URINARY EXCRETION OF ANDROGENIC SUBSTANCES AFTER
INTRAMUSCULAR AND ORAL ADMINISTRATION OF
TESTOSTERONE PROPIONATE TO HUMANS

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The urinary excretion of estrogens following administration of female sex hormones has been
compared with the relief obtained from hot flushes and other phenomena accompanying the meno-
pause (1) and has been studied with regard to the percentage of the administered material which
appears in the urine (3 to 12 per cent) (2). Studies of the excretion of urinary androgens
following the treatment of patients with testosterone propionate are reported below with re-
gard to (a) the rapidity of the appearance and comparative levels of excretion in oral and intra-
muscular treatment, (b) the percentage of recovery in the urine of administered material, and
(c) the evaluation of urinary levels of androgens as criteria of effective levels of male hormone in
the body.

METHODS, MATERIALS, AND SUBJECTS

All urine samples were collected on a 24-hour basis and extracted with benzene in a continuous extractor
according to the method previously described (3). The assays for androgenic activity contained in the urine
extracts were performed on the day-old chick's comb (4). A minimum of 15 and usually 20 chicks were used
for each assay. The accuracy of the determinations is considered to be ±25 per cent and probably greater as
can be calculated from previous data (4). All values are expressed as international units (I.U.) per 24 hours.
The international unit of androgenic activity is that evoked by 0.1 mgm. of androsterone.

The androgen used was testosterone propionate. In-
tramuscular administration employed a standard dose of
20 mgm. dissolved in 1 cc. of peanut oil, since the
amount and the type of oil influences the absorption (5).
Injections were made intramuscularly into the buttocks
or upper arms.

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the Fluid Research Fund of Yale University School of
Medicine, the General Education Board of the Rocke-
feller Foundation, and The International Cancer Research
Foundation.

2 Furnished through the courtesy of the Ciba Company,
under the trade name Perandren.

Crystalline testosterone propionate was given per os
in disc-shaped tablets 0.5 cm. in width, 0.3 cm. in diam-
er. Tablets were taken on an empty stomach at spaced
intervals during waking hours: ½ hour before break-
fast, 1 hour before lunch, 1 hour before dinner, and in
the evening before retiring.

The subjects are 2 men with organic and functional
evidence of deficient testicular secretion. Case 1 is a
35-year-old man who had lived until the age of 35 with
a single descended testis. He has a wife and 2 children.
Following surgical removal of the testis at the age of
35, castration phenomena (6), including the following,
appeared: atrophy of the genitalia, muscular weakness
and fatigue (7), absence of penile erection (8, 9), pre-

cence of vasomotor phenomena such as may be found in
women at the menopause, skin pallor, and inability to
tan (10). Before androgenic administration, the level
of output of male hormone in 4 urine assays covering 7
days, taken from a period of weeks, showed a daily
average of 13.5 I.U. This is far below the levels of
androgen excretion of normal adult men (11, 12, 3).
Still further evidence that if any abdominal testis were
present it did not secrete large amounts of androgens
was the negative clinical response obtained with 16 in-
jections of large amounts of anterior pituitary-like sub-
stance, 100 rat units (R.U.) given three times weekly.

Case 2 is a 27-year-old sexually underdeveloped man,
described in detail elsewhere (8), who exhibited a like
condition save that the genitalia instead of undergoing
atrophy had never developed. The testes did not re-

cpond to injections of 100 R.U. of anterior pituitary-
like material given 3 times weekly for 9 weeks.

RESULTS

A. Intramuscular administration. Intramuscu-
lar injections of 20 mgm. of testosterone propio-

nate were given daily to Case 1 for 30 succes-
sive days. The resultant increased excretion of
urinary androgens is shown in Table I. The
average daily output prior to injections was 13.5
I.U. By the 3d day of injection, the output was
50 I.U. at which time some of the clinical symp-
toms of androgen deficiency remained, particularly
the vasomotor phenomena. This is perhaps cor-
related with the fact that the longer the period
of androgen injection the greater the suppression of pituitary hyperactivity (13).

This high level of androgen excretion was maintained throughout the period of treatment, as shown by assays of 11 out of the 16 remaining days of injection. The average daily output of 12 days, ranging from the 3d to the 30th day was 68.9 I.U., an increase of 510 per cent. In all determinations this level remained within the range of 40 to 100 I.U. which is the range for normal men as determined by several groups of workers (11, 12, 3). After the first few days castration symptoms remained absent throughout the course of treatment. Thus, with the intramuscular injections which maintained normal ranges of urinary androgens, the patient remained symptom-free.

Eight to 9 days after the cessation of injections, the daily output was still 38 I.U. Although this was an increase of 281 per cent over the castrate level and can hardly be considered lower than normal levels, castration signs were again of considerable severity.

Determinations done 30 and 37 days respectively after the end of treatment showed 13 I.U., a return to low pre-injection levels.

B. Oral administration. Daily oral administration of 60 mgm. of testosterone propionate was given to Case 1. An immediate high level of urinary androgen was obtained. A 24-hour specimen between the 36th and 60th hour after ingestion of the first tablet contained 137 I.U., an increase of 124.5 I.U. over the pre-treatment level of daily excretion (Table II). This increase of

<table>
<thead>
<tr>
<th>Day of treatment</th>
<th>Day after cessation of treatment</th>
<th>Urinary androgenic activity</th>
<th>I.U. per day</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td></td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td></td>
<td>21</td>
<td>0</td>
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<tr>
<td>3</td>
<td></td>
<td>50</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td></td>
<td>93</td>
<td>0</td>
</tr>
<tr>
<td>5, 6</td>
<td></td>
<td>47</td>
<td>0</td>
</tr>
<tr>
<td>9, 10</td>
<td></td>
<td>78</td>
<td>0</td>
</tr>
<tr>
<td>12, 13, 25, 26, 27</td>
<td></td>
<td>68</td>
<td>0</td>
</tr>
<tr>
<td>30</td>
<td></td>
<td>94</td>
<td>0</td>
</tr>
</tbody>
</table>

* Intramuscular injections of 20 mgm. of testosterone propionate (equivalent to 1110 I.U.) in 1 cc. of peanut oil were given Case 1 daily for 30 successive days. The urinary output of comb growth-stimulating materials reached the levels of normal men the 3d day of injection and remained so during the 30 days of treatment. Thirty days after the end of injections, the amount in the urine was again down to a pre-injection level. Castration signs disappeared during this treatment.

922 per cent in urinary androgen activity was accompanied by slight, if any, clinical improvement. The amount ingested was then raised to 120 mgm. of testosterone propionate daily and after 3 days the daily androgen excretion was 500 I.U., an increase of 3703 per cent over the pre-treatment level. Castration symptoms still continued.

Case 2 had an average urinary output of 16 I.U. before oral administration was begun. By the 2d day of daily oral administration of 60 mgm. of testosterone propionate, 38 I.U. were excreted in a 24-hour specimen of urine (Table II). Pooled samples for the 7th, 9th, 11th and 12th days of treatment gave an average of 94 I.U. By the 14th day 264 I.U. were excreted, an increase of 1560 per cent over the level before treatment. A moderate amount of clinical relief occurred but it was not equal to that obtained
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with 1/6 of this dose given subcutaneously in injections of 20 mgm. 3 times weekly.

If the samples assayed could be considered representative, a very rough estimate in Case 1 of the percentage of recovery in the urine of comb growth-stimulating material with daily subcutaneous injections might be made as in Table III.

<table>
<thead>
<tr>
<th>TABLE III</th>
<th>Percentage of administered androgenic material recovered in the urine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average urinary androgen in 24 hours</td>
</tr>
<tr>
<td>I.U.</td>
<td>I.U.</td>
</tr>
<tr>
<td>Days 1 to 10 of treatment</td>
<td>67.00</td>
</tr>
<tr>
<td>Days 11 to 30 of treatment</td>
<td>72.33</td>
</tr>
<tr>
<td>Days 1 to 15 after treatment</td>
<td>38.00</td>
</tr>
<tr>
<td>Total excretion</td>
<td>2686</td>
</tr>
<tr>
<td>Total excretion expected over these 45 days without treatment (at 13.5 I.U. daily)</td>
<td>607</td>
</tr>
<tr>
<td>Net excretion</td>
<td>2079</td>
</tr>
</tbody>
</table>

The 600 mgm. of administered testosterone propionate is equivalent to 502 mgm. of testosterone.

If urinary androgen is testosterone:

Since 1 I.U. of testosterone = 0.015 mgm. (cf. Koch (17)) then 2079 I.U. (net excretion) = 31.190 mgm.

Percentage recovery = \( \frac{31.190}{502} \times 100 = 6.2 \text{ per cent} \)

If urinary androgen is androsterone:

Since 1 I.U. of androsterone = 0.100 mgm. then 2079 I.U. (net excretion) = 207.900 mgm.

Percentage recovery = \( \frac{207.9}{502} \times 100 = 41.4 \text{ per cent} \)

If urinary androgen is dehydroisoandrosterone:

Since 1 I.U. of dehydroisoandrosterone = 0.300 mgm. then 2079 I.U. (net excretion) = 623.700 mgm.

Percentage recovery = \( \frac{623.7}{502} \times 100 = 124.2 \text{ per cent} \)

If urinary androgens are an equal mixture of dehydroisoandrosterone and androsterone:

Percentage recovery = \( \frac{313.5}{502} \times 100 = 62.4 \text{ per cent} \)

If the androgenically-active material is excreted in the form of testosterone, the percentage of recovery is 6.2. If it is excreted as androsterone, the recovery is 41.4 per cent, as dehydroisoandrosterone 124.0 per cent, and as equal mixtures of androsterone and dehydroisoandrosterone 62.4 per cent.

The possibility that all of the testosterone propionate is converted into dehydroisoandrosterone is unlikely since, as shown in Table III, this would mean an excretion of more material than was administered.

There is no proof that the pre-treatment level of male hormone secretion by the body was maintained during the injection of the testosterone propionate, especially since this intramuscular administration of androgen greatly decreased the urinary level of gonadotropic (follicle-stimulating) substance in this patient (14). If none of the excreted androgen were of endogenous origin, more of the amount excreted would be considered as recovered from the administered hormone. This would make estimates of the amount recovered as much as ½ higher.

**DISCUSSION**

Even if the figures for recovered hormone are no higher than listed, there remain the obvious facts that (1) daily intramuscular injections of 20 mgm. of testosterone propionate maintained excretion levels at approximately those determined for normal men and that castration symptoms disappeared during this treatment, and (2) orally-administered crystalline testosterone propionate was readily absorbed from the gastro-intestinal tract and eliminated in large amounts through the kidneys; since, however, no satisfactory clinical relief was afforded even when absorption was continued during 16 hours of the day, it seems likely that the material was eliminated with great rapidity. The figures indicate that increasing the oral dose of crystalline testosterone propionate can be expected to result in only exceedingly brief levels of hormone in the fluids and tissues of the body. In the maintenance of effective levels in the body by oral means there must be a balance between slow absorption to avoid prompt elimination by the kidneys and ready absorption to prevent waste in the feces. Furthermore, damage (15, 16) from large doses of male hormone may be incurred, especially in the kidney and the vascular system.

The transient presence of orally-administered testosterone propionate within the body and the rapid appearance of large quantities of androgenic material in the urine raise the question of threshold levels. It may be that when a certain threshold is reached, the hormone above this level is
excreted in the urine. In this connection it is interesting to note that in adrenal virilism in women, enormous titers of androgen activity in the urine may be observed, but the masculinization does not proceed beyond that of the normal male.

The urinary level of androgen is an accurate reflection of the level in the tissues and fluids of the body only when the supply in circulation is at a maintained rate. Temporary presence of a large amount in the body would produce a large urinary excretion. In other words, evaluation of levels of androgens in the body by comparison of one urine titer with another would be misleading if the androgen precursors were not present over the same general length of time. Further, measurements of urinary androgenic activity should be expected to account only with inaccuracy if there were great variations in the amount present in the body during this time.

If the androgens secreted by the body were from the testes and in the form of testosterone and had a percentage of recovery in the urine like that from intramuscular injections in Case 1, there would be secreted in 25 gram testes of a man, 22 I.U. per gram every 24 hours.

SUMMARY

1. In the patient with low levels of testicular secretion both intramuscular and oral administration of testosterone propionate produce an increase in the androgenic activity of the urine.

2. With intramuscular injections of 20 mgm. daily, Case 1 showed an increase in output to normal levels and clinically a disappearance of castration phenomena. A 24-hour titer of 50 I.U. was obtained on the 3d day of administration. The average of 6 assays covering 12 of the 30 days of injection was 68.9 I.U., with a range from 47 to 94 I.U.

3. Tablets per os of 60 to 120 mgm. daily gave 24-hour urine readings as high as 500 I.U. in Case 1, and 264 I.U. in Case 2. These large excretions were not accompanied by as good clinical relief as obtained with 1/6 to 1/4 the amount taken intramuscularly and with lesser androgenic activity of the urine.

4. Absorption of the large amounts of androgen can take place through the gastro-intestinal tract with what appears to be rapid elimination through the kidneys. It is suggested that there may be a threshold for the substance in the body and that rapid disposition is made of an excess. Oral means of administration should be considered from the standpoint of material lost not only through the feces but also excreted in the urine.

5. Urine assays as an indication of the presence of hormone in the body may be misleading if the hormone in the body is present only irregularly. For example, enormous quantities of urinary androgen were found with an oral method that appeared to give only transient levels in the body.

6. A rough estimation of the percentage of the androgenic material recovered in the urine is 6.2 per cent, if it be in the form of testosterone, 41.4 per cent if androsterone, 62.4 per cent if an equal mixture of androsterone and dehydroisoandrosterone.

BIBLIOGRAPHY


11. Dingemanse, E., Borchardt, H. and Laqueur, E., Capon comb growth-promoting substances ("male
16. Hamilton, J. B. and Dorfman, R. I., Influence of various solvents upon the length and strength of the action of synthetic male hormone, testosterone propionate. (To be published.)