ABSTRACT

Hormonally induced azoospermia is an effective, reversible form of male contraception; however, some men treated with weekly intramuscular injections of testosterone enanthate (TE) injections fail to become azoospermic. As weekly injections cause wide fluctuations in serum testosterone levels, we tested the hypothesis that more stable, physiological testosterone levels would consistently produce azoospermia. Using a depot testosterone formulation which provides stable, physiological range testosterone levels for up to 6 months, we studied nine men before and after insertion of six 200 mg testosterone implants under the abdominal wall skin and compared the results with 38 men treated in a previous study with weekly intramuscular injections of 200 mg TE. Testosterone implants suppressed sperm output to near-azoospermia between the second and fourth postimplant months returning to normal by the sixth postimplant month. The fall in sperm output at the first month was greater after testosterone implants than TE injections (mean 68%, se 17%, \( P = 0.011 \)) but similar proportions of men became azoospermic (9% vs. 15%) or severely oligozoospermic (<1 million/ml; 4% vs. 7%). Plasma testosterone and estradiol levels remained mostly within the eugonadal range after implants but were markedly supraphysiological during TE injections. Both treatments suppressed immunoreactive LH and FSH to undetectable levels by ultrasensitive fluorimmunoassay. Sex hormone-binding globulin levels were decreased and PRL levels increased by TE injections but neither was changed by testosterone implants. Prostate-specific antigen demonstrated a small rise of marginal significance \( (P = 0.065) \) after testosterone implants. Fewer men experienced acne after implants (\% vs. 2, \( P = 0.0004 \)). Therefore a depot testosterone preparation with quasi-zero-order release demonstrates higher dose efficiency with similar (but not uniform) efficacy at inducing azoospermia but may cause fewer androgenic side-effects than weekly TE injections. (J Clin Endocrinol Metab 175: 1326-1332, 1992)

A male contraceptive must reduce the number of fertile sperm ejaculated sufficient to prevent fertilization. Hormonal methods aim to reduce sperm output reversibly through inhibiting pituitary gonadotropin secretion and consequent depletion of intratesticular testosterone. Testosterone-induced azoospermia provides highly effective and sustainable contraception (1); however only 65% of men treated with weekly testosterone enanthate (TE) injections become consistently azoospermic whereas the remainder exhibited various degrees of oligozoospermia (1-3). The explanation for the nonuniform induction of azoospermia by testosterone remains unknown however it has been proposed (4) that unfavorable pharmacokinetics of TE, the prototype androgen employed in most male contraceptive studies, lead to supraphysiological circulating testosterone levels that prevent the adequate depletion of intratesticular testosterone. The practical requirement for longer interinjection interval led to development of new long-acting depot testosterone preparations (5) but their effects on human spermatogenesis have not yet been determined. This study aimed to determine the effects of a long-acting depot testosterone preparation (6-8) on human spermatogenesis and test the hypothesis that steady-state testosterone levels within the physiological range might achieve azoospermia uniformly in fertile men.

Materials and Methods

Study design and procedures

The study aimed 1) to describe prospectively the effects of testosterone implants and 2) to compare retrospectively with men receiving weekly injections of 200 mg TE (Testoviron, Schering AG, Berlin, Germany) in the Sydney centre of a World Health Organization multicenter efficacy study of hormonal male contraception (1). The two studies drew from the same pool of volunteers but were consecutive in sequence rather than randomized. Recruitment for the implant study commenced immediately after termination of intake to the WHO study. The design and overall outcomes for the WHO study have been described in detail elsewhere (1). In the implant study subjects provided two baseline sets of semen and blood samples at least 2 weeks apart before implantation of six 200 mg (total dose 1200 mg, release rate 9 mg/day) pellets of fused crystalline testosterone (Organon, Sydney, Australia) subdermally in the lower abdominal wall under local anaesthesia as described (7, 8). The pellets, being composed of only crystalline testosterone, are fully biodegradable and do not require removal. Subsequently subjects provided monthly semen and blood samples until their sperm density reached pretreatment baseline geometric mean or consistently normal levels (>20 million sperm/mL). In men receiving weekly TE injections, blood was sampled at 7 days after the last injection during the visit to receive the next dose. Both studies were approved by the Royal Prince Alfred Hospital Human Ethics Review Committee.

Subjects

Healthy men aged between 21 and 50 yr of age free of chronic medical illness, not taking regular medication and having normal testicular function were recruited. Exclusion criteria were any history of gonadal dysfunction (including infertility) or other abnormalities in...
medical screening tests. Subjects were advised to continue reliable contraception throughout this study to avoid conception.

Assays

Semen collected by masturbation was analyzed within 60 min according to methods described in the WHO Semen Manual (9) using a Makler chamber. Assays of total and free testosterone, estradiol, PRL, and sex hormone-binding globulin (SHBG) were performed by standard immunoassays described previously (7, 10). LH and FSH were assayed by an ultrasensitive time-resolved fluoroimmunoassay (Delfia, Pharmacia, Piscataway, NJ) with a detection limit of 0.1 U/L for both LH and FSH. Prostate-specific antigen (PSA) was measured by immunoassay (DPC, Los Angeles, CA). Between-assay coefficients of variation ranged from 6–12% for all assays. Eugonadal ranges for semen and hormone assays were determined from a contemporary and ongoing study of men (n = 394) screened as potential sperm donors (11).

Data analysis

Results are expressed as mean ± SEM. Data were analyzed by a two-way analysis of variance or Fisher's test as appropriate (12). Semen data were cube-root transformed and hormonal data log-transformed to normalize the distribution and stabilize variance. The degree of suppression of sperm output (azoospermia or oligozoospermia) was defined on the basis of the lowest recorded monthly sperm density.

Results

Subjects

The men entering the testosterone implant (n = 9) and WHO TE injection (n = 38) studies were similar in height, weight, body surface area, body mass index, and testis size and neither group differed significantly in these anthropometric variables from the control group of healthy men (n = 394) screened as potential sperm donors (data not shown).

Implantation of testosterone pellets was very well tolerated (7, 8) and none of the men complained of any side-effects including changes in sexual function, bleeding, infection, or extrusion. Fused crystalline implants do not cause irritation or fibrosis unlike the compressed, cholesterol-containing pellets originally used in the United States (8). Among men having weekly TE injections, four discontinued within 4 months for medical reasons including two due to discomfort and intolerance of injection frequency and two due to acne. Acne (defined as any degree of typical pimples on back, chest, or face) was not observed in men having implants whereas it was common in men receiving TE injections (%

<table>
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<tr>
<th>Assay</th>
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<th>During</th>
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<td>781 ± 65</td>
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* Significant (P < 0.05) difference from baseline. Baseline is defined as the mean of the two baseline measurements. During treatment is defined as measurements made at 3 months after the start of either treatment. After treatment is defined as measurements made at 3 months after cessation of testosterone enanthate injections or at 8 months after testosterone implantation. Results expressed as mean and SEM.

Reproductive hormones and SHBG

After testosterone implantation, plasma testosterone levels increased (Fig. 1) which remained at near zero levels between the 2nd and 4th month after implantation. Essentially identical patterns were observed whether expressed as concentrations or total output of motile, morphologically normal, or of all sperm. The fall in sperm output was similar to that observed among men having TE injections except that the rate of fall (expressed as a percentage of geometric mean baseline sperm density) was significantly greater at the first month in men bearing implants (17 ± 6% vs. 58 ± 7%, P = 0.011). After the 5th month after testosterone implantation, sperm output returned to normal whereas mean sperm density remained suppressed on men continuing on TE injections. Similar proportions of men achieved azoospermia [% (56%) vs. %, 66% (66%)] or severe oligozoospermia [<1 million sperm/mL, % (100%) vs. % (97%)]. The implant study provided a power of 80% to exclude the hypothesis that testosterone implants consistently induce azoospermia but, due to the limited study sample size, the power was too low however to exclude an improved rate of achieving azoospermia but to less than 100%.

TABLE 1. Effects of testosterone on reproductive hormones from two studies.
Fig. 1. Time course of sperm output (expressed as sperm density in million sperm per milliliter) before and after either implantation of six 200 mg (1200 mg total) testosterone pellets in nine healthy fertile men in the implant study (filled circles) and in a previous WHO study of weekly IM injection of 200 mg testosterone enanthate (open squares). Time of commencement of treatment is indicated by the triangular symbol. Results expressed as mean and SEM.

Fig. 2. Plasma total (upper panel) and free (middle panel) testosterone and gonadotropins (lower panel) before and after either implantation of six 200 mg (1200 mg total) testosterone pellets in nine healthy fertile men. In lower panel plasma LH is indicated by filled circles and FSH by open circles. Dashed lines indicate the eugonadal range as determined from healthy volunteers (n = 394) screened as potential sperm donors. Results expressed as mean and SEM.

(1015 pm) when levels just exceeded the upper limit of the eugonadal range (35 nm for total, 1000 pm for free testosterone). Plasma LH and FSH were suppressed to undetectable levels from the 1st until the 4th month when levels increased significantly by the 5th and returned to baseline by the 6th month. Plasma estradiol levels (Fig. 3) rose to peak at the 1st month but remained within the eugonadal range. Neither PRL nor SHBG levels changed significantly after testosterone implantation. After TE injections, plasma total and free testosterone and estradiol were markedly higher whereas PRL levels increased and SHBG levels fell (Table 1). Plasma LH and FSH were both suppressed to virtually undetectable levels during continued treatment.

Metabolic effects of testosterone

The biochemical effects of testosterone on lipids, urea, calcium, albumin total protein, and hemoglobin were similar in magnitude for implants and TE injections; however due to larger sample size, these effects were statistically significant for TE injections only (Table 2). There were no significant effects of either testosterone treatment on electrolytes.
Fig. 3. Plasma estradiol (upper panel), SHBG (middle panel), and PRL (lower panel) before and after either implantation of six 200 mg (1200 mg total) testosterone pellets in nine healthy fertile men. All values remain within the eugonadal range at all times. Results expressed as mean and SEM.

(Na, K, Cl, HCO₃), glucose, phosphate, liver function tests (bilirubin, alkaline phosphatase, 3 transaminases), creatinine, uric acid, leukocyte, or platelet counts.

Discussion

A recent WHO multicenter study demonstrated that testosterone-induced azoospermia provides highly effective and reversible contraception with minimal side effects (1). In countries such as Indonesia (14), India (15), and China (1) where testosterone induces azoospermia in virtually all fertile men, testosterone can constitute the basis of a reliable, reversible hormonal male contraceptive. In developed countries, however, weekly injections of 200 mg TE induces azoospermia in only 60% of fertile men (1-3) with the remaining men exhibiting various degrees of severe oligozoospermia (1-3). Although residual sperm are unable to penetrate zona-free hamster oocytes in vitro (16, 17) and azoospermia may therefore not be necessary for reliable contraception (18), until the contraceptive efficacy of testosterone-induced oligozoospermia is firmly established it remains an unexplained limitation of a testosterone-based regimen that not all men become azoospermic.

Virtually all previous studies of the effects of exogenous steroids on human spermatogenesis have used either injectable and/or oral androgens (1-3, 19), most commonly weekly IM doses of 200 mg TE. The failure to produce azoospermia uniformly among fertile men treated with TE cannot be attributed to the dose or frequency of injections. Increasing the weekly dose above 200 mg does not further increase the proportion achieving azoospermia (3) whereas prolonging the interinjection interval fails to sustain suppression of spermatogenesis (2). From the single-dose pharmacokinetics of TE (20), where peak testosterone reaches supraphysiological levels at 48 h and returns to baseline levels over 10-14 days, both simulation (21) and empirical (22) studies indicate that chronic weekly administration results in marked fluctuations and frequently supraphysiological plasma testosterone levels. Such supraphysiological circulating testosterone levels might counteract the depletion of intratesticular testosterone necessary for effective hormonal suppression of spermatogenesis (4). If supraphysiological peripheral testosterone levels raised the interstitial and seminiferous tubular testosterone levels (23), this might maintain sufficient androgen effects on Sertoli cells to continue low levels of spermatogenesis. Although sex steroid levels were considerably higher in men having TE injections, these kinetic considerations also indicate that the nadir levels reported in this study at 7 days after the last TE injection actually underestimate the mean integrated testosterone levels to which subjects were exposed throughout the week during regular weekly TE injections. Whereas the testosterone levels at 3 months after testosterone pellet implantation are also lower that peak levels (7, 8), this fall-off from peak testosterone levels is much less and therefore the discrepancy between integrated testosterone and estradiol levels on the two regimens is even greater than indicated by Table 1.

Testosterone implants provide the first opportunity to test the effects of steady-state testosterone levels on normal human spermatogenesis. These implants provide virtually
zero order release kinetics ensuring stable, physiological testosterone levels for up to 6 months after subdermal implantation (6-8). This duration of action is significantly longer than other depot testosterone formulations including testosterone microcapsules (24, 25) or testosterone buccinate (5) which would need to be administered in multiple overlapping doses to maintain suppression for more than the full duration of a human spermatogenic cycle. The present study shows that a long-acting depot testosterone formulation can effectively and reversibly suppress human spermatogenesis. This suppression of spermatogenesis is achieved without mean testosterone levels exceeding the eugonadal range and at a lower daily testosterone dose than either conventional (1-3) or even lower TE dose (3) injections. Furthermore lower and less fluctuating testosterone levels from the depot preparation caused fewer hormonal side-effects while effecting an equivalent degree of spermatogenic suppression and similar biochemical effects as that of weekly TE injections. Nevertheless testosterone implants still did not produce azoospermia uniformly in all men. This finding suggests it is unlikely that other newly developed depot testosterone formulations such as biodegradable testosterone microspheres (24, 25) and new long-acting testosterone esters (5) would solve the problem of the nonuniformity of induction of azoospermia in testosterone-based regimens. One caveat is however that a smaller dose of testosterone implanted with a lower daily release rate, if equally effective at maintaining gonadotrophin suppression, may yet produce uniform azoospermia and further studies at a lower dose would be required to clarify this issue.

This study underlines the fact that the testicular mechanisms involved in the testosterone-induced suppression of human spermatogenesis are still poorly understood. Classically, endocrine effects on spermatogenesis are mediated via the action of pituitary FSH and testosterone on Sertoli cells (26) and thereby indirectly on the later stages of spermatogenesis (27). Exogenous testosterone suppresses hypothalamic GnRH, and consequently pituitary LH and FSH, secretion through negative feedback inhibition and the cessation of LH secretion reduces Leydig cell testosterone biosynthesis causing a depletion of intratesticular testosterone. By this means exogenous testosterone causes indirectly a potentially reversible withdrawal of both major hormonal stimuli, FSH, and intratesticular testosterone, which provide the dual endocrine drive to Sertoli cells in their role of maintaining, nourishing, and coordinating spermatogenesis. Empirically however testosterone-induced suppression of spermatogenesis is more complex. In rats, testosterone has biphasic effects on spermatogenesis with low doses causing suppression while higher doses continue to maintain sperm production in the absence of gonadotrophin secretion (26, 27) suggesting that sufficiently high intratesticular testosterone levels may substitute for some FSH effects required to maintain spermatogenesis (26).

Although the usually prevailing intratesticular testosterone levels in the human testis may be too high to reproduce by exogenous injections of testosterone (27, 28), intratesticular testosterone levels obligatory for maintenance of rat spermatogenesis are only 5-20% of those normally present in the testis (23, 29). Thus if a hormonal regimen caused depletion of intratesticular testosterone to just below threshold levels required to maintain spermatogenesis (30, 31), any accompanying supraphysiological circulating testosterone levels could then approach or exceed the intratesticular threshold to reinitiate spermatogenesis. This mechanism has been invoked to explain how testosterone injections can partially induce spermatogenesis in immature (32), stalk-sectioned (33), or hypophysectomized (34) monkeys and that the coadministration of testosterone with a GnRH analog can retard the fall of sperm output in nonhuman primates (4) and men (35). The present study however does not support this hypothesis in men since steady-state testosterone levels which stayed mostly within the eugonadal range failed to achieve higher rates of azoospermia than did testosterone injections. If the high-normal testosterone levels maintained by the testosterone implants in this study are still too high, then it is unclear if lower testosterone levels can be achieved that would still provide adequate androgen replacement and complete gonadotrophin suppression required for near-total inhibition of spermatogenesis. The failure of stable, physiological testosterone levels to produce azoospermia uniformly therefore argues that the reasons for the inconsistency of testosterone-induced azoospermia is more likely to reflect intrinsic differences in spermatogenesis rather than general metabolic actions of testosterone.

Another interesting finding from this study is that TE
injections induced more hormonal changes than did the
testosterone implants. This is evident in the greater preval-
ance of acne as well as the decrease in SHBG and increase in
PRL exhibited by men receiving TE injections compared
with men having testosterone implants. In contrast the bio-
chemical effects were generally of similar magnitude. Where
hormonal effects were greater, this may relate to the supra-
physiological testosterone and/or higher estradiol levels pro-
duced by regular TE injections compared with the more
physiological levels maintained by the testosterone implants.
It remains unclear whether these effects are due to testoster-
one itself or to its 5-α reduced metabolite, 5-α dihydrotest-
osterone (DHT), acting upon androgen receptors and/or upon
estrogen receptors following aromatization of testosterone to
estradiol. DHT levels were not measured in this study how-
ever it is unlikely that they differed between regimens. The
subdermal implants are placed well below hair follicles which
contain dermal 5-α reductase and transdermal passage of
testosterone across trunci skin does not increase circulating
DHT levels (36) in contrast with passage across scrotal skin
where 5-α reductase levels are high (37). In any case our
findings do indicate that the lower effective daily dose of
testosterone produced by the depot testosterone preparation
has the advantage of less hormonal side-effects while exhib-
iting the same degree of suppression of spermatogenesis.
This dose-sparing effect of a steady-state formulation may
reduce the risks of potential long-term side-effects of andro-
gens such as cardiovascular or prostatic disease which need
to be considered in relation to testosterone therapy.

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