LONG-TERM EFFECTS OF FINASTERIDE ON PROSTATE TISSUE COMPOSITION

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ABSTRACT

Objectives. To determine the long-term effects of finasteride treatment on prostate tissue composition; to relate these effects to clinical outcomes; and to test the hypothesis that finasteride exerts a selective or preferential action on the transition zone.

Methods. Nineteen men with symptomatic benign prostatic hyperplasia (BPH) who completed a 6-month double-blind trial of finasteride were enrolled in a 24-month open-label extension study of drug responders. Magnetic resonance imaging and prostate biopsy for morphometric analysis were performed together 70 times: at baseline (n = 19), after treatment periods of intermediate duration (6 to 18 months, n = 32), and after long-term drug treatment (24 to 30 months, n = 19). At baseline, prostate volume averaged 51 cc, of which 57% was transition zone.

Results. Decreases in symptom score, dihydrotestosterone and prostate-specific antigen levels, and prostate volume occurred at 6 months (P < 0.01), stabilized, and were maintained without further long-term decreases. Prostate epithelium contracted progressively from baseline (19.2% tissue composition; 6.0-cc volume; 3.2 stroma/epithelial ratio) to intermediate (12.5%, 3.3 cc, and 5.6, respectively) to long-term treatment (6.4%, 2.0 cc, and 17.4, respectively, P < 0.01 for all). Percent epithelial contraction was similar in the peripheral and transition zones (P = NS). The transition zone remained a relatively constant proportion (53% to 58%) of whole-prostate volume from baseline to long-term observation.

Conclusions. Long-term finasteride treatment (24 to 30 months) results in a marked involution of the prostate epithelium, which continues to progress for many months after clinical effects stabilize. The effect on the epithelium is similar in the peripheral and transition zones for both morphometric and volumetric changes. Progressive contraction of the prostate epithelium appears to constitute the underlying mechanism for sustained action of finasteride.

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of action of the drug. Furthermore, finasteride-treated men remain at risk of developing prostate cancer, which requires a biopsy diagnosis; hence, information about finasteride-induced tissue changes in men with BPH (ie, “control” histologic data) would be important in biopsy interpretation.8,9

Thus, to help clarify the long-term histologic effects of finasteride, we quantified prostatic tissue components in serial biopsies of men with BPH before, during, and after 24 to 30 months of finasteride treatment. By analyzing inner and outer gland tissue separately, the present study was also designed to test the “transition zone hypothesis,” a concept that finasteride may exert a selective or preferential effect on the inner prostate.6,10,11

MATERIAL AND METHODS

STUDY DESIGN

The study design is presented schematically in Figure 1. The study was an open-label extension of a 6-month double-blind trial that was previously published.6 In the original study, men with symptomatic BPH were randomized to receive placebo (n = 15) or finasteride (n = 26) for 6 months. On completion of the 6-month placebo-controlled period, all participants were invited to enroll in a long-term open-label extension study of active finasteride therapy.

Of the original group of 41 men, 39 completed the 6-month double-blind study. Twenty-seven men completed a 1-year extension, and 19 men—the present study group—completed a 2-year extension. Of the 19 long-term participants, 6 were originally randomized to receive placebo for 6 months; as their baseline for the present report, we used those data obtained just before starting active drug. Thus, 6 men were receiving finasteride for 24 months, and 13 were receiving finasteride for 30 months. Study dropouts are explained in Figure 1.

Patients

Baseline demographic data for the 19 men reported herein are presented in Table I. At entry, the patients were middle-aged, with chronic, uncomplicated prostatism, exhibiting some degree of prostate enlargement on rectal examination; an International Prostate Symptom Score (IPSS) of 9 or more; and a serum PSA level less than 10 ng/mL. Exclusion criteria included concurrent use of alpha-adrenergic blocking agents; history of neurogenic bladder or urethral stricture; urinary infection; or any previous invasive treatment for BPH. At the outset of the extension, a consent form, approved by the Western Institutional Review Board, Portland, Oregon, was signed by each participant.

The 19 men were thus highly selected, that is, men who were satisfied with results of treatment for 24 to 30 months, who had no intolerable side effects of finasteride, who developed no cancer or BPH complications mandating surgical intervention, and who remained evaluable throughout the repetitive magnetic resonance imaging (MRI) and biopsy components of the study.

OUTCOME MEASURES

At baseline and at 6, 18, and 30 months, participants had the following tests: voiding symptom score (IPSS); maximal uroflow; serum dihydrotestosterone (DHT) and PSA levels; prostate MRI for whole and zonal volumetrics; and transrectal ultrasound-guided sextant biopsy (18 gauge) for routine histologic evaluation and quantitative image analysis or morphometric analysis. Morphometric analysis was performed in a blinded manner by a study author (A.W.P.) using a technique of computer-assisted color image analysis.6 The deep portion of each biopsy core was stained with India ink to identify the transition zone, a method validated in a separate study reported elsewhere.6

TABLE I. Demographic characteristics of 19 patients at baseline

| Age (yr) | 62.8 ± 6.7 |
| Race     |            |
| White    | 12         |
| African-American | 5         |
| Asian    | 2          |
| IPSS     | 15.9 ± 6.7 |
| Max uroflow (cc/s) | 13.2 ± 4.1 |
| Serum PSA (ng/mL) | 3.0 ± 2.5  |
| Serum DHT (ng/dL) | 45.4 ± 21.0 |
| Prostate volume on MRI (cc) |            |
| Whole gland | 51 ± 28    |
| Transition zone | 29 ± 20   |
| Prostatic tissue composition (%) |            |
| Epithelium | 19.2 ± 7.7  |
| Lumen     | 31.7 ± 6.4 |
| Stroma    | 49.2 ± 9.8 |

Key: DHT = dihydrotestosterone; IPSS = International Prostate Symptom Score; Max = maximal; MRI = magnetic resonance imaging; PSA = prostate-specific antigen.

Data presented are mean value ± SD or number of patients.
men, and percent stroma, as previously described. Tissue volumes were calculated by multiplying the morphometrically determined percent tissue composition times the MRI-determined volume of the prostate or one of its zones. Independent observers, working without knowledge of the other results, obtained the clinical (L.S.M.), volumetric (J.B.G.), histologic (J.I.E.), and morphometric (A.W.P.) data. Methodology for all outcome measures was constant from baseline through the 30-month end point. Detailed methodology of outcome measures used in the present report has been published previously.

DATA PROCESSING AND STATISTICAL ANALYSIS

Because of the crossover study design, which limited the number of patient entries at each point, data were analyzed and reported only at baseline (n = 19) and two end points (“intermediate” and “long term”). The intermediate end point includes measurements made after 6 months (n = 13), after 12 months (n = 6), and after 18 months of finasteride treatment (n = 13). The long-term end point includes measurements made after 24 (n = 6) or 30 months of treatment (n = 13). Thus, 19 data points are averaged for each outcome measure at baseline; 32 at the intermediate point; and 19 at long term, for a total of 70 end-point assessments. In 5 patients MRI data were not available at baseline, and for these patients the MRI volumes were imputed from the ultrasound measurements, according to a regression validated elsewhere. A detailed analysis of the 6-month placebo-controlled data (n = 39) was reported previously.

All data were entered into an IBM-compatible personal computer through Excel spreadsheets and analyzed using the Stata statistical package (Stata Corp., College Station, Tex). The paired t test was used when evaluating test results before versus after treatment. Repeated-measures analysis of variance was used when evaluating serial changes. The relation between continuous variables was tested using the nonparametric Kendall tau correlation. Multivariate analysis was performed using linear regression. Statistical significance was defined a priori as P < 0.05.

RESULTS

Demographic characteristics of the study group at baseline are shown in Table I. In general, the patients were middle-aged men with chronic, uncomplicated, moderate prostatism, who continued to respond favorably to finasteride during 24 to 30 months of treatment.

Symptom score and serum levels of PSA and DHT all decreased during the study (Table II). The decrease from baseline was statistically significant at the intermediate term (P < 0.01) and was maintained thereafter. For these three measures, a slight further decrease observed at the long-term point was not significantly different from that seen at the intermediate point (P = NS). Maximal urinary flow rate increased during the treatment, but the change was not statistically significant in this group of 19 patients.

In Figure 2, prostate volume changes are depicted. Whole-prostate volume decreased from an average of 51.3 cc at baseline to 42.2 cc at the intermediate point, a 17.8% decrease (P < 0.01) and was maintained thereafter. For these three measures, a slight further decrease observed at the long-term point was not significantly different from that seen at the intermediate point (P = NS). Maximal urinary flow rate increased during the treatment, but the change was not statistically significant in this group of 19 patients.

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The effect of finasteride on three different measures of prostate epithelium—percent, volume, and stroma/epithelial ratio—is shown in Table III. At baseline, the average prostate gland was morphometrically determined to be 19.2% epithelial and contained a volume of 6.0 cc of epithelium; the stroma/epithelial ratio (percent stroma/percent epithelium) was 3.2. All three measures of prostate epithelium are seen to change progressively from baseline to the intermediate point and even further from the intermediate to the long-term point. Each

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**TABLE II. Clinical and chemical changes during finasteride treatment**

<table>
<thead>
<tr>
<th></th>
<th>Baseline (mean ± SD)</th>
<th>Intermediate (mean ± SD)</th>
<th>Long-Term (mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPSS</td>
<td>15.9 ± 6.7</td>
<td>9.2 ± 5.8*</td>
<td>6.8 ± 4.1*</td>
</tr>
<tr>
<td>Max uroflow (cc/s)</td>
<td>13.2 ± 4.1</td>
<td>16.5 ± 7.4</td>
<td>14.3 ± 4.5</td>
</tr>
<tr>
<td>PSA (ng/mL)</td>
<td>3.0 ± 2.5</td>
<td>1.5 ± 1.3*</td>
<td>1.2 ± 1.0*</td>
</tr>
<tr>
<td>DHT (ng/dL)</td>
<td>45.2 ± 21.0</td>
<td>11.4 ± 6.2*</td>
<td>9.5 ± 5.8*</td>
</tr>
</tbody>
</table>

Key: Baseline = 19 measurements; Intermediate = 32 measurements at 6 to 18 months; Long-Term = 19 measurements at 24 to 30 months; other abbreviations as in Table I.

* Statistically significant change from baseline (P < 0.01); no changes between intermediate and long-term values are statistically significant.

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**FIGURE 2. Prostate volume changes, determined by magnetic resonance imaging (MRI), during long-term finasteride treatment.** Transition zone volume is determined separately and is seen to approximately parallel changes seen in whole prostate. Changes at the intermediate point are significantly different from baseline values (P < 0.01). Changes at the long-term point remain significantly different from baseline values (P < 0.05) but not from intermediate values.
TABLE III. Changes in prostate epithelium during finasteride treatment

<table>
<thead>
<tr>
<th></th>
<th>Baseline (mean ± SD)</th>
<th>Intermediate (mean ± SD)</th>
<th>Long-Term (mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Epithelium</td>
<td>19.2 ± 7.7</td>
<td>12.5 ± 5.5*</td>
<td>6.4 ± 4.0*</td>
</tr>
<tr>
<td>Epithelial volume (cc)</td>
<td>6.0 ± 2.8</td>
<td>3.3 ± 1.6*</td>
<td>2.0 ± 1.6*</td>
</tr>
<tr>
<td>Stroma/epithelial ratio</td>
<td>3.2 ± 1.8</td>
<td>5.6 ± 3.1*</td>
<td>17.4 ± 13.4*</td>
</tr>
</tbody>
</table>

Key: Abbreviations as in Table II. * Statistically significant change from preceding value (P < 0.01).

During treatment, epithelial and luminal tissues contracted progressively from baseline to intermediate to long-term follow-up. Luminal epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. Luminal epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. Luminal epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. In the transition zone, the epithelial volume decreased 71%, from an average of 5.9 cc before treatment to 1.8 cc at long-term follow-up. The transition zone epithelium demonstrated a 71% decrease, from an average of 5.9 cc before treatment to 1.8 cc at long-term follow-up. The transition zone epithelium demonstrated a 71% decrease, from an average of 5.9 cc before treatment to 1.8 cc at long-term follow-up. stroma/epithelial ratio increased progressively and dramatically during finasteride administration, from 3.2 at baseline to 5.6 at the intermediate point to 17.4 at the long-term point (P < 0.001).

The effect of finasteride on tissue volumes of the separate zones of the prostate is shown in Figure 3. At baseline, the average peripheral zone contained 5.9 cc of epithelium (percent times volume) and 7.8 cc of lumen. The average transition zone contained 2.8 cc of epithelium and 7.1 cc of lumen. During treatment, epithelial and luminal tissues both contracted progressively from baseline to intermediate to long-term (P < 0.01); the effect is statistically significant in both the peripheral and transition zone (P < 0.01). Peripheral zone epithelium demonstrated a 71% decrease, from an average of 5.9 cc before treatment to 1.8 cc at long-term follow-up. In the transition zone, the epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. Luminal epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. Luminal epithelial volume decreased 64%, from 2.8 cc at baseline to 1.0 cc at long-term follow-up. The patient-to-patient and temporal variability of epithelial response is illustrated in Figure 4. In Figure 4A, the most typical response pattern, the epithelium was suppressed in a progressive manner, that is, lower at the intermediate point than at baseline and lower at the long-term point than at the intermediate point (n = 7). In Figure 4B, there was epithelial suppression at the intermediate point, with little additional change at the long-term point (n = 5). In Figure 4C, epithelial suppression occurred as a delayed response and was notable only after the intermediate point (n = 5). In Figure 4D, no change was observed (n = 2). Ultimately, reduction in epithelial composition by 50% or more (baseline versus long-term end point) was found in 17 of the 19 patients.

In a multivariate regression analysis, we found that at baseline, prostate volume was directly correlated with serum PSA level and both percent and volume of epithelium, that is, the large prostate glands were associated with high serum PSA levels and with both a relative and an absolute epithelial richness. Long-term change in prostate volume was directly correlated with baseline prostate volume, serum PSA level, and both percent and volume of epithelium. For all these associations, correlation coefficients ranged between 0.39 and 0.83 (P < 0.05). When transition zone measures were substituted for whole-prostate measures in this analysis, the correlations were similar or weaker than when whole prostate was used. In other words, baseline characteristics of the transition zone were no better an index of finasteride response than those of the whole prostate.

### COMMENT

Huggins and Stevens were among the first to quantify the effect of androgen deprivation (AD) on human prostatic tissue. Using an ocular micrometer, these investigators found that prostatic epithelial cells shrink and acini decrease in size after castration therapy for symptomatic BPH. Herein we confirm and expand those observations from 60 years ago, using contemporary morphometric analysis to study finasteride treatment, which allows prostatic AD without the systemic side effects of castration. In Figure 5, landmark photomicrographs from the work of Huggins and Stevens are shown alongside comparison photomicrographs from the present work.

Using a variety of treatments and methods, others have subsequently attempted to quantify the...
effects of AD on human BPH histologic features.\textsuperscript{6,14–20} The prostatic effects of AD have also been demonstrated in rats\textsuperscript{21} and dogs\textsuperscript{22} treated with finasteride. Consistently, the major effect appears to be involution of the epithelium. However, using conventional histologic evaluation, the characteristics of epithelial atrophy are spotty and non-specific.\textsuperscript{23} Thus, contemporary morphometric analysis,\textsuperscript{24–26} used here to measure field change rather than individual cells, has provided a degree of quantitation not available with other methods of tissue analysis. The validity of using biopsy specimens to index prostatic tissue composition has been demonstrated previously.\textsuperscript{27}

The clinical, chemical, and volumetric observations made in the present group of long-term finasteride responders are consistent with previous reports.\textsuperscript{1–5} After 6 months of treatment, prostate volume decreases by approximately 18% to 20%; serum levels of DHT and PSA decrease by approximately 70% and 50%, respectively; voiding symptoms and flow rates are improved; and the new levels are maintained for years thereafter. Thus, the present group of patients, although highly selected (see Material and Methods, Patients), appears to be representative of the thousands of men studied in other clinical trials. Epithelial contraction, which we found to be statistically significant at 6 months,\textsuperscript{6} clearly continues further to the intermediate observation and still further to long-term follow-up (24 to 30 months). Thus, the tissue changes induced in the prostatic gland of finasteride responders are dramatic, enduring, and, in fact, progressive during several years of treatment. Progressive epithelial atrophy appears to be a basic mechanism of action of the drug and may explain the important benefits—55% to 57% risk reduction for operation or acute urinary retention—seen in men treated with long-term finasteride therapy.\textsuperscript{1}

Indirect evidence, based on serum PSA levels, indicates that finasteride may exert a preferential effect on the transition zone. The transition zone, not the whole prostate, appears to be the major source of PSA in the serum of men with BPH.\textsuperscript{28–31} In men with relatively high serum PSA levels, the response to finasteride is better than in those with low PSA levels.\textsuperscript{32} Furthermore, finasteride treatment results in a reduction of serum PSA levels out of proportion to the reduction of whole-prostate volume,\textsuperscript{8,13} suggesting the possibility of a preferential effect on the transition zone. In more direct studies, Tempany \textit{et al.},\textsuperscript{10} using MRI volumetrics,
and Tewari et al., using transrectal ultrasound, found that finasteride treatment seemed to shrink the transition zone more than the whole prostate. However, neither the study by Tempany et al. nor that by Tewari et al. is regarded as conclusive because of limited numbers of patients, only a single evaluation after treatment, large numbers of dropouts, and/or conflicting data.

The present work was designed in part to test the transition zone hypothesis of finasteride action and differs from previous reports in two important ways: (1) MRI and morphometric tissue analysis were used to assess the intact prostate before and serially during long-term finasteride treatment. (2) In addition to whole-gland studies, the peripheral and transition zones of the prostate were evaluated separately. Throughout the present study, the transition zone remained a constant percent of whole-prostate volume, and epithelial suppression was parallel in the transition and peripheral zones. Thus, in contrast to the placebo-controlled results found at 6 months, we found no evidence for a long-term preferential effect of finasteride on the inner prostate in treatment responders. In the present patients, prostate volume averaged 51 g at baseline. Results might be different in men with an even larger prostate gland and/or a larger transition zone because response to finasteride is directly correlated with prostate volume.

Substantial variability was seen in the epithelial response to finasteride, both between patients and temporally (Fig. 4). Although 17 of the 19 men ultimately exhibited a dramatic (50% or more) reduction in epithelial tissue, some men exhibited progressive suppression; others showed a delayed response; and in 2 patients, no change was seen. Whether use of a dual inhibitor of 5-alpha reductase (5-AR) would lead to faster or more uniform epithelial suppression than that seen with finasteride is unknown and would be of interest.

**CONCLUSIONS**

Taken altogether, the present data indicate that long-term finasteride treatment (24 to 30 months) results in marked suppression of the prostate epithelium. The effect is proportionately similar in both peripheral and transition zones for both volumetric and morphometric changes. Three different measures of prostate epithelium—percent tissue composition, volume, and stroma/epithelial ratio—all reflected an epithelial involution that continues to progress for many months after clinical changes have stabilized. Progressive contraction of

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**FIGURE 5.** (A and B) Photomicrographs of prostatic tissue before (A) and 86 days after (B) castration treatment for benign prostatic hyperplasia. (Original magnification ×235). Reprinted, with permission, from Huggins C, and Stevens RA: The effect of castration on benign hypertrophy of the prostate in man. J Urol 43: 705–714, 1940. (C and D) Color photomicrographs of prostatic tissue before (C) and after (D) 24 months of finasteride treatment. Enlarged from ×100. Marked flattening of epithelium and decrease in acinar diameter are noted after both treatments.
the prostate epithelium appears to constitute the underlying mechanism for the sustained action of finasteride.

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REFERENCES