Therapeutic Class Overview Androgens (testosterone)

Therapeutic Class

Overview/Summary: The topical testosterone products listed in Table 1 are approved by the Food and Drug Administration for testosterone replacement therapy in males with primary hypogonadism (congenital or acquired) or hypogonadotropic hypogonadism (congenital or acquired) with testosterone pellets also having an indication to stimulate puberty in carefully selected males with clearly delayed puberty. 1-9 There are few differences between the topical testosterone products with the exception of formulation and site of administration. Androderm® is the only testosterone product available as a transdermal patch. AndroGel[®], Fortesta[®], Testim[®], and Vogelxo[®] are available in gel preparations, while Axiron® is formulated as a topical solution. These products are available as metered-dose pumps or single-use packets/tubes. Striant® is a mucoadhesive buccal tablet system that is placed on the gum for 12 hours and applied twice a day, once in the morning and once in the evening. Testopel[®] is an implantable pellet that consists of crystalline testosterone. It is a cylindrically shaped pellet, 3.2mm (1/8 inch) in diameter and approximately 8-9mm in length. When implanted subcutaneously, the pellet(s) slowly release the hormone over three to six months for a long acting androgenic effect. Androderm® is applied at night, while the topical gels and solution are generally applied in the morning. 1-9 A higher incidence of skin pruritus is associated with the transdermal patch compared to the topical gels; however, the use of hydrocortisone cream, may reduce skin irritations that develop. The labeling of testosterone solution and gels include a Black Box Warning regarding the risk of virilization of female sexual partners that has been reported with male use of topical testosterone gels and solution.²⁻⁷ The occlusive backing film on Androderm[®] prevents the partner from coming in contact with the active material in the system, and therefore the warning is not included on this product. Currently, only AndroGel® has an A-rated generic formulation.

Hypogonadism refers to a defect of the reproductive system resulting in a lack of gonad function. Hypogonadism is classified based on the level of the defect within the reproductive axis. Primary hypogonadism results from a defect of the gonads and occurs when the serum testosterone concentration and/or sperm counts are below normal, and the serum luteinizing hormone (LH) and/or follicle-stimulating hormone (FSH) concentrations are above normal. Secondary hypogonadism, known as hypogonadotropic hypogonadism, results from defects in the hypothalamus or pituitary. This occurs when the serum testosterone concentration and/or sperm counts are below normal, and the serum LH and/or FSH concentrations are normal or reduced. Combined primary and secondary hypogonadism may occur and results in below-normal testosterone concentrations and variable LH and/or FSH concentrations, depending upon which clinical condition predominates. Male hypogonadism may manifest as testosterone deficiency with or without infertility. Clinical signs and symptoms depend primarily on the age at the onset of the condition. Postpubertal hypogonadism usually results in slowly evolving clinical manifestations that may include a progressive decrease in muscle mass, loss of libido, impotence, oligospermia or azoospermia, poor concentration, and an increase in the risk of osteoporosis and fractures.

Table 1. Current Medications Available in the Therapeutic Class 1-9

Generic (Trade Name)	Food and Drug Administration Approved Indications	Dosage Form/Strength	Generic Availability
Testosterone	Hypogonadism in males, primary	Androderm [®] :	
(Androderm [®])	(congenital or acquired) and	2 mg/day patch	
	hypogonadotropic hypogonadism in	4 mg/day patch	-
	males (congenital or acquired)		
Testosterone	Hypogonadism in males, primary	AndroGel [®] 1%:	
(AndroGel ^{®*})	(congenital or acquired) and	Metered-dose pump:	
	hypogonadotropic hypogonadism in	12.5 mg testosterone/actuation	а
	males (congenital or acquired)		





Generic (Trade Name)	Food and Drug Administration Approved Indications	Dosage Form/Strength	Generic Availability
(made mame)	7.pp. 0.704	Unit-dose packet: 50 mg testosterone/packet	7 country
		AndroGel® 1.62%: Metered-dose pump: 20.25 mg/actuation	
		Unit-dose packet: 20.25 mg/packet	
Testosterone (Axiron®)	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired)	Axiron®: Metered-dose pump: 30 mg/actuation	
Testosterone (Fortesta®)	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired)	Fortesta [®] : Metered-dose pump: 10 mg/actuation	-
Testosterone (Striant®)	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired)	Striant®: Buccal mucoadhesive system: 30 mg	-
Testosterone (Testim [®])	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired)	Testim [®] 1%: Unit-dose tubes: 50 mg/tube)	-
Testosterone (Testopel [®])	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired); stimulate puberty in carefully selected males with clearly delayed puberty	Testopel [®] : Implantable pellet: 30 mg	-
Testosterone (Vogelxo [®])	Hypogonadism in males, primary (congenital or acquired) and hypogonadotropic hypogonadism in males (congenital or acquired)	Vogelxo [®] : Metered-dose pump: 12.5 mg/actuation Unit-dose packet:	_
		50 mg/packet Unit-dose tube: 50 mg/tube	

^{*}A-rated generic available in at least one dosage form or strength

Evidence-based Medicine

- Topical and miscellaneous testosterone products have been evaluated in several clinical trials. 18-30
- The safety and efficacy of Striant® (testosterone buccal tablet) was evaluated in a 12 week, open-label, multicenter, phase III clinical trial involving 98 hypogonadal men. At the conclusion of the trial, 86.6% of patients with sufficient data for full analysis had mean serum testosterone concentration values within the physiologic range. The mean (± standard deviation) serum testosterone concentration at the end of the study was 520 (±205) ng/dL compared with a mean of 149 (±99) ng/dL at baseline.8





- The clinical trials evaluating the safety and effectiveness that were used to obtain FDA approval of testosterone pellets are not available. However, a literature search identified a phase IV clinical trial by Kaminetsky et al. Mean testosterone significantly increased and luteinizing hormone (LH) levels significantly decreased from pre-implantation values at week one, week four and week 12 visits, and had returned to pre-implantation levels by week 24 (P<0.001 for mean testosterone and LH levels at week one, week four and week 12 visits; P=0.58 and P=0.87 for mean testosterone and LH at week 24 respectively). Prostate-specific antigen levels remained unchanged for the duration of the study.¹⁸
- Several clinical studies have shown that the transdermal patch and gels all restore serum testosterone
 concentrations to within normal limits and maintain sexual characteristics, sexual behavior, mood, and
 muscle development, and improve bone mineral density in hypogonadal men. The results of these
 head-to-head trials favored the use of the gel over the patch.
- In an open-label study, Axiron[®] topical solution applied to the axilla provided a serum testosterone level in the normal range for 84.1% of patients after 120 days of treatment.¹⁷ Results from a second openlabel study reported that 76.2% of men achieved a mean serum testosterone level within the normal physiologic range following 35 days of treatment with Fortesta[®].²⁶
- In an open label extension study Kaufman et al evaluated efficacy of testosterone 1.62% gel up to one year of therapy.²⁹ Results from the study show that testosterone 1.62% is effective in replacement therapy with 78% (95% CI, 70.0% to 84.6%) and 87.0% (95% CI, 66.4% to 97.2%) of the different dosing regimens reaching therapeutic levels of testosterone.
- Blick et al evaluated the use of testosterone replacement therapy in human immunodeficiency virus infection/acquired immune deficiency syndrome (HIV/AIDS). In this prospective cohort study the effects of replacement therapy with testosterone 1% (Testim®) were evaluated in HIV/AIDS patients. During the twelve month study, but non-HIV/AIDS patients and HIV/AIDS cohorts had significant increases in total testosterone and free testosterone to within normal limits along with increased sexual function and improved and decreased antidepressant use. Body composition profiles improved significantly in men without HIV/AIDS (P≤0.05) and remained stable in men with HIV/AIDS during the twelve months of follow-up. ³⁰
- A meta-analysis of 16 studies evaluating testosterone supplementation for the diagnosis or erectile dysfunction was conducted by Jain et al. The overall response rate was 57% ± 2.3% (203 of 356 cases). Among the studies with stratified results, 75 of 117 (64% ± 4%) men with a primary etiology responded and 53 of 120 (44% ± 2.9%) men with a secondary etiology responded, which was determined to be statistically significant (P<0.001).</p>

Key Points within the Medication Class

- According to Current Clinical Guidelines 13-16:
 - o Intramuscular and topical testosterone preparations are generally recommended for the management of hypogonadism in adult male patients.
 - The oral alkylated androgens are not recommended due to poor androgen effects, adverse lipid changes, and hepatic side effects, but may be considered when other agents are not suitable.
 - The selection of testosterone replacement therapy should be a joint decision between the
 patient and physician and should be made after consideration of patient preferences, the
 pharmacokinetic profiles of the respective agents, treatment burden and cost.
 - The short-acting preparations may be preferred over long-acting depot preparations when initiating treatment in patients with late-onset hypogonadism due to the potential development of an adverse event that may require rapid discontinuation of testosterone replacement therapy. Treatment guidelines do not recommend one topical preparation over another.

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Therapeutic Class Review Androgens (testosterone)

Overview/Summary

Testosterone products are available in a number of dosage forms including oral administration, intramuscular injection, topical gel, transdermal patch, a topical solution, a subcutaneous implantable pellet and a buccal delivery system. This review will focus on the topically administered testosterone products including Androderm®, AndroGel®, Axiron®, Fortesta®, Striant®, Testim® and Vogelxo® and the implant pellet Testopel®. All of these products are approved by the Food and Drug Administration (FDA) for testosterone replacement therapy in males with primary hypogonadism (congenital or acquired) and hypogonadotropic hypogonadism (congenital or acquired) with Testopel® also being indicated for stimulation of puberty in males who clearly have delayed puberty. All testosterone products are controlled substances and have all been assigned as Schedule III products.¹⁻⁹

Hypogonadism refers to a defect of the reproductive system resulting in a lack of gonad (testes) function. 11-15 Hypogonadism is classified based on the level of the defect within the reproductive axis. Primary hypogonadism results from a defect of the gonads and occurs when the serum testosterone concentration and/or sperm counts are below normal, and the serum luteinizing hormone (LH) and/or follicle-stimulating hormone (FSH) concentrations are above normal. ¹² Secondary hypogonadism (hypogonadotropic) results from defects in the hypothalamus or pituitary and occurs when the serum testosterone concentration and/or sperm counts are below normal, and the serum LH and/or FSH concentrations are normal or reduced. 12 Combined primary and secondary hypogonadism may occur, and results in below-normal testosterone concentrations and variable LH and/or FSH concentrations, depending upon which clinical condition predominates.¹⁴ Male hypogonadism may manifest as testosterone deficiency with or without infertility. As a result, appropriate disease classification is necessary since fertility can be restored with appropriate androgen stimulation in individuals with secondary hypogonadism, but not in most individuals diagnosed with primary hypogonadism. ¹⁴ Clinical signs and symptoms depend primarily on the age at the onset of the condition. Postpubertal hypogonadism usually results in slowly evolving clinical manifestations that may include a progressive decrease in muscle mass, loss of libido, impotence, oligospermia or azoospermia, poor concentration, and an increase in the risk of osteoporosis and fractures. 11-16

There are few differentiating factors between the topical testosterone products with the exception of formulation and site of administration. Androderm® is the only testosterone product that is available as a once-daily transdermal patch that is applied at night. AndroGel®, Testim®, Fortesta® and Vogelxo® are available in gel preparations and Axiron® is formulated as a topical solution. These products are available as meter-dosed pumps and single-use tubes and are all applied once daily, generally in the morning. Striant® is formulated as a buccal mucoadhesive system that is placed on the gum for 12 hours and applied twice a day, once in the morning and once in the evening. Testopel® is a pellet that consists of crystalline testosterone. It is cylindrically shaped, 3.2mm (1/8 inch) in diameter and approximately 8 to 9 mm in length. When implanted subcutaneously, the pellet(s) slowly release the hormone for a long acting androgenic effect. A higher incidence of skin pruritus is associated with the transdermal patch compared to the topical gels; however, the use of hydrocortisone cream, applied after the transdermal system has been removed, may reduce skin irritations that may develop. Currently, only AndroGel® has an A-rated generic formulation.

According to current consensus guidelines, intramuscular (IM) and topical testosterone preparations are generally recommended for the management of hypogonadism in adult male patients. ¹³⁻¹⁶ The selection of testosterone replacement therapy should be a joint decision between the patient and physician and should be made after consideration of patient preferences, the pharmacokinetic profiles of the respective agents, treatment burden, and cost. The short-acting preparations may be preferred over long-acting depot preparations when initiating treatment in patients with late-onset hypogonadism due to the potential development of an adverse event that may require rapid discontinuation of testosterone replacement therapy. Moreover, the guidelines do not recommend one topical preparation over another.

Medications

Table 1. Medications Included Within Class Review

Generic Name (Trade name)	Medication Class	Generic Availability
Testosterone (Androderm [®] , AndroGel [®] *, Axiron [®] , Fortesta [®] , Striant [®] , Testim [®] , Testopel [®] , Vogelxo [®])	Androgens	а

^{*}A-rated generic exists in at least one formulation or strength

Indications

Table 2. Food and Drug Administration Approved Indications 1-9

-	Testosterone		
Indication	Androderm [®] , AndroGel [®] , Axiron [®] , Fortesta [®] , Striant [®] , Testim [®] , Testopel [®] , Vogelxo [®]		
Hypogonadism, primary	а		
(congenital or acquired in males)	(all)		
Hypogonadotropic hypogonadism in males	а		
(congenital or acquired)	(all)		
Stimulate puberty in carefully selected males with	a ,		
clearly delayed puberty	(Testopel [®])		

In addition to the Food and Drug Administration-approved indications, testosterone has been used off-label for male infertility, osteoporosis and weight gain. Testosterone has also been used concomitantly with estrogens for the management of vasomotor symptoms associated with menopause and in postmenopausal women with decreased sexual desire.¹⁰

Because of their anabolic and androgenic effects on performance and physique, androgens have been misused and abused by athletes, bodybuilders, and others.¹⁷ Due to the potential risk of serious adverse health effects, androgens should not be used to enhance athletic performance. Testosterone replacement therapy is also not indicated for the treatment of erectile dysfunction in men with normal serum testosterone concentrations.

Pharmacokinetics

Table 3. Pharmacokinetics 1-10

Drug	Bioavailability (%)	Absorption (%)	Renal Excretion (%)	Active Metabolites	Serum Half- Life (hours)
Testosterone, transdermal (buccal system, gels, implant, patch, solution) [†]	10 (gel)	2 to 8 (gel); 8 (patch);	Urine (90) [‡]	Estradiol, Dihydro- testosterone	0.2 to 1.7*

^{*} Half-life not reported for all products but range of 10 to 100 minutes referenced.





[†] Any product not listed did not have a value reported.

DHT=dihydrotestosterone.

[‡] Based on intramuscular administration.

Clinical Trials

Topical and miscellaneous testosterone products have been evaluated in several clinical trials and are summarized in Table 4. 18-30

The clinical trials evaluating the safety and effectiveness that were used to obtain FDA approval of testosterone pellets are not available. However, a literature search identified a phase IV clinical trial by Kaminetsky et al. Results from the open-label trial showed that mean testosterone levels significantly increased from pre-implantation values at week one, week four and week 12 visits (P<0.001 at all time points) and had returned to pre-implantation levels by week 24 (P=0.58). In addition, luteinizing hormone (LH) levels significantly decreased from pre-implantation values at week one, week four and week 12 visits (P<0.001 at all time points) and returned to pre-implantation levels by week 24 (P=0.87). Prostate-specific antigen levels remained unchanged for the duration of the study. Improvements in symptoms were determined with multiple questionnaires including International Index of Erectile Function (IIEF)-erectile function domain and International Prostate Symptom Score (IPSS). Mean IIEF scores were not significantly different at the end of the study when compared with baseline (P=0.56). Although the severity of voiding symptoms, as assessed by IPSS, decreased at all time points compared with pre-implantation scores, there was not a statistically significant difference (P=0.76, P=0.92, P=0.68, respectively). Overall, implanted testosterone pellets were found to be well tolerated.¹⁸

Several clinical studies have shown that the transdermal patch and gels all restore serum testosterone concentrations to within normal limits and maintain sexual characteristics, sexual behavior, mood, and muscle development, and improve bone mineral density in hypogonadal men. The results of these head-to-head trials favored the use of the gel over the patch. 19-22

In a randomized, multidose, multicenter, active-controlled study comparing two doses of testosterone gel (Testim® 50 mg and 100 mg) and a transdermal testosterone system, Testim® 100 mg produced significantly higher serum levels of testosterone, free testosterone and dihydrotestosterone (DHT). All three treatments produced significant increases in lean body mass (LBM) while only Testim® 100 mg produced significant decreases in percentage of fat. Significant differences between treatment groups were seen in the alleviation of negative mood and improvements in spontaneous erections favoring Testim® over transdermal testosterone for both measures. All three treatment groups produced significant improvements in sexual motivation, sexual desire and sexual performance. The transdermal testosterone system was associated with a higher incidence of treatment-emergent adverse events. In a second study comparing two doses of Testim®, a transdermal testosterone patch (Androderm®) and placebo, all treatment groups produced similar increases in serum testosterone and DHT levels. All treatment groups produced increases in LBM, however the Testim® 100 mg group increased LBM to a significantly greater degree compared to the Androderm® and placebo groups (*P*<0.05 for each measure). The use of both Testim® and Androderm® resulted in significant decreases in fat mass compared to placebo. Only Testim® 100 mg produced significant improvements in sexual function over placebo. There were no significant differences among treatment groups in improving mood, and Androderm® was associated with more treatment-emergent adverse events.

When two doses of a testosterone gel (AndroGel[®]) were compared to Androderm[®], AndroGel[®] 100 mg was associated with significantly higher levels of testosterone and free testosterone compared to AndroGel[®] 50 mg and Androderm[®]. There were significant increases in serum DHT levels with both doses of AndroGel[®] compared to Androderm[®]. The discontinuation rate, mostly due to adverse skin reactions, was significantly greater in the Androderm[®] group. In a study by Wang et al, AndroGel[®] and Androderm[®] average serum testosterone levels increased greatest with AndroGel[®] 100 mg (*P* values not reported).²² A decrease in percent body fat and total fat mass occurred in all treatment groups, however, this was only significant for AndroGel[®]. All treatment groups produced significant improvements in sexual function. Treatment with AndroGel[®] resulted in significant increases in prostate specific antigen levels. Skin irritation at the application site occurred in 65.8, 5.3 and 5.7% of patients in the Androderm[®], AndroGel[®] 100 mg and 50 mg groups. This study also demonstrated that all treatments caused a significant increase in hemoglobin (Hgb) and hematocrit (Hct) but had no overall effects on lipid profiles or blood chemistries.





In an extension study, patients treated with three doses of AndroGel[®] were observed for a period of 36 months.²³ Long-term treatment with AndroGel[®] maintained increased levels of serum testosterone and improvements in sexual function, positive mood and body composition. A gradual, but significant improvement in hip and spine bone mineral density was also observed. Increases in Hgb and Hct plateaued at 12 months and clinically insignificant increases in high-density lipoprotein cholesterol, serum creatinine and total bilirubin were seen. Serum levels of prostate specific antigen showed no further significant increases past six months of treatment. Treatment-emergent adverse events included application site reactions (7.4%), acne (7.4%) and gynecomastia developed in eight patients.

Grober et al evaluated the efficacy of changing from one testosterone gel preparation to another after suboptimal response. ²⁴ Of the 370 hypogonadal men on testosterone replacement therapy, 20% of men underwent a brand substitution due to initial suboptimal response. Among men switching from AndroGel® to Testim® a total of 69, 58 and 65% experienced improvements in libido, erectile function and energy levels, respectively. The rates of improvement for these same parameters among men switching from Testim® to AndroGel® were 46, 39 and 46%, respectively. Changing from AndroGel® to Testim® was reported to have resulted in improved clinical and biochemical responsiveness. Changing from Testim® to AndroGel® eliminated or minimized unwanted side effects (primarily scent).

The safety and efficacy of Striant[®] (testosterone buccal tablet) was evaluated in a 12 week, open-label, multicenter, phase III clinical trial involving 98 hypogonadal men. At the conclusion of the trial, 86.6% of patients with sufficient data for full analysis had mean serum testosterone concentration values within the physiologic range. The mean (± standard deviation) serum testosterone concentration at the end of the study was 520 (±205) ng/dL compared with a mean of 149 (±99) ng/dL at baseline.⁸

In a multicenter, randomized control trial by Korbonits et al, testosterone buccal 30 mg applied twice daily was compared to the testosterone transdermal patch (Andropatch® [not commercially available in the U.S.] or Androderm®) 5 mg once-daily for seven days. The investigators concluded (results not reported) testosterone buccal was non-inferior to the testosterone patch formulation. At all measured time points, the mean testosterone levels were within the established physiological range among patients receiving the buccal formulation compared to five measured time points falling outside of this range among patients receiving the patch formulation. Also, the proportion of patients with levels outside the physiological range was lower in the buccal group compared to the patch group for both the mean (0 to 24 hour) and minimum testosterone levels (the differences; *P*<0.001 for each). The serum testosterone concentrations over the 24-hour period were higher for patients receiving buccal testosterone compared to those receiving the patch (*P*<0.00001). The mean maximum and mean minimum 24-hour testosterone levels were within the physiological range for the testosterone buccal group; whereas only the mean maximum 24-hour testosterone level was within the physiological range for the testosterone patch group. A total of 84.8% of patients in the buccal group were within the physiological range over 24 hours compared to 55.1% of patients in the patch group. The most common adverse events reported among both groups were application site reactions.

In an open-label efficacy trial (N=155), Wang et al evaluated varying doses of testosterone 2% topical solution (Axiron®) applied to the axilla once daily. During the open-label phase of the trial, the mean serum testosterone level before and after application of the testosterone solution was within the adult male range over the 24-hour measurement period on days 15, 60 and 120. Among subjects who were responders at study endpoint (day 120), the geometric mean of serum testosterone values for subjects on any dose was 16.86 nmol/L. Additionally, the proportion of patients completing the study with an average testosterone concentration (Cavg) in the normal range was 76.1% on day 15/16, 84.8% on day 60/61, and 84.1% at day 120. Serum DHT levels and serum free testosterone remained relatively stable over the 24 hours following dosing. The DHT/testosterone ratio values among patients completing the study and among responders remained relatively constant from baseline. Improvements in sexual desire and activity were apparent 15 days after application of testosterone solution and were sustained throughout the study. Statistically significant changes from baseline were seen in sexual desire, sexual activity, positive mood and negative mood as assessed by the Psychosexual Daily Questionnaire (PDQ) domain for the seven days prior to visits one, 15, 60 and 120. Mean changes from day 1 to 120 in the SF-36 Physical Component and SF-36





Mental Component scores were also statistically significant. Treatment-emergent adverse events in the open-label study included application site irritation, application site erythema, headache, increased hematocrit, nasopharyngitis, diarrhea, and vomiting.

Dobs et al evaluated the efficacy of testosterone topical gel (Fortesta $^{\$}$) 40 mg applied to the thighs once daily in varying doses depending upon serum testosterone response in a multicenter, open-label, non-comparative trial. At study endpoint (day 90), the mean serum total testosterone concentration over 24 hours (C_{avg} 0 to 24hr \pm SD) for the 129 individuals with data available for analysis, was 438.56 \pm 162.51 ng/dL, a total of 77.5% of patients achieving a mean serum testosterone level within the pre-defined normal physiological range of \geq 300 and \leq 1140 ng/dL (95% CI, 70.3% to 84.7%). By day 35, 76.2% (95% CI, 68.8% to 83.6%) of patients had reached the primary endpoint and on day 90, 22.5% of patients had a total testosterone level <300 ng/dL. The most commonly reported adverse events were skin reactions, upper respiratory infections, and sinusitis. Skin reactions considered possibly/probably related to study medication were reported in 16.1% of patients, of which 79.2% were determined to be mild in severity.

A meta-analysis of 16 studies evaluating testosterone supplementation for the diagnosis or erectile dysfunction was conducted by Jain et al²⁸. The overall response rate was $57\% \pm 2.3\%$ (203 of 356 cases). The etiology of impotence was reported in 11 of the articles; of which nine included stratified response rates based upon primary versus secondary etiology. Among the studies with stratified results, 75 of 117 (64% ± 4%) men with a primary etiology responded and 53 of 120 (44% ± 2.9%) men with a secondary etiology responded, which was determined to be statistically significant (P<0.001). Further analysis evaluated the delivery method [transdermal patch, intramuscular injection, and oral routes of administration] and found that intramuscular and oral formulation were similar with a response rate of 51.2% ± 2.9% versus 53.2% ± 5.6, respectively (independent sample z test for proportions weighted by study sample size; P=0.86). Conversely, the transdermal formulation was significantly different than intramuscular formulation with a response rate of 80.9% ± 5.9% (independent sample z test for proportions weighted by study sample size; P<0.001). The response rate for transdermal delivery was also significantly different from oral delivery (independent sample z test for proportions weighted by study sample size; P<0.001). Only five of the 16 trials evaluated reported response rates for both placebo and testosterone and had randomized crossover evaluations. There was a mean response of 16.7% versus 65.4% for the placebo and testosterone arms, respectively (two-sample z test for proportions weighted by study sample size z=5.9; P<0.0001). The observed difference was 48.7% (range 16.7% to 65.4%, 95% CI, 32.6 to 64.8) in favor of testosterone.

In an open label extension study Kaufman et al evaluated efficacy of testosterone 1.62% gel up to one year of therapy. ²⁹ Results from the study show that testosterone 1.62% is effective in replacement therapy with 78% (95% CI, 70.0% to 84.6%) and 87.0% (95% CI, 66.4% to 97.2%) of the different dosing regimens reaching therapeutic levels of testosterone. This study also showed that >50% men require doses larger than the traditional starting dose, which is in agreement with previous data.

Blick et al recently evaluated the use of testosterone replacement therapy in human immunodeficiency virus infection/acquired immune deficiency syndrome (HIV/AIDS) patients utilizing the Testim Registry in the United States (TRiUS)³⁰ In this prospective cohort study the effects of replacement therapy with testosterone 1% (Testim[®]) were evaluated in HIV/AIDS patients. During the twelve month study, both non-HIV/AIDS patients and HIV/AIDS cohorts had significant increases in total testosterone and free testosterone to within normal limits along with increased sexual function and improved and decreased antidepressant use. Body composition profiles improved significantly in men without HIV/AIDS (P≤0.05) and remained stable in men with HIV/AIDS during the twelve months of follow-up.





Table 4. Clinical Trials

Study and Drug Regimen	Study Design	Sample Size	End Points	Results
Study and Drug Regimen	and	and Study	Liid Foliits	Results
	Demographics	Duration		
Treatment of Hypogonadis		Duration		
Kaminetsky et al ¹⁸	(UUA215)	(UUA215)	Primary:	Primary:
(UUA215)	(00A213) OL	N=30	Mean testosterone,	(UUA215)
Testosterone pellets	OL	14-50	LH, IIEF score, IPSS	The preimplantation mean testosterone level was 216 ng/dL. Mean
implanted dose based on	Men ≥18 years of	24 weeks	score and adverse	testosterone levels were significantly higher at the week one, week four,
baseline testosterone level	age with primary	Z+ WCCR3	events	and week 12 visits (845 ng/dL, 838 ng/dL, 524 ng/dL, respectively)
and BMI	or secondary	(UUA216)	CVCIIIO	compared with the preimplantation level (P<0.0001 at all time points).
and Dim	hypogonadism,	N=24	Secondary:	Mean testosterone at the conclusion of the study (week 24, or earlier for
(UUA216)	historical serum		Not reported	subjects who opted for a second implant when testosterone levels were
Testosterone pellets	testosterone	24 weeks		<315 ng/dL) had returned to preimplantation levels (232 ng/dL, P=0.58).
implanted dose based on	concentration of			
peak testosterone level	≤315 ng/dL and ≥			Mean LH was reduced from a preimplantation level of 5.1 ng/dL to 1.3
during UUA215	three months of			ng/dL, 0.2 ng/dL, and 0.6 ng/dL at week one, week four, and week 12,
	testosterone			respectively (P<0.0001 at all time points). By the end of the study, mean
	replacement			LH had returned to pre-implantation level (5.2 ng/dL, P=0.87).
	therapy			
				Mean IIEF scores were not significantly higher compared with baseline
	(UUA216)			(15.9) at the end of the study (18.5, P=0.56). However, there was a
	ES, OL			significant difference in IIEF scores compared with baseline at week four
				(20.1, P=0.003) and week 12 (20.9, P=0.001).
	Patients who			TI 11 (11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	enrolled in			The severity of voiding symptoms, as assessed by IPSS, decreased at
	UUA215 and had			all time points compared with pre-implantation scores, but did not reach
	a total			statistical significance (P =0.76, P =0.92, P =0.68 at weeks 4, 16 and 24,
	testosterone level			respectively).
	≤315 ng/dL at the end of the study			(UUA216)
	end of the study			Mean testosterone levels increased from 201 ng/dL at the time of
				implant to 743 ng/dL at week four (P <0.0001), and all subjects had
				increased testosterone levels at this time point compared with baseline.
				Although mean testosterone levels had fallen below 315 ng/dL in the 22
				subjects for whom week 16 data are available, they were still
				significantly higher at this time point compared with the time of implant
				(200 ng/dL vs 275 ng/dL, P=0.003). Mean testosterone levels at the end
				of the study were similar to those at the time of implant (200 ng/dL vs





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
McNicholas et al ¹⁹ Testosterone gel (Testim [®]) 50 mg daily in the morning vs testosterone gel (Testim [®]) 100 mg daily in the morning vs testosterone patch (Andropatch [®] *) 2.5 mg two patches daily in the morning	AC, DB, MC, OL, RCT Hypogonadal men, 31 to 80 years old, morning serum testosterone level ≤10.4 nmol/L at screening with one or more symptoms of low testosterone	N=208 90 days	Primary: 24-hour PK profiles at 30, 60 and 90 days; treatment effectiveness as measured by body composition, mood, and sexual function data at 30, 60 and 90 days; safety Secondary: Not reported	214 ng/dL, P=0.53). All subjects had testosterone levels >315 ng/dL at week four, and nearly a third (31.8%) were still above 315 ng/dL at week 16. (UUA215 and UUA216) Testosterone pellets were generally well tolerated. Most investigator-reported adverse events were mild and transient, and included pain, tenderness, erythema/redness, swelling, and ecchymosis. In both the UUA215 and UUA216 protocols, these symptoms were most commonly observed on the day of implantation and at week one visit. Secondary: Not reported Primary: At 90 days, mean increases in serum testosterone levels were significant for testosterone gel 100 mg (12.41 nmol/L) over testosterone gel 50 mg (6.54 nmol/L; P<0.05) and testosterone patch (3.82 nmol/L; P<0.001). Results at 30 and 60 days were consistent with those at 90 days. The same results were also seen with the mean increase from baseline in free testosterone levels. At 90 days, the mean change in DHT levels with testosterone gel 100 mg were significant over testosterone gel 50 mg (P<0.05) and testosterone patch (P<0.001). In addition, the mean change in DHT levels with testosterone patch at 90 days (P<0.001). Results at 30 and 60 days were consistent with those at 90 days. Significant within-treatment group changes in LBM were seen for all three treatment groups; 0.9 kg (P<0.05), 1.5 kg (P<0.001) and 1.0 kg (P<0.05) for testosterone gel 50 mg, testosterone gel 100 mg, and testosterone patch, respectively. Significant within-treatment group mean changes in percentage fat were only seen with testosterone gel 100 mg (−0.7; P<0.05). There were no statistically significant changes in BMD within any of the three treatment groups.





Study and Drug Regimen	Study Design and	Sample Size and Study	End Points	Results
	Demographics	Duration		
				No significant differences in improvement in positive mood were seen among the three treatment groups. There were significant differences between treatment groups at 90 days in the alleviation of negative mood favoring testosterone gel over the testosterone patch (<i>P</i> <0.05). At 90 days there were significant within-treatment group improvements from baseline in all three groups in sexual motivation, sexual desire, and sexual performance (<i>P</i> <0.05). Both testosterone gel groups had a statistically significant within-treatment improvement in spontaneous erections at all times from baseline (<i>P</i> <0.05). Testosterone patch produced no significant improvement in spontaneous erections at any time. The incidence of treatment-emergent adverse events was 35% for testosterone gel 50 mg, 29% for testosterone gel 100 mg, and 63% for testosterone patch groups. The most commonly reported adverse events were erythema, irritation, and reactions at the application site.
				Secondary: Not reported
Steidle et al ²⁰ Testosterone gel (Testim [®]) 50 mg daily in the morning	AC, DB, MC, OL, PC, RCT Hypogonadal men, 20 to 80	N=406 90 days	Primary: Periodic 24-hour PK profiles; effect of normalizing serum testosterone on body	Primary: At 30 days, all treatment groups had increased mean serum testosterone and DHT concentrations. Testosterone gel 100 mg had a significant increase in mean changes in testosterone concentrations over the testosterone patch (<i>P</i> <0.001). Testosterone gel 50 mg and 100
vs testosterone gel (Testim®) 100 mg daily in the morning	years old, morning serum testosterone level ≤10.4 nmol/L at		composition, sexual function, mood and BMD; safety Secondary:	mg resulted in significant increases in mean changes in DHT concentrations compared to the testosterone patch (<i>P</i> <0.001 for each comparison). By 90 days, similar results were seen across treatment groups.
vs testosterone patch (Androderm®) 2.5 mg 2 patches daily in the	screening with one or more symptoms of low testosterone		Not reported	At 90 days, mean change in LBM was 1.5±4.5, 1.7±2.6, 0.9±1.8 and 0.6±1.8 kg for testosterone gel 50 mg, testosterone gel 100 mg, testosterone patch, and placebo, respectively. Increases in LBM were significantly higher for testosterone gel 100 mg than the testosterone patch and placebo (<i>P</i> <0.05 for each comparison). With the exception of placebo





Study and Drug Regimen	Study Design and	Sample Size and Study	End Points	Results
	Demographics	Duration		
morning	•			treatment, all treatments resulted in a significant decrease in FM
				compared to placebo (<i>P</i> <0.01).
VS				At 90 days, when compared to placebo, testosterone gel 100 mg had
placebo				significant improvements in spontaneous erections (<i>P</i> <0.001), sexual motivation (<i>P</i> <0.05), sexual desire (<i>P</i> <0.01), and sexual performance (<i>P</i> <0.05). No other treatment groups had significant improvements compared to placebo.
				All treatments resulted in mean improvements from baseline in both positive and negative mood scores with no significant differences among the treatment groups.
				The incidence of treatment-related adverse events was 29.1, 36.9, 62.7, and 40.4% for testosterone gel 50 mg, testosterone gel 100 mg, testosterone patch, and placebo, respectively.
				At 90 days, clinically notable decreases in total-C, LDL-C, and HDL-C were seen with testosterone gel 100 mg (<i>P</i> value not reported). Increases in Hgb and Hct were the highest with testosterone gel compared to The testosterone patch and placebo. Increases in PSA values were highest in the testosterone patch group (6.6%).
				Secondary: Not reported
Swerdloff et al ²¹	DB, MC, OL, PG, RCT	N=227	Primary: Serum testosterone	Primary: At 30 and 90 days, testosterone gel 100 mg produced significantly
Testosterone gel	. 5, 101	180 days	and free testosterone	higher C _{avg} testosterone levels over testosterone 50 mg and
(AndroGel®) 50 mg daily	Hypogonadal men, 19 to 68	, ,	levels at 0, 1, 30, 90, and 180 days; safety;	testosterone patch (27.46±1.12 nmol/L vs 19.17±1.06 and 14.46±0.68 nmol/L, respectively; <i>P</i> =0.0001). At 180 days, serum testosterone levels
vs	years old,		serum DHT, E ₂ , FSH,	and PK parameters were similar to those on days 30 and 90 in those
	morning serum		LH, SHBG levels on	patients who continued their initial randomized treatment. Patients
testosterone gel	testosterone		0, 30, 60, 90, 120,	switched to testosterone gel 75 mg had a C _{avg} testosterone level of
(AndroGel [®]) 100 mg daily	level ≤10.4		150 and 180 days	20.84±1.76 nmol/L at 180 days. This value was between the 180 day
vs	nmol/L at screening		Secondary:	C _{avg} testosterone levels achieved with testosterone gel 50 mg (19.24±1.18) and testosterone gel 100 mg (24.72±1.05).





Study and Drug Regimen	Study Design	Sample Size	End Points	Results
otady and Drug Regimen	and	and Study	Liid i Oilits	Results
	Demographics	Duration		
			Not reported	
testosterone patch (Androderm®) 2.5 mg 2 patches daily At 60 days, men with serum testosterone levels <10.4 nmol/L who were applying AndroGel® 50 mg and men with serum testosterone levels >34.7 nmol/L who were applying AndroGel® 100 mg were			·	PK parameters of serum free testosterone levels on days one, 30, 90 and 180 mirrored those of serum testosterone levels. The free testosterone levels in the testosterone gel 100 mg group was 1.4- and 1.7-fold higher than the testosterone gel 50 mg and testosterone patch groups (<i>P</i> =0.001). The discontinuation rate at 90 days for the testosterone patch (27.6%) was significantly higher than testosterone gel 50 and 100 mg (8.2% and 6.4%, respectively; <i>P</i> =0.0002). Most patients discontinued treatment due to adverse skin reactions. Throughout the 180 days, increases in serum DHT levels were
instructed to apply AndroGel® 75 mg once daily for days 91 through 180.				significant with testosterone gel 50 and 100 mg over the testosterone patch (P =0.0001). Mean serum increases to stable levels of E $_2$ occurred in 9.2, 30.9, and 45.5% of patients in the testosterone patch, testosterone gel 50, and testosterone gel 100 mg groups, respectively (P =0.001). All three treatment groups showed a small decrease in serum SHBG
				levels (<i>P</i> =0.0046).
				The mean percent suppression of serum LH levels was the smallest with testosterone patch (30 to 40%), intermediate with testosterone gel 50 mg (55 to 60%), and greatest with testosterone gel 100 mg (80 to 85%; P <0.01). The suppression of serum FSH paralleled that of serum LH levels.
				Secondary: Not reported
Wang et al ²²	DB, MC, OL, PG, RCT	N=227	Primary: Mean change from	Primary: On day 90 the average serum testosterone concentration with
Testosterone gel (AndroGel [®]) 50 mg daily	Hypogonadal men, 19 to 68	180 days	baseline in serum testosterone concentrations, body	testosterone gel 100 mg (27.46±1.12 nmol/L) was 1.4-fold higher than testosterone gel 50 mg (19.17±1.06 nmol/L) and 1.9-fold higher than the testosterone patch (14.46±0.68 nmol/L; <i>P</i> value not reported). On day
VS	years old,		composition, and	180 average serum testosterone concentrations for the treatment groups





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
testosterone gel (AndroGel®) 100 mg daily vs testosterone patch (Androderm®) 2.5 mg two patches daily At 90 days, dose adjustments were made in the AndroGel® groups based on the preapplication serum testosterone levels on day 60. Twenty subjects in the AndroGel® 50 mg group had their dose increased to 75 mg and 20 subjects in the AndroGel® 100 mg group had their dose reduced to 75 mg.	morning serum testosterone level ≤10.4 nmol/L at screening		muscle strength at 90 and 180 days; mean change from baseline in sexual function and mood at 30, 60, 90, 120, 150 and 180 days; degree of skin irritation; mean change from baseline in serum PSA levels at 30 and 90 days; mean change from baseline in Hgb, Hct, lipid profiles and blood chemistries Secondary: Not reported	were 24.72±1.05 nmol/L, 19.24±1.18 nmol/L and 14.14±0.88 nmol/L, respectively. The percent body fat and FM decreased in all treatment groups but was only significant with testosterone gel. At 90 days the total FM was significantly decreased with testosterone gel 50 mg and testosterone gel 100 mg (<i>P</i> =0.0065 and <i>P</i> =0.0001, respectively). At 180 days the total FM decreased further with testosterone gel 100 mg (<i>P</i> =0.008). At 90 days, the percent body fat was significantly decreased with testosterone gel 50 mg and testosterone gel 100 mg (<i>P</i> =0.0018 and <i>P</i> =0.001) and remained significant at 180 days. Significant increases in arm and leg muscle strength were seen in all three treatment groups without intergroup differences on days 90 and 180 (<i>P</i> values compared to baseline ranged between 0.0001 to 0.08). All subjects, regardless of treatment group, showed significant improvement in sexual motivation (<i>P</i> =0.0001), sexual desire (<i>P</i> =0.0001), sexual performance (<i>P</i> =0.0001), self-assessment of satisfaction of erection (<i>P</i> =0.0001) and percentage of full erection (<i>P</i> =0.0001). All three treatment groups showed significant improvement in positive mood scores (<i>P</i> =0.0001) and a decrease in negative mood scores (<i>P</i> =0.0001) without significant between-group differences. Minimal skin irritation at the application site was seen in 5.7 and 5.3% of patients in the testosterone gel 50 mg and 100 mg group. Minimal to severe skin irritation occurred in 65.8% of patients in the testosterone patch group. Mean serum PSA levels significantly increased with testosterone gel 100 mg (<i>P</i> =0.008) and testosterone gel 50 mg (<i>P</i> =0.05) with no significant increase in the testosterone patch group. As a group, both Hgb and Hct increased (<i>P</i> =0.0001). There were no overall treatment effects or intergroup differences in serum concentrations of





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
Wang et al ²³ Testosterone gel (AndroGel [®]) 50 mg daily vs testosterone gel (AndroGel [®]) 75 mg daily vs testosterone gel (AndroGel [®]) 100 mg daily		•	Primary: Mean changes from baseline in serum testosterone, free testosterone, DHT, E2, SHBG, LH and FSH; mean changes from baseline in sexual function and mood, body composition, bone turnover markers, muscle strength and BMD; mean changes from baseline in Hgb, Hct, lipid profiles and blood chemistries; mean changes from baseline in serum PSA and prostate disease; safety Secondary: Not reported	total-C, HDL-C, LDL-C or TG (data not provided). Secondary: Not reported Primary: Mean serum testosterone levels were significantly different (<i>P</i> =0.012) between dosing groups at baseline (six months of TRT). At 12 months, differences among the dosing groups became smaller but remained significant (<i>P</i> =0.042). Serum free testosterone levels followed the same pattern as testosterone. Mean serum DHT levels were different in the three dosing groups at 12 (<i>P</i> =0.0031) and 24 (<i>P</i> =0.018) months with the highest levels seen with testosterone gel 100 mg. Mean serum E ₂ levels progressively increased from 6 to 24 months (<i>P</i> =0.0001) with significant differences between treatment groups. The highest levels of serum E ₂ were seen with testosterone gel 100 mg. No significant change in SHBG was seen. Suppression of LH and FSH was maintained throughout with no significant changes after six months. The suppression was more pronounced with testosterone gel 100 mg. Significant improvements in sexual desire, enjoyment with or without a partner, percent full erection, and self-assessment of satisfaction with erections were maintained as a group throughout the study period. Positive mood scores were improved with treatment and were sustained (<i>P</i> =0.0022). Negative mood parameters were decreased and remained significantly lower (<i>P</i> =0.0013) than baseline without further changes after six months.
				(<i>P</i> =0.0157) and did not significantly change with continued therapy. LBM increased significantly (<i>P</i> =0.0001) from baseline and remained increased throughout the study. A significant decrease in FM was seen at 30 months (<i>P</i> =0.088) without significant differences between doses.





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
				Serum PTH levels significantly increased from baseline (P =0.0001) and continued to increase from six (P =0.0002) until 12 months when it remained stable throughout the rest of the treatment period. Serum SALP levels followed the same pattern (P =0.001). At 12 months serum osteocalcin was significantly elevated and remained elevated throughout treatment (P =0.0001). Serum procollagen levels transiently increased then steadily increased from six months to reach significant levels by 36 months (P =0.0001).
				Muscle strength increased but did not reach significance over time due to the large variation in patients.
				BMD of the hip (<i>P</i> =0.0004) and spine (<i>P</i> =0.0001) showed a gradual and progressive increase with treatment. No significant differences among treatment doses or older and younger patients were observed.
				Serum Hgb and Hct concentrations increased, compared with month zero (<i>P</i> =0.0001) and month six (<i>P</i> =0.001) and plateaued at 12 months.
				Small statistically significant increases in serum HDL-C levels (<i>P</i> <0.001), creatinine (<i>P</i> <0.001), and total bilirubin (<i>P</i> =0.001) were seen but were not clinically significant. No significant changes in total-C, LDL-C, serum liver enzymes, or other clinical chemistry parameters were observed.
				The mean serum PSA was 1.11+0.08 at six months and showed no further significant increases with continued treatment.
				Application-site reactions occurred in 12 of the 163 (7.4%) patients. Acne occurred in 12 (7.4%) of patients and gynecomastia was observed in eight more patients.
				Secondary: Not reported
Grober et al ²⁴	OL	N=370	Primary: Reasons for brand	Primary: Of the 370 hypogonadal men using testosterone gel, 20% underwent a
AndroGel [®] 5 to 10 g	Hypogonadal	Treatment	substitution, total and	brand substitution. The reasons for switching from AndroGel® to Testim®





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
vs Testim [®] 5 to 10 g	men on testosterone gel who underwent a brand substitution due to initial suboptimal biochemical or symptomatic response, mean age of men switched to Testim® was 60 years, mean age of men switched to AndroGel® was 52 years	duration after switch, 4 weeks	free testosterone, presence of hypogonadal symptoms Secondary: Not reported	(N=62) were poor efficacy (92%), hypertension (2%), skin reaction (2%), worsening symptoms (2%), and insurance coverage (2%). The reasons for switching from Testim® to AndroGel® (N=13) were scent (46%), poor efficacy (30%), fear of transfer to partner (8%), flushing (8%) and skin reaction (8%). Prior to substitution, patients initially treated with AndroGel®, had mean total and free testosterone levels of 311 ng/dL and 10.4 pg/mL, respectively. Total testosterone levels were <300 ng/dL in 58% of these patients. Following a change to Testim®, mean total and free testosterone levels increased to 484 ng/dL (<i>P</i> <0.001) and 14.6 pg/mL (<i>P</i> =0.01), respectively. Total testosterone levels remained <300 ng/dL in 17% of these patients. Among patients initially treated with Testim®, the mean total and free testosterone levels were 544 ng/dL and 18.0 pg/dL, respectively. Total testosterone levels were <300 ng/dL in 15% of men. Following a change to AndroGel®, mean total and free testosterone levels were 522 ng/dL (<i>P</i> =0.7) and 16.1 pg/mL (<i>P</i> =0.6), respectively. Total testosterone levels remained <300 ng/dL in 27% of these patients.
Korbonits et al ²⁵	IT, MC, RCT	N=66	Primary:	Secondary: Not reported Primary:
Testosterone buccal 30 mg BID (Striant®) vs Andropatch®* or Androderm® TD patch 5 mg once daily	Men with testosterone deficiency with a morning serum testosterone < 6.94 nmol/L, normal agerelated PSA levels, and Hct < 50	7 days	Non-inferiority analysis (endpoints not defined) Secondary: Efficacy analysis of superiority (endpoints not defined)	Investigators concluded that non-inferiority was established (results not reported). Secondary: In the buccal testosterone group, the mean testosterone concentrations at all measured time points (days three, four, six, seven and eight) were within the physiological range; whereas mean concentrations at five time points were outside of the physiological range among patients in the testosterone patch group. For both mean (0 to24 hour) and minimum testosterone levels, the proportion of patients with levels outside the physiological range was





Study and Drug Regimen	Study Design	Sample Size	End Points	Results
	and Demographics	and Study Duration		
	Demographics	Duration		lower in the buccal group than in the patch group (the differences; <i>P</i> <0.001 for each).
				The serum testosterone concentrations over the 24-hour period were higher for patients receiving buccal testosterone compared to those receiving the patch (mean AUC \pm SD; 451.31 \pm 140.71 h*nmol/L vs. 304.63 \pm 134.46 h*nmol/L; 95% CI, 1.25 to 1.91; P <0.00001).
				The mean maximum and mean minimum 24-hour testosterone levels were within the physiological range for the testosterone buccal group. Comparatively, the mean maximum 24-hour testosterone level was within the physiological range for the testosterone patch group; however, the mean minimum 24-hour testosterone level was below the physiological range. A total of 84.8% of patients in the buccal group were within the physiological range over 24 hours compared to 55.1% of patients in the patch group.
				Testosterone concentrations were within the physiological range in the buccal group for a significantly greater portion of the 24-hour treatment period compared to the patch group (84.9 vs 54.9%; <i>P</i> <0.001).
				Mean DHT levels were within the normal range (1.03 to 2.92 nmol/L) for both the buccal group (2.36 \pm 0.99 nmol/liter) and the patch group (1.2 \pm 0.57 nmol/L).
				The median estradiol concentrations increased from baseline to day seven, but returned to baseline levels at the follow-up visit. The median increase from baseline to day seven was greater in the buccal group (55.07 pmol/liter) compared to the patch group (34.87 pmol/liter; $P < 0.001$).
26				A total of 51.5% of patients in the buccal group reported an adverse event compared to 47.1% in the patch group. The most commonly reported adverse events among both groups were application site disorders.
Wang et al ²⁶	OL with	N=155 OL	Primary:	Primary:





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
Testosterone 60mg topical solution applied to each axilla once daily (Axiron®)	extension study Men ≥18 years with androgen deficiency (diagnosis of hypogonadism) and a BMI <35.0 kg/m² with testosterone levels on two consecutive samples < 10.4 nmol/L and a baseline Hgb level ≥ 110.5 g/L.	study 120 days N=71 extension study 60 days	Total testosterone and DHT (OL phase) Secondary: PDQ domain assessing sexual desire, enjoyment and performance, sexual activity, and mood, SF-36 health survey (extension phase)	At day 120, the proportion of patients completing the study with an average testosterone concentration (C _{avg}) in the normal range was 84.1%. Also, 76.1% and 84.8% of patients completed the study with a C _{avg} in the responder range on days 15/16 and 60/61, respectively. The mean serum testosterone level before and after dosing was within the adult male range over the 24-hour period on days 15, 60 and 120. The geometric mean of serum testosterone over 24 hours was 15.62 nmol/L (coefficient of variation [CV]; 38%). Among subjects who were responders at day 120, the geometric mean of serum testosterone values for subjects on any dose was 16.86 nmol/L. Serum DHT levels and serum free testosterone remained relatively stable over the 24-hours following dosing. The mean day 15 baseline pre-dose DHT/T ratio was 0.23, and the mean DHT/T ratio remained between 0.17 to 0.26 throughout the 24-hour period. The ratio values among patients completing the study and among responders remained relatively constant from baseline. Secondary: Improvements in sexual desire and activity were apparent 15 days after application of testosterone and were sustained throughout the study. Statistically significant changes from baseline were seen in sexual desire, sexual activity, positive mood and negative mood as assessed by the PDQ domain for the seven days prior to visits one, 15, 60 and 120. Significant mean changes from day one to 120 for SF-36 Physical Component and SF-36 Mental Component scores were 1.55 (SD=7.72; P=0.0254) and 4.54 (SD=9.20; P<0.0001), respectively. Treatment-emergent adverse events occurring in >2% of patients receiving at least one dose of testosterone in the open-label study included: application site irritation, application site erythema, headache, increased hematocrit, nasopharyngitis, diarrhea, and vomiting. Three patients withdrew from the open-label phase of the study due to adverse events, including superficial thrombophlebitis, effects on lability/anger, and malignant melanoma; while two patients with





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
				extension phase of the study due to application site irritation and application site erythema.
Dobs et al ²⁷ Testosterone gel 40 mg applied to the thighs once daily (Fortesta [®]) Dose adjustments allowed for a downward titration to a minimum of 10 mg daily and an upward titration to 70 mg daily.	MC, NC, OL Men 18 to 75 years, with primary or secondary hypo- gonadism (defined as a single serum testosterone concentration <250 ng/dL or two consecutive serum testosterone levels <300 ng/dL at least one week apart) and a BMI ≥22 kg/m² and <35 kg/m²	N=149 90 days	Primary: The average serum total testosterone concentration over 24 hours (C _{avg} 0 to 24h) on Day 90 Secondary: The maximum serum testosterone concentration (C _{max}) on Day 90	Primary: Of the 129 patients with available data for analysis, the mean C _{avg} over 24 hours was 438.56 ± 162.51 ng/dL with 77.5% of patients achieving a mean serum testosterone level within the pre-defined normal physiological range (≥300 and ≤1140 ng/dL) (95% CI, 70.3 to 84.7%). By day 35, 76.2% (95% CI, 68.8 to 83.6%) of patients had reached the primary endpoint. On day 90, 22.5% of patients had a total testosterone level <300 ng/dL. Secondary: The C _{max} ± SD was 827.6 ± 356.5 ng/dL on day 90. At endpoint, a total of 94.6% of patients achieved a C _{max} ≤1500 ng/dL, 1.6% of patients had levels between 1880 and 2500 ng/dL, and no patients had levels >2500 ng/dL. This C _{max} was evident by treatment day 35. Adverse events were reported in 46.3% of patients; however on 22.8% were considered related to the study medication. The most commonly reported adverse events were skin reactions, upper respiratory infections and sinusitis. Skin reactions were considered 'possibly' or 'probably' related to study medication in 16.1% of patients, of which 79.2% were mild in severity.
Kaufman et al ²⁸ Testosterone 1.62% titrated to therapeutic dose vs testosterone 1.62% titrated to a specific serum testosterone level and then continued at dose for the remainder of the study	OL,ES Males 18 to 80 years of age with hypogonadism who completed a six month double blind study that elected to continue	N=191 182 days	Primary: Percentage of subjects achieving an average serum total testosterone concentration in the normal range of 300 to 1,000 ng/dL Secondary: Measurement of SHBG, LH, FSH, and selected serum	Primary: At the end of the study (day 364) 77.9% (95% CI, 70.0% to 84.6%) of subjects continuing on active testosterone treatment had Cav values within the normal range with 87.0% (95% CI, 66.4% to 97.2%) of the Formerly Placebo group reaching Cav values within in the normal range. A combined 79.2% (95% CI, 72.1% to 85.3%) of patients in both groups reached a Cav value within the normal range. Secondary: SHBG levels increased significantly from baseline on day 266 (P<0.0001) and on day 364 (P<0.0166) for the Continuing Active group but not for the Formerly Placebo group.





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
			inflammatory and cardiovascular risk markers, waist-to-hip ratio, and serum markers of bone	LH levels decreased significantly from baseline on day 266 and day 364 with 1.62% testosterone treatment for the Continuing Active group (P<0.0001 for both days) and for the Formerly Placebo group (P<0.0054 and P<0.0309, respectively).
			metabolism; quality of life	FSH levels decreased significantly from baseline on day 266 and day 364 for the Continuing Active group (P<0.0001 for both days) and Formerly Placebo group (P<0.0001 and P<0.0087, respectively).
				Interleukin-10 decreased significantly from baseline on day 364 in the Continuing Active group (P<0.001) and on day 266 for the Formerly Placebo group (P<0.0089).
				MMP-9 levels decreased significantly from baseline for the Continuing Active group on both day 266 (P<0.0080) and day 364 (P<0.0055) but not for the Formerly Placebo group (P>0.05).
				Alkaline phosphatase values for bone-specific alkaline phosphatase significantly (P<0.0001) increased from baseline on day 266 for both groups, although no significant changes were seen on day 364.
				Values for type 1 cross-linked C-telopeptide decreased significantly from baseline on day 266 and day 364 for the Continuing Active group (P<0.001 both days) but not for the Formerly Placebo group (P > 0.05 both days).
20				Scores on the SF-36 remained stable throughout the treatment period.
Miner et al ²⁹ (abstract)	Cohort , PRO	N=849	Primary: Total testosterone,	Primary: Mean total testosterone and free testosterone levels increased
Testosterone 1%	Men in the Testim Registry in the United States (TRiUS) –	12 months	free testosterone, prostate specific antigen, sexual function,	significantly after three months of therapy. For mean total testosterone level of 16.8 \pm 9.87 nmol/L (P<0.001) and mean free testosterone level 286.3 \pm 224.9 pmol/L (P<0.001).
	hypogonadal men who were prescribed TRT		mood/depression, and cardiometabolic and anthropometric criteria	Mean PSA levels increased significantly (P=0.004) from 1.12 \pm 1.11 μ g/L at baseline to 1.26 \pm 1.22 μ g/L after 12 months of TRT, although changes were within guidelines (< 1.4 μ g/L/year increase).





Study and Drug Regimen	Study Design and Demographics	Sample Size and Study Duration	End Points	Results
			before and after therapy Secondary: Not reported	Significant improvements were seen in sexual function and mood/depression at three months and in metabolic parameters at 12 months.
Blick et al (abstract) ³⁰	Cohort, PRO	N=849	Primary: Total testosterone,	Primary: During the 12 months, both the HIV/AIDS and non-HIV/AIDS cohorts
Testosterone 1% in HIV/AIDS patients	Men in the Testim Registry in the United States	12 months	free testosterone, sexual function, depression and body	experienced significant elevations in total testosterone and free testosterone levels to within normal ranges.
VS	(TRiUS) – hypogonadal men		composition profiles	Sexual function and depression scores improved and antidepressant medication use decreased in both cohorts.
Testosterone 1% in non- HIV/AIDS patients	who were prescribed TRT broken up by HIV status for this study		Secondary: Not reported	Body composition profiles improved significantly (P≤0.05) in men without HIV/AIDS and remained stable in men with HIV/AIDS during the 12 months of follow-up.
*Agent not available in the United Str				Secondary: Not reported

^{*}Agent not available in the United States.

Study abbreviations: AC=active-controlled, DB=double-blind, ES=extension study, IT=international, MA=meta-analysis, MC=multicenter, NC=non-comparative, OL=open-label, PC=placebo-controlled, PG=parallel-group, PK=pharmacokinetic, PRO=prospective trial, RCT=randomized controlled trial, RETRO=retrospective, SA=single-arm

Miscellaneous abbreviations: AFS=American Fertility Society, BMD=bone mineral density, BMI=body mass index, C=cholesterol, Cavg=average concentration, DHT=dihydrotestosterone,
E2=Estradiol, FM=fat mass, FSH=follicle-stimulating hormone, Hct=hematocrit, HDL=high density lipoprotein, Hgb=hemoglobin, IIEF=International Index of Erectile Function-erectile function domain, IPSS= International Prostate Symptom Score, LBM=lean body mass, LDL=low density lipoprotein, LH=luteinizing hormone, PK=pharmacokinetics, PSA=prostate specific antigen,
PTH=parathyroid hormone, SALP=bone-specific alkaline phosphatase, SHBG=sex hormone-binding globulin, T=testosterone, TG=triglycerides, TRT=testosterone replacement therapy





Special Populations

Table 5. Special Populations¹⁻⁹

Generic		Population	and Precaution		
Name	Elderly/	Renal	Hepatic	Pregnancy	Excreted in
	Children	Dysfunction	Dysfunction .	Category	Breast Milk
Testosterone	No dosage adjustment is	Use with	Use with	X	Contra-
buccal	required in the elderly.	caution, not	caution, not		indicated
mucoadhesive		studied in	studied in		
system	Elderly patients treated	renal	hepatic		
	with androgens may be	dysfunction.	dysfunction.		
	at increased risk for				
	development of prostatic	It appears	Testosterone		
	hypertrophy and	that no	use has been		
	prostatic carcinoma.	dosage	associated		
	Cofety and office as in	adjustment is	with the		
	Safety and efficacy in	required.	development		
	males <18 years have		of severe		
Tootootorono	not been established.	Use with	hepatotoxicity. Use with	X	Contra-
Testosterone gel	No dosage adjustment is required in the elderly.	caution, not	caution, not	^	indicated
gei	required in the elderry.	studied in	studied in		indicated
	Elderly patients treated	renal	hepatic		
	with androgens may be	dysfunction.	dysfunction.		
	at increased risk for	ayorarromorr.	ayorarronorr.		
	development of prostatic	It appears	Testosterone		
	hypertrophy and	that no	use has been		
	prostatic carcinoma.	dosage	associated		
		adjustment is	with the		
	Safety and efficacy in	required.	development		
	males <18 years have		of severe		
	not been established.		hepatotoxicity.		
Testosterone	No dosage adjustment is	Use with	Use with	X	Contra-
implant pellet	required in the elderly.	caution, not	caution, not		indicated
		studied in	studied in		
	Elderly patients treated with androgens may be	renal dysfunction.	hepatic		
	at increased risk for	dysidifiction.	dysfunction.		
	development of prostatic	It appears	Testosterone		
	hypertrophy and	that no	use has been		
	prostatic carcinoma.	dosage	associated		
	Processing Community	adjustment is	with the		
	Indicated for the	required.	development		
	stimulation of puberty in		of severe		
	selected males with		hepatotoxicity.		
	clearly delayed puberty.				
	No age is specified.				
Testosterone	No dosage adjustment is	Use with	Use with	X	Contra-
patch	required in the elderly.	caution, not	caution, not		indicated
	Flands a - the state ()	studied in	studied in		
	Elderly patients treated	renal	hepatic		
	with androgens may be	dysfunction.	dysfunction.		
	at increased risk for	It appears	Testosterone		
	development of prostatic hypertrophy and	It appears that no	use has been		
	prostatic carcinoma.	dosage	associated		
	prostatio carcinoma.	dosage	นออบเสเซน		



Generic	Population and Precaution							
Name	Elderly/	Renal	Hepatic	Pregnancy	Excreted in			
	Children	Dysfunction	Dysfunction	Category	Breast Milk			
		adjustment is	with the					
	Safety and efficacy in	required.	development					
	males <18 years have		of severe					
	not been established.		hepatotoxicity					
Testosterone	No dosage adjustment is	Use with	Use with	X	Contra-			
solution	required in the elderly.	caution, not	caution, not		indicated			
		studied in	studied in					
	Elderly patients treated	renal	hepatic					
	with androgens may be at increased risk for	dysfunction.	dysfunction.					
	development of prostatic	It appears	Testosterone					
	hypertrophy and	that no	use has been					
	prostatic carcinoma.	dosage	associated					
		adjustment is	with the					
	Safety and efficacy in	required.	development					
	males <18 years have		of severe					
	not been established.		hepatotoxicity					



Adverse Drug Events

Table 6. Adverse Drug Events (%)¹⁻⁹

Adverse Event	Androderm®	AndroGel®	Axiron®	Fortesta [®]	Striant [®]	Testim [®]	Vogelxo [®]	Testopel [®]
Central Nervous System	•							
Abnormal dreams	-	-	-	1.3	-	-	-	-
Anxiety	-	-	а	-	-	-	-	а
Asthenia	-	<3	5 -	-	-		-	-
Depression	-	1	_	-	_	-	-	а
Dizziness	-	-	_	а	-	-	-	-
Emotional lability (including anger)	-	2.6 to 3	а	-	-	-	-	-
Headache	<4	<4	5 to 6	-	3.1	1	1	а
Insomnia			-	-	-	1	1	-
Libido, increased or decreased	-	<3	-	-	-	-	-	а
Migraine	-	-	-	а	-	-	-	-
Mood swings	-	-	_	-	-	1	1	-
Nervousness	-	-	а	-	-	-	-	-
Smell disorder	-	-	_	-	-	1	1	-
Dermatologic								
Acne	-	1 to 3	а	-	-	-	-	а
Allergic contact blistering	12	-	-	-	-	-	-	-
Alopecia	-	1	ı	-	ı	-	-	а
Application site burning	3	-	ı	-	ı	-	-	-
Application site erythema	<7	-	5 to 7	а		-	-	-
Application site edema	-	-	а	-	ı	-	-	-
Application site exfoliation	<3	-	ı	-	ı	-	-	-
Application site induration	3	-	-	-	-	-	-	-
Application site reaction	-	3 to 5	ı	-	ı	2 to 4	4	-
Application site inflammation	-	-	ı	-	ı	-	-	а
Application site irritation	-	-	7 to 8	а	ı	-	-	-
Application site pain	-	-	ı	-	ı	-	-	а
Application site warmth	-	-	а	-	-	-	-	-
Application site vesicles	6	-	-	-	-	-	-	-
Contact dermatitis	-	2.1	-	а	-	_	-	-
Folliculitis	-	-	а	_	-	_	-	-
Pruritus	17 to 37	-	-	а	-	-	-	-



Adverse Event	Androderm [®]	AndroGel®	Axiron®	Fortesta [®]	Striant [®]	Testim [®]	Vogelxo [®]	Testopel [®]
Rash	<3	_	-	а	-	-	-	-
Skin reactions	-	-	-	16.1	-	-	-	-
Endocrine and Urogenital								
Benign prostatic hyperplasia	-	-	-	-	-	1	-	-
Blood testosterone, increased	-	-	а	-	-	-	-	-
Blood testosterone, decreased	-	-	-	-	-	а	-	-
Breast pain	-	<3	-	-	-	-	-	-
Breast tenderness	-	_	а	-	-	-	-	-
Erectile dysfunction	-	-	-	а	-	-	-	-
Gynecomastia	-	<3	-	-	-	1	-	а
Hot flushes	-		ı	-	-	1	1	-
Penile erections, excess frequency and duration	-	-	-	а	-	-	-	а
Penile erection, spontaneous	-	_	-	-	-	1	1	-
Polyuria	<3	_	-	-	_	-	-	-
Prostate abnormalities	5	-	-	-	-	-	-	-
Prostate disorder	-	3 to 5	-	-	-	-	-	-
Prostate enlarged	<3	-	-	-	_	-	-	-
Prostate specific antigen, increased	-	11.1	1 to 4	1.3	-	-	-	-
Testes disorder	-	<3	-	-	-	-	-	-
Urinary symptoms	-	<2	-	-	-	-	-	-
Gastrointestinal		•						
Abdominal symptoms	-	-	ı	а	-	-	-	-
Cholestatic jaundice	-	-	-	-	-	-	-	а
Diarrhea	<3	-	3 to 4	-	-	-	-	-
Gastrointestinal bleeding	<3	-	ı	-	-	-	-	-
Gastroesophageal reflux disease	<3	-	ı	-	-	-	-	1
Vomiting	-	-	3 to 4	-	_	-	-	-
Hematologic								
Bleeding	<3	-	-	-	-	-	-	-
Hematocrit/ hemoglobin increased	1	2.1	4 to 7	а	-	2	2	-





Adverse Event	Androderm [®]	AndroGel [®]	Axiron®	Fortesta [®]	Striant [®]	Testim [®]	Vogelxo [®]	Testopel [®]		
Polycythemia	-	-	-	а	-	-	-	-		
Red blood cell count,		_		_	-			-		
elevation	_	_	а	_		_	_			
Metabolic	Metabolic									
Blood glucose, increased	-	-	а	-	-	-	-	-		
Cholesterol, increased	-	<2	-	-	-	-	-	-		
Other										
Back pain	6	-	-	-	-	-	-	-		
Blood pressure increase	-	<4	а	-	-	1	1	-		
Fatigue	<3	-	-	а	-	-	-	-		
Gum edema	-	-	-	-	2.0	-	-	-		
Gum or mouth irritation	-	-	-	-	9.2	-	-	-		
Gum pain	-	-	-	-	3.1	-	-	-		
Gum tenderness	-	-	-	-	3.1	-	-	-		
Influenza like illness/malaise	-	-	-	а	-	-	-	-		
Laboratory test, abnormal	-	3 to 6	-	-	-	-	-	-		
Lacrimation, increased	-	-	а	-	-	1	-	-		
Nasopharyngitis	_	-	а	-	-	_	-	-		
Pain in extremities	_	-	-	а	-	_	-	-		
Pelvic pain	<3	-	-	-	-	_	-	-		
Taste sense, diminished	-	-	-	-	2.0	1	-	-		
Taste bitter	-	-	-	-	4.1	-	-	-		
Vitreous detachment	-	-	-	а	-	-	-	-		

a Frequency of adverse event not reported.
- Incidence ≤1% or not reported.

Contraindications

Table 7. Contraindications 1-9

Contraindications	Testosterone		
Contraindications	Androderm [®] , AndroGel [®] , Axiron [®] , Fortesta [®] , Striant [®] , Testim [®] , Testopel [®] , Vogelxo [®]		
Men with carcinoma of the breast or known or	a (all)		
suspected carcinoma of the prostate	- (- /		
Women who are, or who may become pregnant, or who are breastfeeding.	a (all)		
Hypersensitivity to testosterone or any component of the product	a (all)		





Precautions/Warnings

Table 8. Precautions/Warnings¹⁻⁹

Mouning/Draggistian	Testosterone		
Warning/Precaution	Androderm [®] , AndroGel [®] , Axiron [®] , Fortesta [®] , Striant [®] , Testim [®] , Testopel [®] , Vogelxo [®]		
Worsening of Benign Prostatic Hyperplasia and	a (all)		
Potential Risk of Prostate Cancer	a (a ii)		
Polycythemia	a (all)		
Venous Thromboembolism	a (all)		
Use in Women and Children	a (Androderm®)		
Use in Women	a (AndroGel [®] , Axiron [®] , Fortesta [®] , Striant [®] , Testim [®] , Vogelxo [®])		
Potential for Adverse Effects on	2 (a ll)		
Spermatogenesis	a (all)		
Hepatic Adverse Effects	a (all)		
Edema	a (all)		
Gynecomastia	a (all)		
Sleep Apnea	a (all)		
Lipids	a (all)		
Hypercalcemia	a (all)		
Decreased Thyroxine-Binding Globulin	a (all)		
Delayed puberty; use with caution	a (Testopel [®])		
Dosage adjustment less flexible	a (Testopel [®])		
Magnetic Resonance Imaging (MRI)	a (Androderm®)		
Gum-related adverse reactions and limited long-	a (Striant®)		
term information on oral safety	a (Striant)		
Potential for Secondary Exposure to	a (AndroGel [®] , Axiron [®] , Fortesta [®] , Testim [®] , Vogelxo [®])		
Testosterone	, , , , , , , , , , , , , , , , , , ,		
Flammability	a (AndroGel [®] , Axiron [®] , Fortesta [®] , Testim [®] , Vogelxo [®])		



Black Box Warnings Regarding Testosterone Solution and Gels (AndroGel[®], Testim[®], Axiron[®], Vogelxo[®] & Fortesta[®])²⁻⁷

WARNING

Secondary Exposure to Testosterone

Virilization has been reported in children who were secondarily exposed to topical testosterone products.

Children should avoid contact with any unwashed or unclothed application sites in men using testosterone gel/solution.

Healthcare providers should advise patients to strictly adhere to recommended instructions for use.

Drug Interactions

Table 7. Drug Interactions 1-9

Drug	Interacting Medication	Potential Result
Testosterone	Anticoagulants	The concurrent administration of androgens with oral
resiosierone	Articoagularits	anticoagulants may decrease anticoagulant requirements.
Testosterone	Antidiabetic drugs	In diabetic patients, the metabolic effects of androgens may
resiosierone	(including insulin)	decrease blood glucose and insulin requirements.
Tostostorono	ovyphophutazopo	Concurrent administration of oxyphenbutazone and androgens
Testosterone	oxyphenbutazone	may result in elevated serum levels of oxyphenbutazone.
testosterone	adrenocorticotropin &	Concurrent administration of androgens with adreno-
lesiosierone	corticosteroids	corticotropin or corticosteroids may enhance edema formation.
testosterone	propranolol	Administration of testosterone cypionate in a PK study led to an
lesiosierone	propranolol	increased clearance of propranolol.
tootootorono	triamcinolone	Pretreatment of the skin with triamcinolone ointment
testosterone	ointment	significantly reduced testosterone absorption from the patch
patch	Ollittiletit	drug delivery system.

PK=pharmacokinetic

Dosage and Administration

Table 8. Dosing and Administration 1-9

Generic Name	Adult Dose	Pediatric Dose	Availability	
Testosterone buccal mucoadhesive system (CIII)	(congenital or acquired in males) esive or Hypogonadotropic		Buccal mucoadhesive system: Striant®: 30 mg	
Testosterone gel (CIII)	Hypogonadism, primary (congenital or acquired in males) or Hypogonadotropic hypogonadism in males (congenital or acquired):	Safety and efficacy in males <18 years have not been established.	Metered dose pumps: AndroGel® 1%: 12.5 mg/actuation AndroGel® 1.62%:	



Generic Name	c Name Adult Dose		Availability	
	Testim® 1% & AndroGel® 1% gel: Initial: 5 g applied once daily (preferably in the morning); Maintenance: 5 g to 10 g per day; Maximum: 10 g per day AndroGel® 1.62% gel: Initial: 40.5 mg applied once daily (preferably in the morning); Maintenance: 20.25 mg to 81 mg per day; Maximum: 10 g per day Fortesta® gel: Initial: 40 mg applied once daily (preferably in the morning); Maintenance: 10 mg to 70 mg per day; Maximum: 70 mg per day Vogelxo® gel Initial: 50 mg applied once daily (at that same time each day) Maintenance: 50 mg to 100 mg per day Maximum: 100 mg Recommended application sites: Testim®: shoulders and/or upper arms AndroGel 1%: shoulders and/or upper arms and/or abdomen AndroGel 1.62%: upper arms and/or shoulders Fortesta®: thighs Vogelxo®: shoulders and/or upper	Dose	20.25 mg/actuation Fortesta®: 10 mg/actuation Vogelxo® topical gel: 12.5 mg/actuation Unit-dose packets: AndroGel® 1%: 25 mg/pack 50 mg/pack AndroGel® 1.62%: 20.25 mg/pack 40.5 mg/pack Vogelxo® topical gel: 50 mg/pack Unit-dose tubes: Testim® 1%: 50 mg/tube Testosterone 1%: 50 mg/tube Vogelxo® topical gel: 50 mg/tube Vogelxo® topical gel: 50 mg/tube	
Testosterone implant pellet (CIII)	Arms Hypogonadism, primary (congenital or acquired in males) or Hypogonadotropic hypogonadism in males (congenital or acquired): Testopel® implant pellet Initial, Maintenance: 150 to 450 mg (2 to 6 pellets) SQ every 3 to 6 months administered by a health care professional	Safety and efficacy in males <18 years have not been established.	Implant Pellet: Testopel® 75 mg	





Generic Name	Generic Name Adult Dose		Availability
	Delayed puberty in males: Generally dosing is in the lower range of that listed above and, for a limited duration (i.e. 4 to 6 months).		
Testosterone solution (CIII)	Hypogonadism, primary (congenital or acquired in males) or Hypogonadotropic hypogonadism in males (congenital or acquired): Axiron® solution Initial: 60 mg applied once daily to the axilla in the morning; Maintenance: 30 mg to 120 mg once daily; Maximum: 120 mg daily Application site: axilla	Safety and efficacy in males <18 years have not been established.	Meter Dose Pump: Axiron®: 30 mg/pump
testosterone transdermal system (CIII)	Hypogonadism, primary (congenital or acquired in males) or Hypogonadotropic hypogonadism in males (congenital or acquired): Androderm® patch: Initial: 4 mg/day patch applied once nightly; Maintenance: 2 mg/day to 6 mg/day applied at night Application site: back, abdomen, upper arms, or thighs	Safety and efficacy in males <18 years have not been established.	Transdermal system: Androderm®: 2 mg/day patch 4 mg/day patch

Clinical Guidelines

Table 9. Clinical Guidelines Using the Androgens

Clinical Guideline	Recommendations
The American Association of Clinical Endocrinologists (AACE): Medical Guidelines for Clinical Practice for the Evaluation and Treatment of Hypogonadism in Adult Male Patients (2002) ¹³	 Testosterone replacement therapy (TRT) should maintain testosterone levels within the physiologic range (280 and 800 ng/dL). TRT can be used in men with hypogonadism who are not interested in fertility or who are not able to achieve fertility. Treatment of men with hypogonadism with TRT results in increased sexual interest and increased number of spontaneous erections. Secondary sex characteristics (i.e., increased muscle mass, beard growth, growth of pubic and axillary hair, and phallus growth) improve with TRT. In adolescent male patients with hypogonadotropic hypogonadism, TRT increases bone mineral density in comparison with that in male patients with hypogonadism not receiving TRT. In prepubertal-onset hypogonadotropic hypogonadism, diminished bone mass may be only marginally improved by TRT.



Clinical Guideline	Recommendations
	No specific recommendations can be made on the possible normalization of
	growth hormone levels in elderly men with TRT. Further research is needed
	to clarify the potential risks and benefits associated with therapy.
	Whether TRT in men with hypogonadism increases, decreases, or has a neutral effect on cardiovascular risk remains uncertain.
	Orally administered testosterone is quickly metabolized by the liver and appear achieves sufficient blood levels ever time to be useful. The grally
	cannot achieve sufficient blood levels over time to be useful. The orally administered alkylated androgen preparations currently available in the
	Unites States are generally not recommended because of poor androgen
	effects, adverse lipid changes, and hepatic side effects, such as
	hemorrhagic liver cysts, cholestasis, and hepatocellular adenoma.
The Endocrine	TRT is recommended for symptomatic men with classical androgen
Society:	deficiency syndromes to induce and maintain secondary sex characteristics
Clinical Practice	and to improve their sexual function, sense of well-being, muscle mass and
Guidelines:	strength, and bone mineral density.
Testosterone	TRT is not recommended for use in patients with breast or prostate cancer.
Therapy in Adult Men	TRT is not recommended without further urological evaluation in patients
With Androgen	with palpable prostate nodule or induration or a prostate specific antigen
Deficiency	(PSA) 4 ng/mL or PSA 3 ng/mL in men at high risk of prostate cancer (i.e.,
Syndromes (2010) ¹⁴	African Americans or men with first degree relatives with prostate cancer).
	TRT is not recommended in patients with a hematocrit >50%, untreated
	severe sleep apnea, severe lower urinary tract symptoms, uncontrolled or
	poorly controlled heart failure or in those desiring fertility).
	Initiating TRT is recommended with any of the following regimens after
	evaluating patient preference, consideration of pharmacokinetics, treatment
	burden, cost:
	 Testosterone enanthate or cypionate: 75 to 100 mg IM weekly; or 150 to 200 mg IM every two weeks.
	 Testosterone patches: one or two 5-mg non-genital patches applied
	nightly over the skin of the back, thigh, or upper arm, away from
	pressure areas.
	o Testosterone 1% gel: 5 to 10 g applied daily over a covered area of non-
	genital skin (patients should wash hands after application).
	o Testosterone buccal: apply one 30 mg tablet to buccal mucosa every 12
	hours. o Testosterone pellets implanted subcutaneously at intervals of 3 to 6
	months; the dose and regimen vary with the formulation used.
	 Oral testosterone undecanoate, injectable testosterone undecanoate,
	testosterone-in-adhesive matrix patch, and testosterone pellets where
	available. (Note: testosterone undecanoate is not available in the United
	States.)
	Monitoring is advised three to six months after treatment initiation and then
	annually to assess symptom response, the presence of any adverse effects,
	and to check compliance.
	Recommendations aim at achieving serum testosterone levels during
	treatment in the mid-normal range. In men receiving testosterone enanthate or cypionate, aiming for testosterone levels between 400 and 700 ng/dL one
	week after the injection is recommended.
	Hematocrit monitoring is advised at baseline, at three to six months, then
	annually; if exceeds 54% therapy should be discontinued until reduced to a
	safe level.
	Bone mineral density testing of the lumbar spine, femoral neck, and hip
	after one to two years of testosterone therapy is advised in hypogonadal
	men with osteoporosis or low trauma fracture.





Clinical Guideline	Recommendations		
	 Digital rectal exam is advised in men ≥ 40 years with a baseline PSA > 0.6 		
	ng/mL, prior to initiating therapy, at three to six months, and then based		
	upon evidence-based guideline recommendations.		
	Urological consultation is advised if there is an increase in serum or plasma PSA > 1.4 ps/psl, within any 1.2 month period of technological consultation.		
	PSA > 1.4 ng/mL within any 12-month period of testosterone treatment; a PSA velocity of more than 0.4 ng/mL·yr using the PSA level after six		
	months of testosterone administration as the reference (PSA velocity should		
	be used only if there are longitudinal PSA data for more than two years);		
	detection of a prostatic abnormality on digital rectal examination; or a		
	AUA/IPSS score >19.		
	 TRT should be offered to men with low testosterone levels and low libido to 		
	improve libido and to men with erectile dysfunction (ED) who have low		
	testosterone levels after evaluation of underlying causes of ED and		
	consideration of established therapies for ED. TRT should not be offered to all older men with a low testosterone level.		
	 Clinicians should consider offering TRT on an individualized basis to older 		
	men with low testosterone levels on more than one occasion and clinically		
	significant symptoms of androgen deficiency.		
	Short-term TRT may be considered as adjunctive therapy in HIV-infected		
	men with low testosterone levels and weight loss to promote weight		
	maintenance and gains in lean body mass and muscle strength.		
	Short-term TRT may be offered to men receiving high dose glucocorticoids who have law testesteres levels to premote preservation of least hady.		
	who have low testosterone levels to promote preservation of lean body mass and bone mineral density.		
International Society	Late-onset hypogonadism is a clinical and biochemical syndrome associated		
of Andrology (ISA),	with advancing age and characterized by symptoms and a deficiency in		
International Society	serum testosterone levels (below the young healthy adult male reference		
for the Study of the	range). This condition may result in significant detriment in the quality of life		
Aging Male (ISSAM),	and adversely affect the function of multiple organ systems.		
European Association of Urology (EAU),	Response to TRT should be assessed. If there is no improvement of signs waste as within a reasonable time interval (three to give months in adequate).		
European Academy of	symptoms within a reasonable time interval (three to six months is adequate for libido and sexual function, muscle function, and improved body fat; a		
Andrology (EAA),	longer interval is required to see improvement in bone mineral density),		
American Society of	TRT should be withdrawn. Further investigation for other causes of		
Andrology (ASA):	symptoms is then mandatory.		
ISA, ISSAM, EAU,	TRT improves body composition (i.e., decrease of fat mass, increase of		
EAA, and ASA	lean body mass) in men with hypogonadal values of testosterone.		
Recommendations: Investigation,	Secondary benefits of these changes of body composition on strength,		
Treatment, and	muscle function, metabolic, and cardiovascular dysfunction are suggested by available data but require confirmation by large-scale studies.		
Monitoring of Late-	Osteopenia, osteoporosis and fracture prevalence rates are greater in		
Onset Hypogonadism	hypogonadal younger and older men. Bone density in hypogonadal men of		
in Males (2009) ¹⁵	all ages increases under TRT. Fracture data are not yet available and thus		
	the long-term benefit of TRT requires further investigation.		
	Men with erectile dysfunction (ED) and/or diminished libido and		
	documented testosterone deficiency are candidates for TRT. In the		
	presence of a clinical picture of testosterone deficiency and borderline serum testosterone levels, a short (i.e., three months) therapeutic trial may		
	be justified. An absence of response calls for discontinuation of TRT. There		
	is evidence suggesting therapeutic synergism with combined use of TRT		
	and phosphodiesterase-5 (PDE5) inhibitors in hypogonadal or borderline		
	eugonadal men; however, these observations require additional study. The		
	combination treatment should be considered in hypogonadal patients with		
	ED failing to respond to either treatment alone. It is unclear whether men		





Clinical Guideline	Recommendations
Official Guideline	with hypogonadism and ED should be treated initially with testosterone,
	PDE5 inhibitors, or the combination.
	Currently available intramuscular (IM), subdermal, transdermal, oral, and
	buccal preparations of testosterone are safe and effective. The treating
	physician should have sufficient knowledge and adequate understanding of
	the pharmacokinetics as well as of the advantages and drawbacks of each
	preparation. The selection of the preparation should be a joint decision of
	an informed patient and physician.
	Short-acting preparations may be preferred over long-acting depot
	preparations in the initial treatment of patients with late-onset
	hypogonadism because of the possible development of an adverse event
	that may require rapid discontinuation of TRT.
	Inadequate data are available to determine the optimal serum testosterone
	level for efficacy and safety. For the present time, mid-to-lower young adult
	male serum testosterone levels seem appropriate as the therapeutic goal.
	Sustained supraphysiological levels should be avoided. No evidence exists
	for or against the need to maintain the physiological circadian rhythm of
	serum testosterone levels.
	The 17-α-alkylated androgen preparations such as methyltestosterone are
	obsolete because of their potential liver toxicity and should no longer be
	prescribed.
	Due to insufficient data regarding the therapeutic and adverse effects of
	human chorionic gonadotropin treatment in older men and its higher cost, the treatment cannot be recommended in late-onset hypogonadism except when
	fertility is an issue. Antiestrogens and aromatase inhibitors have been shown
	to increase endogenous testosterone levels. Adequate evidence does not
	exist to recommend their use.
	TRT is contraindicated in men with prostate or breast cancer. TRT is
	relatively contraindicated in men at high risk of developing prostate cancer.
	It is unclear whether localized low-grade prostate cancer represents a
	relative or absolute contraindication for treatment.
	Men with significant erythrocytosis, untreated obstructive sleep apnea, and
	untreated severe congestive heart failure should not be started on TRT
	without prior resolution of the comorbid condition.
	Age is not a contraindication to initiate TRT. Individual assessment of
	comorbidities (as possible causes of symptoms) and potential risks versus
	benefits of TRT is particularly important in elderly men.
American College of	Treatment with a phosphodiesterase type 5 (PDE5) inhibitor should be
Physicians: Hormonal	initiated in men who seek treatment for erectile dysfunction and who do not
Testing and	have a contraindication to therapy.
Pharmacologic Treatment of Erectile	The clinical benefit associated with the use of PDE5 inhibitors was
Dysfunction (2009) ¹⁶	demonstrated regardless of the cause (such as diabetes, depression, or prostate cancer) or baseline severity of erectile dysfunction.
Dysidifiction (2003)	 Improvement in erectile functioning was related to higher doses for sildenafil
	and vardenafil but not for tadalafil; however, higher doses were associated
	with a greater risk for adverse events.
	There is insufficient evidence to compare the efficacy and adverse events of
	the different PDE5 inhibitor agents.
	The choice of which PDE5 inhibitor to administer should be made based on
	the individual preferences of men with erectile dysfunction, including the
	ease of use, cost, and tolerability.
	Due to inconclusive evidence, there are no recommendations against or for
	routine use of hormonal blood tests or hormonal treatment
	(i.e., testosterone oral, injection, gel, patch, and cream) in the management





Clinical Guideline	Recommendations		
	 of erectile dysfunction. Clinicians should individualize decisions to measure hormone levels on the basis of clinical presentation and physical findings that suggest hormonal abnormality. There is insufficient evidence to determine whether PDE5 inhibitors are associated with an increased risk for non-arteritic anterior ischemic optic neuropathy. 		

Conclusions

The testosterone products included in this review are Androderm[®], AndroGel[®], Axiron[®], Fortesta[®], Striant[®], Testim[®], Testopel[®] and Vogelxo[®]. These agents primarily differ in their formulations and site of administration. Different formulations include the topical gels, solutions and transdermal patches in addition to a mucoadhesive buccal tablet and an implantable pellet. Currently, only AndroGel® has an A-rated generic formulation. All of the products are indicated for testosterone replacement therapy in males with primary hypogonadism (congenital or acquired) or hypogonadotropic hypogonadism (congenital or acquired) with Testopel® (testosterone) implantable pellets also having an indication to stimulate puberty in certain carefully selected males with clearly delayed puberty. 1-5

Available head-to-head studies suggest that Testim[®] and AndroGel[®] may produce higher serum testosterone concentrations, and reduce body fat more so compared to Androdem. ¹⁹⁻²² One study suggests that patients with a suboptimal response to AndroGel® may experience symptomatic improvements in libido, erectile function and energy levels following a switch to Testim[®]. ²³ No studies are available that evaluate Axiron® or Fortesta® compared to other androgens or topical testosterone products. The results from a meta-analysis demonstrated that the transdermal patch showed the greatest rate of erectile response compared to the (intramuscular) IM and oral formulations of testosterone, with the IM and oral products showing essentially equivalent response rates.31

According to current consensus guidelines. IM and topical testosterone preparations are generally recommended for the management of hypogonadism in adult male patients while the oral androgen therapies are generally not recommended for this condition due to poor androgen effects, adverse lipid changes, and hepatic side effects. 13,15 The selection of a specific testosterone replacement therapy should be a joint decision between an informed patient and physician after considering patient preferences, the pharmacokinetic profiles of the respective agents, treatment burden, and cost. Furthermore, currently available guidelines do not give preference to one topical preparation versus another.

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