Endeerinol Metab 68: 971 1989

500. Influence of nandrolonedecanoate on the pituitary-gonadal axis in males. Bijlsma J,Duarsma S,Thijssen J, Huber O.Acta Endocrinol 101 (1982):108-12.

501. Biosynthesis of Estroagns, Gual C, Morato T, Hayano M, Gut M, and Dorfman R. Endocrinology 71 (1962) 920-25.

502. Metabolism of boldenone in marc gas chromatographic/mass spectrometric identification of urinary excreted metabolites and determination of excretion rates. Scharzer, Donike, Bol Mass Spec, 21 (1992):3-16.

503. J. Mathieu. Proc. Intern. Symp. Drug Res. 1967, p 134. Chem. Inst. Can., Montreal, Canada.

504. Unique stemid congruens for recentor studies. Olasco. Rowmand. Cancer Research 38 (1978):4186-98

505. Characterisation of the affinity of different anabolics and synthetic hormones to the human androzen receptor. human sex hormone binding dobulin and to the bovine progestin receptor. Bauer, Meyer et al. Acta Pathol Microbiol Immol Scand Sapol 108 (2000):838-46.

506. Unique steroid congeners for receptor studies. Ojasoo, Raynaud. Cancer Research 38 (1978):4186-98.

507. Disposition of 17 beta-tresholone in humans. Spranger, Metzler. J Chromatogr 564 (1991):485-92.

508. Cholestasis induced by Parabolan successfully treated with the molecular adsorbent recirculating system. Anand JS et al. ASAIO 2006. JanFeb:52(1):117-8

509. Herr, M E, Hogg J A, Levin R H, J Am Chem Soc. 78, 500 (1956)

510. Lyster SC. Lund GH, and Stafford RO, Endocrinology 58, 781 (1956)

511. Testing for flaosymesterone (Halotestin®) administration to man: Identification of urinary metabolites by gas chomatography-mass spectrometry, Karmerer R. Mardink J. Jangels M et al. J Steroid Biochem 36 (1990)659-

512. Eisenberg, E. Modern Trends in Endocrinology (H. Gardiner-Hill, ed) p 46. Hoeber, NY (1961)

513. Methyltestosterone, related steroids, and liver function, deLorimier A,Gilbert G, et al. Arch Intern Med v116 (1965):289,94

514. The offices of fluoromesterone administration on testicular function. Jones TM, Fang VS et al. J Clin Endocrinol Metab 1977 Jar(44(1):121-9.

515. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science CEnsultants, Natick, MA 1982

516. Biosynthesis of Estrogens, Gual C. Morato T. Havano M. Gut M and Dorfman R. Endocrinology 71 (1962):920-25

517. Aromatization of androstenedione and 19-nortestosterone in human placental, liver and adipose tissues (abstract). Nippon Naibunpi Galdai Zasshi 62-18,25 1986

518. Competitive progesterone antagonists: receptor binding and biologic activity of testosterone and 19-nortestosterone derivatives. Red JR, Humphrey RR, Shih YH, Windsor BL, Sakowski R, Creger PL, Edgren RA. Fertil Steril 1979 May:31(5):552-61.

519. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G, Chang M, Zarrow M, Hafez E, Merrill A. December 1956

520. Different nattern of metabolism determine the relative anabolic activity of 19-norandrogens. J Steroid Biochern Mol Bio 53/255/7 1995

521. Relative binding affinities of testosterone, 19-nortestosterone and their 5-alpha reduced derivatives to the indrogen receptor and to other androgen-binding proteins: A suggested role of Saloha-reductive steroid metabolism in the dissociation of "myotropic" and "androgenic" activities of 19- nortestosterone Toth M, Zakar T. J Steroid Biochem 17 (1982):653-60.

522 Metabolic effects of nandrolone decanoate and resistance training in men with HIV Sattler FR, Schroeder ET, Dube MP, Jaque SV, Martinez C, Blanche PJ, Azen S, Krauss RM. Am J Physiol Endocrinol Metab. 2002 Dec:283(6)(E1214-22, Epub 2002 Aug 27,

523. Lipemic and lipoproteinemic effects of natural and synthetic androgens in humans. Crist DM, Peake GT, Stackpole PJ. Clin Exp Pharmacol Physiol 1986 Jul: 13(7):513-8.

524. The administration of pharmacological doses of testosterone or 19-nortestosterone to normal men is not associated with increased insulin secretion or impaired glucose tolerance. Karl E. Friedl et al. J Clin Endocrinol Metab 68: 971, 1989.

525. Inflores of nurdrolonedecanoate on the pituitary-spinadal axis in males. Bijlsma J,Duarsma S,Thijssen J, Huber O.Acta Endocrinol 101 (1982):108-12.

526. 2-Methyl and 2-hydroxymethylene-androstane derivatives. Rinapid HJ et al. J Am Chem Soc 1959;81:427-32. 527. Binauld H. L et al. L org Chem. 21 (1956):1432

528. Herstellung des 17-metyl-testosterons und anderer Androsten- und Androstanderivate. Ruzicka L, Goldberg MW. Et al. Hely Chim Acta 18 (1935):1487-98.

529. Effect of anabolic steroids on liver function tests and creatine excretion. Manquardt G. H. et al. JAMA 175 (Mar 11, 1961):851-3

530. Motholto-stosterone, related steroids, and liver function. deLorimier A, Gilbert G, et al. Arch Intern Med v116 (1965):289-94

531. Hish-density Incorotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl KE, Haman CI Ir. Jones RF. Phymate SR. Metabolism 1990 Jan 39(1):69-74.

532. Anabolic Steroids and Sports Volume II, James E, Wright, Sports Science Consultants, Natick, MA 1982. 533. Bucicka L et al. Hely Chim. Acta 18 (1935) 1487

534. Anabolic Steroids and Sports Volume II, James E, Wright, Sports Science Consultants, Natick, MA 1982

535. Liver toxicity of a new anabolic agent: methy trienolone (17-alpha-methyl-4.9, 11-estrutriene-17 beta-ol-3-one). Kruskemper, Noell, Steroids, 1966 Jul 2011:13-24

536. T. Fevel-Cabanes, Compt. Rend. Soc. Biol. 157, 1428 (1963)

537. Protein anabolism produced in man by a new steroid: methyltrienolone. Tremolieres J, Pequignot E. Presse Med. 1965 Nov 6c73(47):2655-8

538. Specific binding of [3H]-methyltrienolone to both progestin and androgen binding components in human benign specific bilang of [Srij-neity interioue to both progen and indrogen bilang components in italian being mostatic hypertrondy (BPH) Asselin 1 Melancon R. Goundeau V Labrie F. Barne C. Raynard IP. 1 Steroid Biochem 1979 May 10(Sr483.6

Liver toxicity of a new anabolic agent: methylrhienolone (17-alpha-methyl-4,9,11-estratriene-17 beta-ol-3-one). Kruskemper, Noell. Steroids. 1966 Jul;2(1):13-24.

540. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982

541. M. Shimizu, G. Ohta et al. Chem. & Pharm. Bull. (Tokyo) 13,895 (1965)

542. Pharmacological studies on experimental nephritic rats. (4) Improvement of hyperlipernic models in rats utilizing anti-rat kidney rabbit serum and effects of anti-hyperlipemic agents on serum lipid levels. Suzuki Y, Honda Y, Ito M. Jpn J Pharmacol. 1978 Oct;28(5):729-38.

543. The use of coundrolone in hyperlinidemia Dovie AF. Pinkus NB. Green J. Med J. Aust. 1974 Feb 2:1(5):127.

544. Clinical evaluation of a new anabolic agent 7a,17a-dimethyltestosterone (bolasterone). Korst, D. R., Bowers, C. Y. Flokstra, J. H., McMahon, F.G. Clin, Pharmacol, Thenn, 4:734-9 (1963).

545. The synthesis of some 7alpha and 7beta methyl steroid hormones. Campbell JA, Babcock JC. J Am Chern Soc 81:4069.74 (1959)

546. Anabolic and androgenic activities of 7a,17a-dimethyltestosterone (U-19,763), a new anabolic steroid. Stucki J. C. Duncan G. W. Lyster SC.

547. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

548. NERIDO Prescribing Information (2004). Schering AG Germany.

549. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F 1976 Clin Pharmacol Ther 20:233-237

550. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Karl Friedl, Charles Hannan et al. Metabolism 39(1)

1000-60 74

551. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

552. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Atam Singh, Stanley Hsia, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

553. Colton F.B., Nysted L.N., Riegel B., et al. J Am Chem Soc. 79 (1954), 1123

554. Studies of the biological activity of certain 19-nor steroids in female animals. Pincus G., Chang M., Zarrow M., Hafez E, Merrill A. December 1956.

555. Anabolic Steroids and Sports Volume II. James E. Wright, Sports Science Consultants, Natick, MA 1982.

556. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F 1976 Clin Pharmacol Ther 20:233-237

557. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Karl Friedl, Charles Haman et al. Metabolism 39(1) 1000-60 74

558. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

599. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolyopproteins, and C-reactive protein in healthy young men. Atam Singh, Stanley Hsia, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

560. Metabolism of anabolic steroid drugs in man and the marmoset monkey (callithrix jacchus)-I. Nilevar and Orabolin, Ward R, Lawson A.M., Shackleton C.L.H. J Steroid Biochem 8 (1977):1057-63.

561. De Winter, M. S. et al. Chem. and Ind. (London) 1959, 905.

562. Overbeek G.A. et al. Acta Endicrin. (Kbh) 40 (1962):133.

563. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

564. Doemer G and Schubert A. Proc. Intern. Congr. Hormonal Steroids, Milan 1962, Excerpta Med. Intern. Congr. Ser. No. 51, 210

565. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

566. Miescher, Wettstein & Tschopp (1936) Schweiz. Med. Wschr. 66, 310.

567. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F 1976 Clin N._____ TTL _ 30.997 9

Pharmacol Ther 20235-257.

568. Hish-density lipontotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Hannan C et al Metabolism 39(1) 1990

(9.74

 Testosterone dose-response relationships in healthy young men. Blusin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001

570. The effects of varying docs of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in heithv young men. Atam Singh, Stanley Hsia, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

571. J. Mathieu Proc. Intern. Symp. Drug Res. 1967, p 134. Chem. Inst. Can., Montreal. Canada.

572. Unique steroid congeners for receptor studies. Ojasoo, Raynaud. Cancer Research 38 (1978):4186-98

573. Characterisation of the affinity of different anabolics and synthetic hormones to the human androgen receptor. human sex hormone binding dobulin and to the bovine progestin receptor. Bauer, Meyer et al. Acta Pathol Microbiol Immol Scand Sapol 108 (2000):838-46.

574. Unique steroid congeners for recentor studies. Olasoo. Raynaud. Cancer Research 38 (1978):4186-98.

575. Disposition of 17 beta-trenholone in humans. Summer: Metzler: 1 Chromatour: 564 (1991):485-97

576. Cholestasis induced by Parabolan successfully treated with the molecular adsorbent recirculating system. Anand JS et al. ASAIO 2006. JanFeb:52(1):117-8

577. Mischort R et al Chem Ber 93 (1960):1710

578. Motowokne acetate. Summary of information for clinical investigators, New Branswick, NJ.The Spabb Institute for Medical Research, May 30, 1962

579. Anabolic effects of methenolone enanthate and methenolone acetate in underweight premature infants and children New York State Journal of Medicine March 1 1965 645.8

580. Kruskemper H L et al, Esc Medica (Amsterd.) Congr. Ser. No. 51 (1962), 209.

581. Weller O Arzneimittelforsch 12 (1967) 234

582. Proc. Intern. Const. Hormonal Stenaids. Milan. 1967. Excentia Med. Intern. Const. Ser. No. 51. n. 209 Eventa Med Found Amsterdam 1967

583. Failure of non-17-alleviated anabolic steroids to produce abnormal liver function tests. J Clin Endeerinol Metab. 1964 Dec:24:1334.6

584. Fatal outcome of a patient with severe aplastic anemia after treatment with metenolone acetate. Ann Hematol. 1993 hi/6711/d1,3 Taikamoto N Li/biyama T Takyuchi T Sato S Nanise T Nakazato Y

585. Comparative studies about the influence of metenoloneacetate and mesterolone on hypophysis and male sonads. Trenkner R. Sense T. Hierz H et al. Arzneimittelforschung. 1970 20(4):545-7.

586. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982. 567. Wisshert R.et al. Chem Ber, 93 (1960):1710

568. Methenolone enanthate, Summary of information for clinical investigators, New Branswick, NJ. The Spabb Institute for Medical Research, April 15 1962

569. Anabolic effects of methenolone enanthate and methenolone acetate in underweight premature infants and children. New York State Journal of Medicine March 1, 1965, 645-8.

570. Proc. Intern. Congr. Hormonal Steroids, Milan, 1962. Excepta Med. Intern. Congr. Ser No. 51, p. 209. Exerota Med Found, Amsterdam 1962

571. Failure of non-17-alkylated anabolic steroids to produce abnormal liver function tests. J Clin Endocrinol Metab 1964 Dec:24:1334.6

572. Relative binding affinity of anabolic-androgenic steroids: comparison of the binding to the androgen receptors in skeletal muscle and in prostate, as well as to sex hormone-binding alobulin. Saartok T. Dahlberg E. Gustafsson JA, Endocrinology, 1984 Jurc114(6):2100-6.

573. Inflatence of various modes of androgen substitution on serum lipids and lipoproteins in hypogenadal men. Jockenhovel F, Bullmann C, Schubert M, Vogel E, Reinhardt W, Reinwein D, Muller-Wieland D, Krone W. Metabolism 1999 May 48/5) 590.6

574. Comparative studies about the influence of metenolonacetate and mesterolone on hypophysis and male sonads. Trenkner R. Sense T. Hienz A. et al. Arzneim-Forsch. (Drug Res) Jahrang 30, Nr. 4 (1970):545-7.

575. The effects of mesterolone, a mile sex hormone in depressed patients (a double blind controlled study). Itil TM, Michael ST, Shapiro DM, Itil KZ. Methods Find Exp Clin Pharmacol. 1984 Jung6(6):331-7.

576 Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F 1976 Clin Pharmacol Ther 20233-237.

577. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Karl Friedl, (harles Hannan et al. Metabolism 39(1)

1990/69.74

578. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81 2001

579. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolypoproteins, and C-reactive protein in healthy young men. Singh A, Hsia S, et al. J Clin Endocrinol Metab 87:136-43, 2002.

580. Enzyme induction by onal testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol They 20/233-237

581. Testosterone replacement, cardiovascular system and risk factors in the aging male. Vigna GB, Bergani E. J Endocrinol Invest, 2005;28(11 Sarol Proceedings):69-74.

582 Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237

583. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Karl Friedl, Charles Hannan et al. Metabolism 39(1) 1000 60 74

584. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

585. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men Sinsh & Heia S et al. 1 Clin Endoyrinol Metab 87: 136-43, 2007

586. Product Data Sheet: Sustanon 250. August 31, 2001. Pharmaco (N.Z.) LTD Auckland New Zealand.

587- Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237.

588. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Haman C et al. Metabolism 39(1) 1990:

69-74.

²⁰⁹⁷. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

590. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Singh A, Hsia S, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

591- History of diethylstilbestrol use in cattle A. P. Raun and R. L. Preston. 2002, American Society of Animal Science.

992. Long-term experience with testosterone replacement through scrotal skin. In Niesching E and Behre HM (eck) Tostosterone. Action, Deficiency, Substitution. Atkinson LE, Chang Y-L and Snyder PJ (1998) Springer, Berlin, Germany 365-388.

503: Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bernet EP, Jorgensen F. 1976 Clin Pharmacel Ther 20233-237.

594. Testosterone replacement, cardiovascular system and risk factors in the aging male. Vigna GB, Bergani E. J Endocrinol Invest. 2005;28(11 Suppl Proceedings):69-74.

 New androgen for retarded action; testosterone phenylpropionate. Klotz HP, Avril Y. Therapie. 1955;10(4):588-50.

596 Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237.

997. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Human C et al. Metabolism 39(1) 100/00/31.

1990:69-74

598. Testosterone dase-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

599. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Singh A, Hsia S, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

600. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237.

 Testostrone replacement, cardiovascular system and risk factors in the aging male. Mgna GB, Bergami E. J Endocrinol Invest. 2005;28(11 Suppl Proceedings):69-74.

602. The effects of depot testosterone therapy on sarum levels of luteinizing hormone and follide-stimularing hormone in patients with Klinedster's syndrome and hypogonadotrophic euracholdism. Fukutani K, Isuragi K et al. J Clin Endcenicol Metai 3938/664.

603. Enzyme induction by oral testosterone. Johnsen SG, Kampmann JP, Bennet EP, Jorgensen F. 1976 Clin Pharmacol Ther 20:233-237.

604. High-density lipoprotein cholesterol is not decreased if an aromatizable androgen is administered. Friedl K, Human C et al. Metabolism 39(1) 100/00/31.

1990:69-74

605. Testosterone dose-response relationships in healthy young men. Bhasin S, Woodhouse L et al. Am J Physiol Endocrinol Metab 281: E1172-81, 2001.

606. The effects of varying doses of T on insulin sensitivity, plasma lipids, apolipoproteins, and C-reactive protein in healthy young men. Singh A, Hsia S, et al. J Clin Endocrinol Metab 87: 136-43, 2002.

607. J. Mathieu, Proc. Intern. Symp. Drug Res. 1967, p 134. Chem. Inst. Can., Montreal, Canada.

608. Unique steroid congeners for receptor studies. Ojasoo, Raynaud. Cancer Research 38 (1978):4186-98.

609. Characterisation of the affinity of different anabolies and synthetic hormones to the human androgen receptor, human sex hormone binding globulin and to the bovine progestin receptor. Bauer, Meyer et al. Acta Pathol Microbiol Humand Scand Suppl 108 (2000):838-466.

610. Unique steroid congeners for receptor studies. Ojasoo, Raynaud. Cancer Research 38 (1978):4186-98.

611. Disposition of 17 beta-trenbolone in humans. Spranger, Metzler. J Chromatogr 564 (1991):485-92.

612. Cholestasis induced by Parabolan successfully treated with the molecular adsorbent recirculating system. Anand JS et al. ASAIO 2006. JanFeb;52(1):117-8.

613. Clinton R. O. et al. J. Amer chem. Soc. 81 (1959):1513.

614. U.S.Patent # 3,030,358.

615. Anabolic-androgenic steroid interaction with rat androgen receptor in vivo and in vitro: a comparative study. Feldioren BL Andersson S. J Steroid Biochem Mol Biol. 2005 Apr:94(5):481-7. Epub 2005 Mar 17.

616. The differential effects of stanozolol on human skin and synovial fibroblasts in vitro: DNA synthesis and receptor binding, Ellis AJ, Cawston TE, Madcie EJ. Agents Actions. 1994 Mar/41(1-2):37-43.

617. Identification of a specific binding site for the anabolic staroid stanozolol in nule rate liver microsomes. Boada LD, Fernandez I. et al. J Pharmacol Exp Ther 1996 Dec;279(3):1123-9.

618. Stanozolol and danazol, unlike natural androgens, interact with the low affinity glucocorticoid-binding sites from male nt liver microsomes. Formandez I, Chinno R, Boada LD, Navarro D, Cabrera N, del Rio I, Diaz-Chico BN. Endocrinology. 1994 Mar;13(3):1401-8.

619. Desaulies P.A.et al.Helv. Med Acta 27 (1960), 479.

620. Sex hormone-binding globulin response to the anabolic steroid stanozolot: Evidence for its suitability as a Biological androgen sensitivity test. G Sinnecker, S Kohler. Journal of Clin Endo Metab. 68: 1195,1989.

621. The influence of 6 months of oral anabolic steroids on body mass and respiratory muscles in undernourished COPD patients. Ivone Martins Ferreira, leda Verreschi et al. CHEST 114 (1) July 1998 19-28.

622 Androgenic/Anabolic steroid-induced toxic hepatitis. Stime D, Milic S, Dintinjam RD, Kovac D, Ristic S J Clin Gistroenterol. 2002 Oct;35(4):350-2.

623. Contrasting effects of testosterone and stanozolol on serum lipoprotein levels. Thompson PD, Cullinane EM,

Sady SP. Chenevert C. Saritelli AL. Sady MA. Herbert PN. JAMA. 1989 Eds 24:261(8):1165;8

624. The effect of intramuscular stanogolol on fibrinolysis and blood linids. Smill M. McAndle BM. Louie GD. Forbes (T) Provide CR. Thromb Res. 1982 Oct.

1:28(1):27,36

123(1)27-30.
625. Alteration of hormone levels in normal males given the anabolic steroid stanozolol. Small M, Beastall GH, Semple CG. Cowan RA, Forbes CD. Clin Endocrinol (Oxf) 1984 Jul 21(1):49-55.

626. Anabolic Steroids and Sports Volume II. James E. Wright. Sports Science Consultants, Natick, MA 1982.

627. The effect of stanoardol on nitrogen retention in the dog. Olson ME, Morek DW, Ouinn KB, Can J Vet Res. 2000 Oct;64(4);246-8

628. Protein synthesis in isolated forelinb muscles. The possible role of metabolites of arachidonic acid in the response to intermittent stretching. Smith Palmer et al. Biochem J. 1983 214 153-61

629. The influence of changes in tension on protein synthesis and prostaglandin release in isolated rabbit muscles. Palvow Reeds et al. Biochem J. 1983

214.1011-14.

630. Protein synthesis and degradation in isolated mascle. Effect of n3 and n6 fatty acids. Palmer, Wahle. Biochem I 1987 242 615-18

631. Romber ownice modulates mascle membrane phospholipid profile in rats. Helse et al. J. Nutr. 1999 129:1636-

632. Exercise training reduces skeletal muscle membrane anchidonate in obese (fu/fa) Zucker rat. Ayre et al. J. Appl. Physiol. 1998 85(5):1898-1902.

633. Effects of physical exercise on phospholipid fatty acid composition in skeletal mascle. Andersson et al. Am. J. Physiol 274 (Endocrinol Metab

37rE437.38 1998

634. Effects of an ormivorous diet compared with a lactoovovegetarian diet on resistance-training-induced changes in body composition and skeletal muscle in older men. W Campbell, M Barton et al. Am J Clin Natr 70(1999):1032-9.

635. The effect of dietary arachidonic acid on plasma lipoprotein distribution, apoproteins, blood lipid levels, and tissue fatty acid composition in humans. G.J. Nelson, P.C. Schmidt et al. Lipids 32: 427-33 (1997).

636. The effect of dietary arachidonic acid on platelet function, platelet fatty acid composition, and blood consulation in humans. Nelson CJ, Schmidt PC et al. Lipids 32(4):421-5 (1997).

637. Effects of dietary arachidonic acid on human immune response. Kelley DS, Taylor PC et al. Lipids 32(4):449-\$6(1997)

638. Dietary intake of N-3 and N-6 fatty acids and the risk of prostate cancer. M Leitzmann, M Stampfer et al. Am J Clin Nutr 80:204-16 (2004)

639. Dietary (n-6) PUFA and intestinal tumorigenesis. J Whelan, M McEntee et al. J Natr 1343421S-268 (2004).

640. Action of herstaminol hydrochloride on contractile properties in flog isolated twitch muscle fibre. Allard B, Jacquemond V, Lenttiri-Chlich F, Pourrias B, Rougier O. Br J Pharmacol. 1991 Nov;104(3):714-8.

641. Heptaminol chlorhydrate: new data. Pourrias B. Ann Pharm Fr. 1991;49(3):127-38.

642. On the mode of action of heptaminol (author's transl)] Grobecker H, Grobecker H. Azzneimittelforschung, 1976/26(17)/2167.71

643. Prostaglandins and the control of muscle protein synthesis and degradation. Prostaglandins Leukot Essent Fatty Acids, 1990 Feb;39(2):95-104

644. Prostadandin F2(alpha)stimulates growth of skeletal muscle cells via an NFATC2-dependent pathway. Horsley V Paylath GK 1 Cell Biol 2003 Apr 14: 161(1):111.8

645. Stretch-induced prostaglandins and protein turnover in cultural skeletal muscle Mandenburgh HH, Hatfaludy S, Sohar I, Shansky J, Am J Physiol, 1990 Aug259(2 Pt 1): C232-40.

646. Skeletal muscle PGF(2)(alpha) and PGE(2) in response to eccentric resistance enercise: influence of ibuprofen acetaminophen. Trappe TA, Flackey JD, White F, Lambert CP, Evans WJ. J Clin Endocrinol Metab. 2001 Oct;86(10):5067-70

645. Prostagandins promote and block adipogenesis through opposing effects on perovisome prolifentor-activated receptor gamma. Reginato MJ, Krakow SL, Bailey ST, Lazar MA. J Biol Chem 1998 Jan 23;273(4):1855-8

646. Preclinical pharmacology of "Arimides" (anastrozole; ZD1033)--a potent, selective aromatase inhibitor. J Steroid Biochem Mol Biol 1996 Jul 58(4):439-45

647. Anastrozole alone or in combination with tamovifen versus tamovifen alone for adjunctive treatment of postmenopausal women with early breast cancer. Frist results of the ATAC randomized trial. Lancet 2002;

648. History and Advancement of Anastrozole in the Treatment of Breast Cancer. Edited by Aman Buzdar & Michael Baum, RSM Press, February 2003

649. Inhibition of adrenal corticosteroid synthesis by aninoplutethimide: Studies on the mechanism of action. Dester RN FishmanI M. Nev RC et al. J Clin Endocrinol 27 (1967) 473-80

650. First generation aromatase inhibitors -aminoglutethimide and testololactone. Cocconi G. Breast Cancer Res Treat 1994;30(1):57-80

651. Stereoselective inhibition of aromatase by enantiomers of aminoplatethimide. Graves PE, Sulhanick HA. Endocrinol 105 (1979) 52-57

652. Beneficial effects of raloxifene and tamovifen in the treatment of pubertal gynecomastia. Lawrence SE, Faught et al. J. Pediatr. 2004 Jul: 145(1):71-6

653. Effects of miosifene on gonadotropins, sex hormones, bone turnover and lipids in healthy elderly men. Eur J Endocrinol. 2004 Apr;150(4):539-46

654. Ralovifene decreases serum IGF-1 in male patients with active acromegaly.Dimanki EV, Symons KV,Barkan AL, Eur J Endocrinol. 2004 Apr;150(4):481-7

655. Fulvestrant, Formerly ICI 182,780. Is as Effective as Anastrozole in Postmenopausal Women With Advanced Breast Cancer Progressing After Prior Endocrine Treatment, Howell A, Robertson JFR, Ougresma Albano J, Aschermannova A. et al. J Clin Oncol. 2002; 1:57.

656. Fulvestrant, an estrogen receptor downregulator, reduces cell tumover index more effectively than tamovifen. Anticoncor Roc. 2007 Ind Acception 2016 (2017) In

AUDICIDEST PUSS 2002 2007000220(4):2311-7.

657. Fulvestrant. Cheung KL, Robertson JF. Expert Opin Investig Drugs 2002 Feb;11(2):303-308

658. Effects of estrogen on the release of gonadotropins and prolactin in male pseudohemaphrodites. Barbarino A, De Darinis L et al. J endocrinol Invest.

1979 Jan-Mar;2(1):41-4

 Estrogen-dependent plasma prolactin response to grandotropin-releasing homone in intact and castrated men. Barbarino A, De-Marinis L. et al. J Clin Endocrinol Metab. 1982 Dec;55(6):1212-6

60. Effects of progesterone administration on folicle-stimulating hormone and prolactin release in estrogen treated expanded adult men. Marcini A, De Marinis L. et al. Andrologia 1991 Sep-Oct;23(5):373-9

661. Inhibitory effect of androgen on estrogen-induced prolactin messenger ribonucleic acid accumulation in the male nat americe pinutary. Tong Y et al. Endocrinology. 1989 Oct. 122(4):1821-8

662. Effects of depot testosterone administration on serum levels of testosterone, FSH, LH and prolactin. Ruiz E. et al. J Endocrinol Invest. 1980 OctDec;3(4):385-8.

663. Response of serum homones to androgen administration in power athletes. Alen M. Reinila M. et al. Med Sci Sports Exerc. 1985 Jun;17(3):354-9.

64. Effect of androgenic anabolic steroids on sperm quality and serum hormone levels in adult male bodybuilders. Torres-Caligi J. et al. Life Sci. 2001 Mar 2;68(15):1769-74.

665. Anabolic steroid-associated hypogenalism in male hemodialysis patients. Maeda Y et al. Clin Nephrol. 1989 Oct;32(4):198-201.

666. Prescribing Information for Dostinex Tablets. Pharmacia & Upjohn 2003.

667. Enhancement of linear growth and weight gain by cyproheptadine in children with hypopituitarism receiving growth hormone therapy. Kaplowitz PB, Jennings S. Pediatr. 1987 Jarc110(1):140-3.

 Megestrol acetate vs cyproheptadine in the treatment of weight loss associated with HIV infection. Sammerbell CD, Youle M, McDonald V, Catalan J, Gazzard BG. Int J STD AIDS. 1992 Jul-Aug3(4):278-80.

669. Cyproheptadine no longer promoted as an appetite stimulant. WHO Drug Info 1994; 8:66

670. Effects of anethum graveolens and garlie on lipid profile in hyperlipidenic patients. Kojari J, Vosoughi AR, Alzarri M. Lipids Health Dis. 2007 Mar 1;65.

671. Epsets of gafie estruct consumption on blocd lipid and oxidant/antioxidant parameters in humans with high blocd cholesterin Damk I, Kawataa M, Ayrag B, Asci A, Devrim E, Ozhek H, Ozilak HS J Narr Biochem. 2004 June 35(0):773-7.

672.8

⁶⁷³. Effect of ground green ten drinking for 2 weeks on the susceptibility of plasma and LDL to the oridation ex vivo in hashby volumeers. Gomikuwa S, Ishikuwa Y, et al. Kobe J Med Sci. 2008 May 23;54(1):E62-72.

674. Cardiovascular protective effects of resventrol. Bradamante S, Barenghi L, Vilia A. Cardiovasc Drug Rev. 2004 Fall:22(3):169-88.

675. Phytosterols/stanols lower cholesterol concentrations in fimilial hypercholesterolemic subjects: a systematic review with meta-analysis. Moraisi KG, Oosthuizen W, Opperman AM. J Am Coll Natr. 2006 Feb;25(1):41-8. Review.

676. Role of policosanols in the prevention and treatment of cardiovascular disease Variady KA, Wang Y, Jones PJ. Natr Rev. 2003 Nov;61(11):376-83. Review.

677. Role of selenium in cytoprotection against cholesterol oxide-induced vascular damage in rats. Huang K, Liu H, Chen Z, Xu H. Atherosclerosis. 2002 May;162(1):137-44.

678. Inositol heuniacinate. Altern Med Rev. 1998 Jun;3(3):222-3.

679. Pownall JH et al., Atheroscherosis, 1999; 143:285-297

680. Harris WS et al. J Cardio Risk 1997; 4:385-391

681. Clinical overview of Omacor: a concentrated formulation of omega-3 polyunsaturated fatty acids. Bays H. Am J Cardiol. 2006 Aug 21;98(4A);71i-76i. Epub 2006 May 30.

682. Gissi P Study, The Lancet 1999; Vol 354:447-55

683. Omega-3 firsty acid supplementation accelerates chylomicron triglyceride clearance. Park Y, Harris WS. J Lipid Res. 2008 Mar;44(3):455-63. Epub 2002 Dec 1.

684. The hypotrighyceridemic effect of eicosupertaenoic acid in rats is reflected in increased mitochondrial fatty acid oxidation followed by diminished lipogenesis. Willamsen N, Skorve J, Heseberg S, Rustan AC, Berge RK. Lipids. 1993 Aug 28(9):683-30.

 Consult-exercic acid (DHA) and hepatic gene transcription. Jump DB, Botolin D, Wang Y, Xu J, Demeure O, Christian B, Chem Phys Lipids. 2008 May;153(1):3-13. Epub 2008 Feb 23.

686. Why do omega-3 fatty acids lower serum triglycerides? Harris WS, Bulchandani D. Curr Opin Lipidol. 2006 Aug;17(4):287-93.

687. Effects of modulinil ingestion on exercise time to exhaustion. Jacobs I, Bell DG. Med Sci Sports Exerc. 2004 Jun;36(6):1078-82.

688. A double-blind clinical trial in weight control. Use of ferthaumine and phentermine alone and in combination. Weintraub M, Haeday JD, Mushlin AL, Lockwood DH. Arch Intern Med. 1984 Jun;144(6):1143-8

689. Modalities of the food intake-reducing effect of sibutramine in humans. Chapelot D, Manuenier C, Thomas F, Hanotin C, Physiol Behav. 2000 Jarc68(3):299-308.

690. Thermogenic effects of sibuttamine and its metabolites. Connoley IP, Liu YL, Frost I, Reckless IP, Heal DJ, Stock MJ. Br J Pharmacol. 1999 Mar;126(6):1487-95.

691. A randomized, double-blind, placebo-controlled, multicenter study on silvatramine in over-weighted and obese subjects. Zhao Y, Wang X, Yan Z. Zhonghua Yu Fang Yi Xue Za Zhi. 2001 Sep;35(5):329-32.

692. Effects of ketotifien and elembatered on beta-adrenergic receptor functions of lymphocytes and on plasma TXB-2 levels of asthrunic patients. Huszar E, Herjavezz I et al. Z Erke Atmangsorgane 1990;175(3):141-6

693. Effect of prednisolone and ketotifen on beta 2-adrenoceptors in asthmatic patients receiving beta 2bronchofilitors. Brodde OE. Howe U et al. Ear J Clin Pharmacol 1988;34(2):145-50

Ketotifen alone or as an additional medication for the long-term control of asthma and wheeze in children. Bassler D, et al. The Cochrane Database of Systematic Reviews; Issue 1. Chichester: John Wiley; 2004.

695. The emergence of endocrinology. Welbourn RB. Gesnerus. 1992;49 Pt 2:137-50.

696. Why do we continue to write for Syntheoid? Volume XI, Issue 8 Harold J. DeMonaco, M.S., Director of Drag Therapy Management, A publication of the Drug Therapy Committee, Massachusetts General Hospital and the MAssachusets General Physicians Oranization.

697. Bioexuivalence of generic and brand-name levothyrosine products in the treatment of hypothyroidism Dong BI Hands WW Gambertoolio IG Gee L White IR Buhn IL Greensnan FS JAMA 1997: 277:1205-1213

698. Detergent effects of sodium deoxycholate are a major feature of an injectable phosphatidylcholine formulation used for localized fat dissolution. Rotunda AM, Suzuki H et al. Dermatol Surg. 2004 Jul;30(7):1001-8

699. Phosnhatidvkholine increases the secretion of triacylglycerol-rich lipoproteins by CaCo-2 cells. Mathar SN, Bom E et al. Biochem 1. 1996 Mar 1/314 (nt 7)/569,75

700. Lipolysis injection with phosphatidylcholine (Lipostabil N) examination of blood values after subcutaneous administration. Dr. Franz Hasenschwandtner, Online Article

701. Clinical experience and safety using phosphatidy leholine injections for the localized reduction of subcutaneous fat: a multicentre retrospective UK study lournal of Cosmetic Dermatology 5(3)/218-276. Sentember 2006 Palmer Mark MD Curran John MD Bouder Patrick MD

702. Stimulation of growth hormone secretion with human growth hormone releasing factors (GRF1-44, GRF1-40, GRF1-49) in normal subjects. Losa M, Schopohl J, Müller OA, von Werder K. Klin Wochenschr. 1984 Dec. 240(02)(1140.2

703. Testing with growth hormone-releasing factor (GRF(1-29)NH2) and somatomedin C measurements for the evaluation of growth hormone deficiency. Ranke MB, Gruhler M, Rosskamp R, Brümman G, Attanasio A, Blum WF, Bierich JR, Eur J Pediatr. 1986 Dec:145(6):485-92.

704. Once daily subcataneous growth hormone-releasing hormone therapy accelerates growth in growth hormonedeficient children during the first year of thermany. Genef International Study Group. Thomas M. Bochiccioli P. Colle-M, Lanes R, Grunt J, Galazka A, Landy H, Eengrand P, Shah S. J Clin Endocrinol Metab. 1996 Mar/81(3):1189-96.

705. Growth hormone responses to growth hormone-releasing hormone (1-29)-NH2 and a D-Ala2 analog in normal men. Barron JL, Coy DH, Millar RP, Pentides, 1985 May-Juno6(3):575-7.

706. A comparative study of growth hormone (GH) and GH-releasing hormone(1-29)-NH2 for stimulation of A comparative study of grown number (Ori) and Ori-releasing number 1-29-Nrt2 for simulation or much in children with GH deficiency. Chen RG, Shen YN, Vei J, Wang CF. Xie DH, Wang XH, Zhou JD, Chen (Y Wu VI, Gumarsson R, et al. Acta Paediate Sunni 1993 Mar 388-32,5' discussion 36

707. Semorelin: a review of its use in the diagnosis and treatment of children with idiopathic growth hormone deficiency. Prakash A. Goa KL, BioDruss.

1999 Aug 12/21:139-57

708. Continuous subcutaneous GHRH(1-29)NH2 promotes growth over 1 year in short, slowly growing children. Brain CE, Hindmarsh PC, Brook CG, Clin Endocrinol (Osf.), 1990 Edv32(7):153-63

709. Once daily subcataneous growth hormone-releasing hormone therapy accelerates growth in growth hormonedeficient children during the first year of therapy, Geref International Study Group. Thomer M. Rochiccioli P. Colle M. Lanes R. et al. J Clin Endocrinol Metab. 1996 Mar:81(3):1189-96

710. Growth hormone (GH) profiles in response to continuous subcutaneous influsion of GH-releasing hormone(1-290.NH2 in children with GH deficiency. Tauber MT Pienkouski C. Pioron P. Catalii M. Rochicvioli P. Acta Paediatr Sannl 1993 Mar 388:28-30: discussion 31

711. Growth response to growth hormone-releasing hormone(1-29)-NH2 compared with growth hormone. Neyzi O, Yordam N et al. Acta Paediatr Suppl.

1993 Mar;388:16-21; discussion 22

712. Growth Hormone-Releasing Hormone in HIV-Infected Men With Lipodystronhy A Randomized Controlled Trial, Polyami Koutka: Bridart Canavan: Jeff Brea: et al. JAMA, July 14, 2004-Vol 292, No. 2.

713. Somatomedin-C mediates growth hormone negative feedback by effects on both the hypothalamus and the nituitary. Berdowitz M. Szabo M, Frohman LA, Firestone S, Chu L, Hintz RL. Science. 1981 Jun 12/212(4500):1279.81

714. Semorelis: a better approach to management of adult-onset growth homone insufficiency? Walker RF. Clin Intery Aging 2006;1(4):307-8.

715. EMD Serono, Inc. to Discontinue Geref Diagnostic 8/5/2008

716. Effects of single nightly injections of growth hormone-scleasing hormone (CHRH 1-29) in healthy elderly men. Vintone J, Blackman MR, Bashy Whitehead J. et al. Metabolism. 1997 Jan;46(1):89-96.

717- Section. 303(e)(1)/FDCA, 21 U.S.C. 333(e)(1)

718. Treatment of pituitary dwarf with human growth hormone. Raben MS. J Clin Endocrinol Metab 18:901-903 1059

719. Long term mortality in the United States cohort of pituitary-derived growth hormone recipients. J Pediat 144:430-436

720. Pioneering recombinant spowth hormone manufacturing pounds produced per mile of height. J Pediat 131:55-

721. Production of Authentic Recombinant Somatropin. Linda Fryklund. Acta Paediatrica Volume 76 Issue s331 Page 5-8, January 1987.

Figs >8, summy res., 722. Five-year follow-up of growth hormone antibodies in growth hormone deficient children treated with recombinant human growth hormone. G Massa et al. Clin Endocrinol 38(2) 137-42 (Feb 1993).

723. Identification of an insulin-responsive element in the promoter of the human gene for insulin-like growth factor binding protein-1. J Biol Chem

268:17063-68.1995

724. Evidence supporting a direct suppressive effect of growth hormone on serum IGFBP-1 levels. Experimental studies in normal, obese and GH-deficient adults. Growth Hormone and IGF Research 9:52-60,1999

725. Growth hormone induced increase in serum IGFBP-3 level is reversed by anabolic steroids in substance abusing power athletes. Clin Endocrinol (Oxf.)

726. United Kingdom multicentre clinical trial of somatrem. Milner RD, Barnes ND, Buckler JM, Carson DJ, Hadden DR, Hughes IA, Johnston DI, Parkin JM,

727. Antigenicity and efficacy of authentic sequence recombinant human growth hormone (somatropin): first-year experience in the United Kinadom, Buzi F. Buchanan CR, Morrell DJ, Preece MA, Clin Endocrinol (Osf) 1989 May:30(5):531-8

778 million and build a second s

— Proneering recommant growth normone manufacturing: pounds produced per mate of neight. J Pediat 131:55-57.

729. Metformircits botanical background. CJ Bailey, C Day. Pract Diab Int April 2004 Vol. 21 No. 3

 Du nouveau dans les antidiabetiques. La NN dimethylamine guaryl guanide (N.N.D.G.). Sterne J. Marce Med 1957; 36: 1295-96.

131. Ghaophage, Ghaophage XR Prescribing Information. Revision January 2009. Brystol-Myers Squbb Company.

732 Protein anabolic actions in the human body. W.M. Bennet et al. Diabetic Medicine 1991, 8 199-207

733. Effect of physiologic hyperinsulinemia on skeletal muscle protein synthesis and breakdown in man. R Gelfand et al. J Clin Invest. 80, July 1987 1-6

734. Insulin action on muscle protein kinetics and amino acid transport during recovery after resistance exercise. G Biolo. B Williams et al. Diabetes Vol 48. May 1999 949-957.

735. Insulin signaling and insulin sensitivity after exercise in human skeletal muscle. Jorgen F. P. Wojtaszewski et al. Datheres Vol 49, March 2000 325-31

⁷³⁶ Intramacular injection of insulin lispro or soluble human insulin: pharmacokinetics and glucodynamics in Type 2 diabetes. Z. Milleceic et al. Diabetes UK. Diabetic Medicine, 18 562-566

²³⁷. Hypotensive effects of hawthom for patients with diabetes taking prescription drugs: a randomised controlled trial. Walker AF, Marakis G, Simpson E, Hope JL, Robinson PA, Hassanein M, Simpson HC. Br J Gen Pract. 2006 Juny 665(27):437-43.

Jauszter Josev, June 2019, 2019 Jan Stranger, Jan Stranger, Jan Stranger, Jan Stranger, Jan Stranger, Santos-Buelga C, Osma A. Pharmacol Rep. 2009 Jan Setts (1):167–75.

739. Effects of nattokinase on blood pressure: a randomized, controlled trial. Kim JY, Gum SN et al. Hypertens Res. 2008 Aug31(8):1583-8.

740. Effects of garlic on blood pressure in patients with and without systolic hypertension: a meta-analysis. Reinhart KM, Coleman CJ, Teevan C, Vachhari P, White CM. Ann Pharmacother. 2008 Dec;42(12):1766-71. Epub 2008 Nov 18.

⁷⁴¹. Prevention of hypertension, cardiovascular damage and endothelial dysfunction with green tea estracts. Antonelio M, Montemarro D, Bolognei M, Di Pascoli M, Piva A, Grego F, Sticchi D, Giuliani L, Garbisa S, Rossi (P. Ann J Hypertens. 2007 Dec;2012):1213-8.

⁷⁴². Effect of Ginkgo bilohu on blood pressure and incidence of hypertension in edderly men and women. Brinkley TE, Lorato JF, Arnold AM, Furberg CD, Kuller LH, Barke GL, Nahin RL, Lopez OL, Yasar S, Williamson JD; Ginkgo Evaluation of Memory (GEM) Study Investigators. Am J Hypertens. 2010 May 22(5):528-33. Epub 2010 Feb 18.

⁷⁴³. Blood pressure lowering effect of an olive leaf estract (Olas europaea) in L-NAME induced hypertension in rats. Khayyal MT, el-Ghazaly MA, Abdallah DM, Nassar NN, Okpanyi SN, Kreater MH. Azzneimittelforschung. 2005;21(1):797-802.

744. Role of coenzyme Q10 (CoQ10) in cardiac disease, hypertension and Meriers-like syndrome. Kumar A, Kaur H, Devi P, Mohan V. Pharmacol Ther.

2009 Dec;124(3):259-68. Epub 2009 Jul 25.

745. The role of endogenous GHRH in arginine, insulin, clouidine- and 1-dopa-induced GH release in normal subjects. Hanew K, Utsumi A. Eur J Endocrinol. 2002 Feb;146(2):197-202.

746. Effect of chronic cloridine administration on GH secretion in adult human subjects. Cataria A, Baldini M et al. Exp Clin Endocrinol. 1991 Mar;97(1):76-80.

¹⁴⁷ Effect of choosic cloudine treatment on urinary growth hormone estretion and linear growth in children with short stature. Summari A, Lucas A, Granda ML, Salinas I, Reverter JL, Cuatrecasas JM, Foz M, Audi L. Horm Res. 1990;34(5):61193-6.

⁷⁴⁸ A new antihypertensive drug clouidine or Catapressan. Amery A, Bossaert H, Deruyttere M, Verstreken G. Acta Clin Belg 1989;24(6):334-55.

^{749.} Medicine and Drugs: Clonidine. Medicinenet.com/clonidine/article.htm. Omadhome Ogbru, Jay W. Marks, MD.

⁷⁵⁰. Multivitamins and phospholipids complex protects the hepatic cells from androgenic-anabolic-steroids-induced toxicity, Pagnisi TA, Koukoulis GN, Hadjichristodoulou CS, Toli PN, Angelepoulos NV. Clinical Toxicel (2008) 46, 57-66.

37.1. Hepatoprotective effects of Liv-52 on ethanol induced liver damage in rats. Indian J Exp Biol. 1999 Aug;37(8):762-6.

752 The effect of the heptoprotective agent LIV 52 on liver damage. Cas Lek Cesk. 1997 Dec 17;136(24):758-60.

753. Hepatoprotective effect of Liv-52 and kumaryasava on carbon tetrachloride induced hepatic damage in rats. Indian J Exp Biol. 1997 Jur;35(6):655-7.

754. Role of Liv-52 in protection against beryllium intoxication. Biol Trace Elem Res. 1994 Jun;41(3):201-15.

755. Alcohol hangover and Liv.52. Chauhan BL, Kulkarni RD. Eur J Clin Pharmacol. 1991;40(2):187-8.

756. Hepstoprotective effects of Liv-52 on ethanol induced liver damage in rats. Sandhir R, Gill KD. Indian J Exp Biol. 1999 Aug;37(8):762-6.

 Himalaya's Liv-52/LiverCare Approved as a Herbol Drug in Switzerland. Hoastan (BW HealthWire) April 4, 2002.

758. Anti-hepatotoxic effects of root and root callus estracts of Cichorium intybus L. Zafar R, Mujahid Ali S. J. Ethnopharmacol. 1998 Dec;63(3):227-31.

759. Aqueous extract of Terminalia arjuna prevents carbon tetrachloride induced hepatic and renal disorders. Manna P, Sinha M, Sil PC. BMC Complement Altern Med. 2006 Sep 30;6:33.

760. Achillea millefolium L. s.I. revisited: recent findings confirm the traditional use. Benedek B, Kopp B. Wien Med Wochenschr. 2007;157(13-14):312-4.

761. N-acetyl-cysteine protects liver from apoptotic death in an animal model of fulminant hepotic fulture. San-Migael B, Alvarez M, Calebras JM, González-Gallego J, Tufón MJ. Apoptosis. 2006 Nov;11(11):1945-57.

762. Toxic effects of anabolic-androgenic steroids in primary nat hepotic cell cultures. Welder AA, Robertson JW, Meichent RB. J Pharmacol Toxicol Methods. 1995 Aux334:187-95.

763. An undated systematic review with meta-analysis for the clinical evidence of silvmarin. Saller R. Brianoli R.

Metzer J, Meier R. Forsch Komplementmed. 2008 Feb: 15(1):9-20.

764. A suffrequencies and operative structures the Nrt2-dependent detoxification pathway. Morimitsu Y, Nakagawa Y et al. J Biol Chem. 2002 Feb

1;277(5):3456-63.Epub 2001 Nov 12

⁷⁶⁵. Marked suppression of dihydrotestosterone in men with benign prostatic hyperplasin by datasteride, a dual 5alpha reductuse inhibitor. Clark RV, Hermann DJ, Curningham GR et al. J Clin Endocrinol Metrab 2004 May/389(5):2179-84

766 Unique preclínical characteristics of GG745, a potent dual inhibitor of 5AR. Branson HN, Hermann D, Bachelor KW, Lee FW, James MK, Fryc SV. J Pharmacol Exp Ther. 1997 Sep;282(3):1496-502.

767. Ecogenous stimulation of corpus lateum formation in the rabbit; influence of estracts of human placenta, decidua, fetus, hydatid mole and corpus lateum on the rabbit ganad. Hirose T 1920 J Jpn Gynecol Sot 16:1055.

⁷⁰⁸. Die Schwangerschaftsdiagnose ausdem Ham durch Nachweis des Hypophysenvorderlappen-hormone. II. Prackrishe und theoretische Ergebreisse aus den harmattersuchungen. Ascheim S, Zondek B 1928 Klin Wechenschr 7:1453-1457.

769. The effect of human chorionic ganadotropin (HCG) in the treatment of obesity by means of the Simeons therapy: a criteria-based meta-analysis. Lijesen GKS et al.Br J Clin Pharmacol 1995; 40: 237–43.

770. A madornized three-way cross-over study in healthy pituitary-suppressed women to compare the bioaraibility of human chorionic gonadotrophin (Prograp) after intramacular and subcutaneous administration. Marmaeris BM, Gearts TB, Odnik J. Hum Reprod. 1988 Jun;13(6):1461-4.

771. Acute stimulation of aromatization in Leydig cells by human chorionic gonadotropin in vitro. Proc Natl Acad Sci USA 76:4460-3,1979.

772. The different mechanisms for suppression of pituitary and testicular function. Sandow J, Engelbart K, von Rechenberg W.Med Biol. 1986;63(5–6):192-200.

Glossary

AAS - Anabolic/Androgenic Steroid

Acute - Short duration. An acute effect is one that occurs rapidly, not after long-term administration.

Aerobic – Refers to a process where oxygen is used to generate energy in the muscles from carbohydrates, fats, and protein, in the form of ATP (adenosine triphosphate). Long distance running is an example of an aerobic activity.

Adipose - Fat tissue.

Adrenoceptor – A type of receptor in the body involved in the regulation of heart rate, metabolism, and thermogenesis. Stimulated by endogenous catecholamines such as epinephrine (adrenaline), noreignephrine (noradrenaline), and dopamine.

Agonist - A substance that initiates a biological response. An estrogen agonist acts as an estrogen in the body.

Ampule – A glass container that holds a single dosage unit of a liquid drug product. An ampule must be broken open before use.

Anabolic – A process that involves the building of tissues such as muscle and bone. AAS are most valued by bodybuilders and athletes for their anabolic properties.

Anaerobic – Refers to a process where energy is generated in the muscles from carbohydrates in the form of ATP (adenosine triphosphate) without the use of oxygen. Weight lifting is an example of an anaerobic activity.

Androgenic – Refers to the musculinizing properties of a substance. AAS stimulate male libido, secondary hair growth, acne, and male pattern hair loss via their androgenic properties.

Antagonist – A substance that inhibits a biological response. An estrogen antagonist blocks the action of estrogen in the body.

Arrhythmia - An irregular heartbeat. Arrhythmias may be life threatening or benign in nature.

Atherogenic - Promoting the formation of plaque deposits on the walls of arteries.

Atherosclerosis - A progressive cardiovascular disease characterized by the buildup of plaque deposits in the arteries. This may obstruct blood flow, causing heart attack or stroke.

Bacteriostatic – Inhibits the growth of bacteria. Bacteriostatic water contains ingredients that prevent bacteria from contaminating the liquid.

Contraindicated - Not advisable for use. A contraindication is a condition that would prevent someone from using a particular drug product.

Diabetogenic - Increases blood sugar.

Diastolic - The phase of blood circulation where the pumping chambers of the heart (ventricles) are being filled. Pressure is at its lowest during the diastolic phase.

Downregulate – To reduce in number. Some cellular receptors downregulate with high levels of drug stimulation, inducing tolerance. Anabole/androgenic steroids generally do not cause a reduction in respective androgen receptor concentrations. Classic downregulation does not occur.

Edema - The retention of excess water in the circulatory system and/or body tissues.

Endogenous - Occurring naturally within the body. Testosterone produced by the testes is an endogenous hormone.

Epiphyses – The growth plates at the end of long bones. The epiphyses plates regulate increases in linear height during development, and fuse at maturity preventing further linear growth.

Erythropoiesis - The process in which red blood cells are produced in the body. Anabolic/androgenic steroids can stimulate erythropoiesis.

Esterified – Refers to a steroid compound that has one or more fatty acids attached to the molecule, usually to show its release from an injection site. Testosterone cypionate is an esterified form of testosterone.

Exogenous - Caused by an agent outside the body.

Glycemic - Relating to blood sugar levels.

Hepatotoxic - Liver toxic. All c-17 alpha alkylated anabolic/androgenic steroids are considered hepatotoxic.

Homeostasis - A state of equilibrium among physiological processes.

Hyperplasia - Growth that occurs via an increase in cell number.

Hypertrophy – Growth that occurs via increases in existing cell size. Anabolic/androgenic steroids produce growth through hypertrophy.

Metabolic Syndrome – A poorly defined common metabolic disorder characterized by abdominal obesity, an atherogenic lipid profile, insulin resistance, elevated blood pressure, a proinflammatry state, and increased risk of cardiovascular disease.

Multi-dose Vial – A vial with a rubber stopper on the top, designed to be pierced repeatedly by a needle (for multiple uses).

Pathological - Involving or caused by physical disease.

Peripheral — Near the surface of the body. Psychoactive drugs are often regarded as centrally acting (brain/central nervous system), while hormones lke anabolic steroids affect both central and peripheral issues (such as muscle and skin).

Prognostic - Serving to predict the likely outcome of a disease.

Pulmonary - Related to the lungs.

Recombinant – Refers to a synthetic manufacturing technology that involves the splicing of genes or DNA segments and inserting them into a cell culture in order to replicate a specific protein. Recombinant DNA technology is used to manufacture many protein-based drug products including human growth hormore, insulin-like growth factors, and human insulin.

Selective - Describes a drug with a very specific effec, and little spillover into other biological systems.

Subcutaneous - Located beneath the skin and above the muscle.

Supraphysiological – In excess of normal biological levels. High doses of testosterone produce supraphysiological levels of hormone in the blood.

Supratherapeutic - In excess of normally defined therapeutic levels.

Systemic - Affecting the entire body through general circulation.

Systolic - The phase of blood circulation where the pumping chambers of the heart (ventricles) are actively pumping. Pressure is at its highest during the systolic phase.

Upregulate – To increase in number. Usually relating to cellular receptor concentrations. Anabolicandrogenic steroids can increase respective androgen receptor concentrations, possibly increasing sensitivity to androgens.