**Effect of dutasteride on clinical progression of benign prostatic hyperplasia in asymptomatic men with enlarged prostate: a post hoc analysis of the REDUCE study**

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**Abstract**

**Objective** To assess the role of dutasteride in preventing clinical progression of benign prostatic hyperplasia in asymptomatic men with larger prostates.

**Design** Post hoc analysis of four year, double blind Reduction by Dutasteride of Prostate Cancer Events (REDUCE) study

**Participants** 1617 men randomised to dutasteride or placebo with a prostate size >40 mL and baseline International Prostate Symptom Score (IPSS) <8. Subjects who took medications for benign prostatic hyperplasia were excluded at study entry.

**Interventions** Placebo or dutasteride 0.5 mg daily.

**Main outcome measures** Comparison of risk of clinical progression of benign prostatic hyperplasia at four years (defined as a ≥4 point worsening on IPSS, acute urinary retention, urinary tract infection, or surgery related to benign prostatic hyperplasia).

**Results** 825 participants took placebo, 792 took dutasteride. A total of 464 (29%) experienced clinical progression benign prostatic hyperplasia, 297 (36%) taking placebo, 167 (21%) taking dutasteride (P<0.001). The relative risk reduction was 41% and the absolute risk reduction 15%, with a number needed to treat (NNT) of 7. Among men who had acute urinary retention and surgery related to benign prostatic hyperplasia, the absolute risk reduction for dutasteride was 6.0% and 3.8%, respectively. On multivariable regression analysis adjusting for covariates, dutasteride significantly reduced clinical progression of benign prostatic hyperplasia with an odds ratio of 0.47 (95% CI 0.37 to 0.59, P<0.001). Analysis of time to first event yielded a hazard ratio of 0.673 (P<0.001) for those taking dutasteride. Sexual adverse events were most common and similar to prior reports.

**Limitations** Further prospective studies may be warranted to demonstrate generalizability of these results.

**Conclusions** This study is the first to explore the benefit of treating asymptomatic or mildly symptomatic men with an enlarged prostate.

Dutasteride significantly decreased the incidence of benign prostatic hyperplasia clinical progression.

**Introduction**

Benign prostatic hyperplasia commonly causes lower urinary tract symptoms among men as they age. The treatment of this progressive condition is often adjusted for prostate size. Progression of prostatic hyperplasia is defined as an aggregate measure of worsening lower urinary tract symptoms, acute urinary retention, and need for prostate surgery. Pivotal phase III trials such as Medical Therapy of Prostatic Symptoms (MTOPS) and Combination of Avodart and tamsulosin (ComBAT) showed that combination medical therapy with a 5α reductase inhibitor and α blocker can halt the progression of benign prostatic hyperplasia among men with moderate to severe lower urinary tract symptoms, with the greatest benefit noted in men with enlarged prostates. An important aspect of both of these trials, as well as other studies showing the preventive benefits of 5α reductase inhibitors, is that they have excluded men with mild lower urinary tract symptoms at screening (MTOPS excluded International Prostate Severity Scale (IPSS) scores <8; ComBAT excluded IPSS scores <12).

Population data on the natural course of benign prostatic hyperplasia from Olmstead County indicate that average prostate size grows at an exponential rate of 1.6% per year. While the rate was not affected by age, it was higher in men with larger prostates. The unadjusted and adjusted relative risk of medical or surgical treatment for prostate volumes >30 mL were 4.2 (95% confidence interval 2.2 to 8.2) and 2.3 (1.1 to 4.7) respectively, while the unadjusted relative risk of acute urinary retention for prostate size >30 mL was 3.0 (1.0 to 9.0). These data were derived from both asymptomatic and symptomatic men. A prior meta-analysis suggested that, as well as predicting...
outcomes, a prostate size >40 mL predicts a significant difference in the magnitude of improvement with a 5α-reductase inhibitor compared with placebo. A prostate size >40 mL is also the volume used to define prostatic enlargement in the recent European Association of Urology guidelines on benign prostatic hyperplasia. The choice of treatment for symptomatic patients is often based on prostate size: α blocker monotherapy for patients with small prostates and combination therapy of 5α reductase inhibitors plus α blockers for patients with enlarged prostates. Obviously, asymptomatic men with small prostates do not require any treatment. However, it is not uncommon in clinical practice to encounter men with enlarged prostates but minimal lower urinary tract symptoms. These men are identified because of digital rectal examination or transrectal ultrasound findings for men with elevated prostate specific antigen levels. Because these patients have not been included in pivotal trials of treatment for benign prostatic hyperplasia, both the risk of clinical progression and the potential benefit of preventive treatment with 5α reductase inhibitors are unknown. The Reduction by Dutasteride of Prostate Cancer Events (REDUCE) cancer prevention trial is a large multinational prospective randomised study that measured prostate volume and urinary symptoms at baseline and included men with minimal or no lower urinary tract symptoms. We can therefore use this dataset to determine if there is any role for 5α reductase inhibitors in the treatment of asymptomatic men with enlarged prostates.

Methods

The REDUCE study was designed to assess the efficacy of dutasteride in decreasing the incidence of biopsy-detected prostate cancer over a four year period in men at increased risk of prostate cancer. Participants were randomised to placebo or dutasteride 0.5 mg daily. Inclusion and exclusion criteria are described in the original publication. This secondary analysis was prompted by the uniqueness of this dataset to answer the relatively common scenario: “How should I manage an asymptomatic man with an enlarged prostate?” This analysis aims to assess the benefit of dutasteride in men with mild or no urinary symptoms and enlarged prostates to prevent clinical progression of benign prostatic hyperplasia. The population assessed in this study includes participants randomised to both trial arms who had an International Prostate Severity Scale (IPSS) score <8 and a prostate size 40-80 mL on initial transrectal ultrasound (men with a prostate size of >80 mL were excluded from the original study). Subjects were excluded if they were taking any prostate related medications.

IPSS questionnaires were completed at baseline and at six-month intervals during the study. Urinary flow rates were measured at study entry (patients with a flow rate <5 ml/sec were excluded from the study) and in selected centres during follow-up. Clinical events—including acute urinary retention, prostate surgery related to benign prostatic hyperplasia, and urinary tract infection—were reported on case report forms every six months, and adverse events were assessed every three months during the study. All episodes of acute urinary retention were documented along with their cause (such as related to benign prostatic hyperplasia).

Similar to the major landmark trials and a Cochrane review, we used a composite index of clinical progression of benign prostatic hyperplasia as our primary end point. Participants who experienced urinary retention related to benign prostatic hyperplasia, surgery for benign prostatic hyperplasia, urinary tract infection, or symptom deterioration of IPSS score ≥4 points during the study were considered to have clinically progressed. A change in IPSS score of ≥4 has been shown to correspond to a change in patients’ global feeling of urination which is clinically meaningful and has been used in several large trials of benign prostatic hyperplasia. We also assessed adverse events between groups.

Statistical analyses

Our power calculations—based on 792 experimental subjects and 825 control subjects, four years of follow-up, and prior data estimating median time to progression of about 14.9 years with placebo—indicated we are able to detect true hazard ratios (relative risks) of placebo versus dutasteride groups of 0.859 or 1.168 with 80% power and α=0.05. Statistical analysis was performed using SAS version 9.2 (SAS Institute, Cary, NC, USA). Student’s t test and χ2 tests were used to compare groups. A multivariable logistic regression analysis assessed the effect of dutasteride on clinical progression of benign prostatic hyperplasia adjusting for age and baseline variables (IPSS, prostate volume, post-void residual urine volume, and peak urinary flow rate). Further, we performed an analysis of time to first event. Symptomatic progression was defined as the time of the first of two consecutive measures of IPSS ≥4 points from the baseline IPSS score (after randomisation). This was done to reduce variation based on only one questionnaire and demonstrate a durable change in symptoms. IPSS questionnaires were administered every six months during the four year study. Cox proportional hazard ratios were used to compare groups.

Results

Our study cohort consisted of 1617 men, 825 randomised to placebo, and 792 randomised to dutasteride. Table 1 shows baseline patient characteristics. The groups were balanced for age, prostate volume, and baseline flow parameters; there was no significant difference between groups.

A total of 464 patients (29%) experienced clinical progression of benign prostatic hyperplasia: 297 (36%) taking placebo, 167 (21%) taking dutasteride (P<0.001 for difference). The relative risk reduction was 41% and absolute risk reduction was 15%, with a number needed to treat of 6.7. Clinical events in each group are summarised in table 2. Seventy six patients (4.7%) had acute urinary retention: 63 taking placebo and 13 taking dutasteride (P<0.001). Of the 46 patients who had surgery related to benign prostatic hyperplasia, 39 were taking placebo and seven were taking dutasteride (P<0.001). One hundred and forty one patients had a urinary tract infection: 87 were taking placebo and 54 taking dutasteride (P<0.001). Figure 1 shows absolute and relative risk reductions for each event and the composite end point. On multivariable logistic regression analysis, dutasteride was the only factor that significantly reduced clinical progression of benign prostatic hyperplasia, with an odds ratio of 0.47 (95% confidence interval 0.37 to 0.59, P<0.001). Excluding patients with a urinary tract infection from the composite end point yielded similar results, with an odds ratio of 0.428 (0.336 to 0.545, P<0.001). The number of prostate cancers with a Gleason score ≥7 was not significantly different between groups (16 for placebo, 12 for dutasteride) in our cohort.

In our time-to-event analysis, 426 patients had two consecutive International Prostate Severity Scale (IPSS) scores ≥4 points greater than baseline score. Figure 2 shows the time to first event indicating progression of benign prostatic hyperplasia (acute urinary retention, surgery related to benign prostatic hyperplasia, or a change in IPSS score of ≥4). We calculated the risk of progression at 4 years and found that the probability of progression was reduced from 41% (placebo) to 25% (dutasteride), which is clinically meaningful and relevant.
hyperplasia, urinary tract infection, or symptomatic progression) for both groups. The hazard ratio for clinical progression among those randomised to dutasteride was 0.673 (P=0.001) relative to placebo. Other significant predictors of reaching the composite end point were baseline prostate volume (hazard ratio 1.007, P=0.008), baseline IPSS score (hazard ratio 0.958, P=0.046), and baseline urinary flow rate (hazard ratio 0.966 P<0.001).

The most common drug related side events were erectile dysfunction, at 5.1% and 9.0% (P=0.02) in the placebo and dutasteride arms respectively, and decreased or no libido, at 2.3% and 6.8% (P=0.001) in the two arms. Table 3 documents all adverse events that occurred with an incidence of >1% in our study population.

Discussion

Medical management goals of benign prostatic hyperplasia mainly consist of alleviating urinary symptoms, but other goals of treatment include improving bladder emptying, decreasing the effects of bladder outlet obstruction, and preventing future events such as haematuria, retention, and need for surgery. To our knowledge, this is the first study to explore the preventive benefit of treating asymptomatic or mildly symptomatic men at risk of progression of benign prostatic hyperplasia. Previous trials, for practical and ethical reasons, have focused on men with moderate to severe lower urinary tract symptoms. Analysis of the placebo arm of some of these trials has demonstrated that an enlarged prostate is a marker for patients at risk for benign prostatic hyperplasia progression. In these men with an enlarged prostate, 5α reductase inhibitors are recognised to be of greatest benefit. This study confirms the prophylactic benefit in asymptomatic or mildly symptomatic men, with the side effect profile as expected.

With men with no lower urinary tract symptoms enrolled, the REDUCE study data uniquely allowed us to demonstrate the benefit of a 5α reductase inhibitor in asymptomatic men at risk of benign prostatic hyperplasia progression due to prostatic enlargement. Careful monitoring and recording of events and causes of urinary retention, surgery, and urinary tract infections during this clinical trial provided accurate data for analysis. Although this is a post-hoc analysis, there are practical barriers to performing a prospective randomised trial in this population, including the need to perform a large number of transrectal ultrasound scans in asymptomatic men. The only other large dataset to our knowledge which contains data for similar patients does not contain routine information on prostate size. Table 4 presents the number needed to treat (NNT) to allow physicians to gauge the preventive benefit to this subset of patients over four years.

Study limitations

Although we used data from a well executed randomised clinical trial, there are some limitations to the present study. The population of men at increased risk of prostate cancer in the REDUCE study may not be representative of the broader population. By selecting men with an IPSS score of <8, there may be some regression to the mean, resulting in higher scores over time. None the less, this statistical effect is balanced by the fact that the changes on the IPSS score are considered more clinically important at lower baseline values. The a priori cut-off value for prostate size that we used (40 mL) is higher than the value of 30 mL often used to categorise large prostates. However, both values are arbitrary, and we felt a cut-off of >40 mL was clinically more representative of larger prostates that would benefit from preventive therapy.

Comparison with other studies

Other studies have noted that using 5α reductase inhibitors in men with enlarged prostates have a larger benefit in men who are more symptomatic. With the Proscar Long-Term Efficacy and Safety Study (PLESS) dataset, it was seen that the relative risk reduction in acute urinary retention and surgery related to benign prostatic hyperplasia was greater in men with higher baseline symptoms: the pLESS data showed a relative risk reduction at four years of 57% and 55% for acute urinary retention and surgery with finasteride. While both finasteride and dutasteride decrease prostate size, the higher relative risk reduction seen in our study may be related to the greater reduction in prostate volume seen with dutasteride. The hazard ratio for progression of benign prostatic hyperplasia in our study is similar to that reported in the secondary analysis of the Prostate Cancer Prevention Trial. However, that study included starting medications for benign prostatic hyperplasia as part of the composite end point, which may have comprised around 50% of the composite end points. The important end points for surgery related to benign prostatic hyperplasia and for urinary retention seem higher in our study, probably because our population is enriched with men with benign prostatic hyperplasia.

In the Medical Therapy of Prostatic Symptoms (MTOPS) study, the NNT with finasteride for men with a prostate size >40 mL or prostate specific antigen level >4 ng/mL was 7.2 to prevent one patient developing clinical progression of benign prostatic hyperplasia. While the NNT for our study is similar (6.7), the differences between the studies are noteworthy. Our study has a larger proportion of urinary tract infections and a lower proportion of acute urinary retention compared with the MTOPS study. This is important, as preventing symptoms with medical therapy has less merit than preventing specific complications. Also, it is important to note that incontinence or renal failure was not included as part of our composite end point. Despite, by definition, fewer symptomatic patients in our study cohort (mean IPSS score 4 v 17) and fewer end points, we did find a significant benefit.

Implications of study results

Treating asymptomatic patients is not an uncommon approach in medicine. Primary prevention with drugs is used to prevent cardiovascular complications and to reduce the risk of cardiac events. While not as lethal, the 10 year cumulative risk of acute urinary retention is estimated to be twice that of stroke or myocardial infarction. Further, research has found an episode of urinary retention to have a substantial impact on patients’ health related quality of life. Lower urinary tract symptoms affect almost three quarters men in their 60s, and direct costs in the US exceed $1bn a year, excluding outpatient prescription costs. For men found to have a large prostate gland during routine rectal examination or imaging, this information may be used to select those for whom preventive medication may improve their quality of life and also yield economic benefits. The trade-offs to the patient in this scenario are the side effects and cost of treatment. Side effects (table 3) include erectile dysfunction, decreased ejaculate volume, decreased libido, and gynaecomastia. Despite a greater patient preference against sexual side effects among those with mild symptoms, trade-off questionnaires suggest men prefer 5α reductase inhibitors compared with no treatment or α blockers. The concern over
risk of high grade prostate cancer remains an issue of controversy, though the incidence of high grade cancer seems to be lower relative to small prostate glands. The current drug costs of dutasteride per patient in the UK are £238 a year (£281, £365), with total direct costs estimated at £325 per year. A recent cost effectiveness study suggested that lifelong monotherapy with dutasteride in unselected patients has a cost per QALY gained of £274. Our data suggest that its preventive use in asymptomatic men with enlarged prostates would have a lower cost per QALY, given the similar relative risk reduction used in this cost analysis and the significantly higher incidence of complications related to benign prostatic hyperplasia in our study population. By comparison, the estimated cost of transurethral resection of the prostate (TURP) in the NHS was £2080, and the estimated cost of one admission for acute urinary retention was £1040. However, cost effectiveness research to assess the use of dutasteride as a preventive agent in this population is unlikely to occur given that its patent expires in 2013. While some patients will not generally be receptive to preventive drug treatment, the magnitude of risk reduction seen in our study warrants further study of patient preferences for choosing optimal management. Certainly, the potential relative harms to each patient need to be discussed and weighed against the expected potential benefits.

Conclusions

This study is the first to explore the benefit of treating asymptomatic or mildly symptomatic men with enlarged prostates. In this cohort, dutasteride significantly decreased the incidence of clinical progression of benign prostatic hyperplasia over four years, with a relative risk reduction of over 50% and an acceptable side effect profile. This post-hoc analysis generates questions on whether 5α-reductase inhibitors should be discussed as preventive treatment for patients with an enlarged prostate.

We thank Lauren Marmor and colleagues at GlaxoSmithKline for their permission to use the data from the Reduction by Dutasteride of Prostate Cancer Events (REDUCE) study (Clinical Trials No NCT00056407). We also thank Karen Chadwick at University Health Network, University of Toronto, in facilitating this. Contributors: PT, DM, and NF initiated the study and designed the statistical analysis plan. PT and DM analysed the data and drafted the paper. GK, AF, AZ, and NF gave critical input on the analysis and revised the paper. PT and NF are the guarantors for the study.

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What is already known on this topic

Dutasteride, a 5α reductase inhibitor, is an established treatment for lower urinary tract symptoms due to benign prostatic hyperplasia, particularly among men with an enlarged prostate. Men with an enlarged prostate are at risk of urinary symptoms and complications such as acute urinary retention, urinary tract infections, and need for surgery. However, none of the major trials of 5α reductase inhibitors have included asymptomatic or minimally symptomatic men.

What this study adds

This post hoc analysis of trial data uniquely estimates the benefit of dutasteride among men with no or minimal symptoms at risk of complications from benign prostatic hyperplasia due to prostate enlargement.

Dutasteride significantly decreased the incidence of clinical progression of benign prostatic hyperplasia over four years, with a relative risk reduction of over 50% and an acceptable side effect profile.

Tables

Table 1  Baseline demographics and characteristics of 1617 asymptomatic men with enlarged prostate glands who were randomised to treatment with dutasteride or placebo. Values are medians (interquartile ranges) unless stated otherwise

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Placebo (n=825)</th>
<th>Dutasteride (n=792)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>64 (59-68)</td>
<td>63 (58-67)</td>
</tr>
<tr>
<td>No (%) of white ethnicity</td>
<td>76 (92)</td>
<td>73 (83)</td>
</tr>
<tr>
<td>Prostate volume (cm³)</td>
<td>51.9 (45.2-60.3)</td>
<td>52.0 (45.5-61.9)</td>
</tr>
<tr>
<td>Serum PSA concentration (ng/mL)</td>
<td>6.0 (4.6-7.5)</td>
<td>5.9 (4.5-7.5)</td>
</tr>
<tr>
<td>Maximum urinary flow rate (mL/sec)</td>
<td>13 (9.7-18)</td>
<td>13 (10-17.6)</td>
</tr>
<tr>
<td>Post-void residual urine volume (mL)</td>
<td>28 (2-70)</td>
<td>30 (5-73)</td>
</tr>
<tr>
<td>No (%) who were sexually active</td>
<td>702 (85%)</td>
<td>663 (83%)</td>
</tr>
<tr>
<td>IPSS score</td>
<td>4.0 (3-6)</td>
<td>4 (3-6)</td>
</tr>
</tbody>
</table>

PSA=prostate specific antigen. IPSS=International Prostate Symptom Score.
<table>
<thead>
<tr>
<th>Clinical event</th>
<th>No (%) of patients</th>
<th>Placebo (n=825)</th>
<th>Dutasteride (n=792)</th>
<th>Odds ratio (95% CI)*</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute urinary retention</td>
<td>76 (4.7)</td>
<td>63 (7.6)</td>
<td>13 (1.6)</td>
<td>0.20 (0.11 to 0.37)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>BPH-related surgery</td>
<td>46 (2.8)</td>
<td>39 (4.7)</td>
<td>7 (0.9)</td>
<td>0.18 (0.08 to 0.40)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td>141 (9.7)</td>
<td>87 (10.5)</td>
<td>54 (6.8)</td>
<td>0.62 (0.43 to 0.88)</td>
<td>0.008</td>
</tr>
<tr>
<td>IPSS score increase of ≥4 points</td>
<td>306 (18.9)</td>
<td>192 (23.3)</td>
<td>114 (14.4)</td>
<td>0.55 (0.40 to 0.72)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

BPH=benign prostatic hyperplasia. IPSS=International Prostate Symptom Score

*Odds ratio from logistic regression analysis.
Table 3  Incidence of drug related adverse events among 1617 asymptomatic men with enlarged prostate glands who were randomised to treatment with dutasteride or placebo

<table>
<thead>
<tr>
<th>Drug related adverse event*</th>
<th>No (%) of patients</th>
<th>Absolute risk (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Placebo (n=825)</td>
<td>Dutasteride (n=792)</td>
<td></td>
</tr>
<tr>
<td>Any</td>
<td>134 (16.2)</td>
<td>276 (34.8)</td>
<td>18.6 (14.4 to 22.7)</td>
</tr>
<tr>
<td>Decreased libido</td>
<td>10 (1.2)</td>
<td>35 (4.4)</td>
<td>3.2 (1.6 to 9.8)</td>
</tr>
<tr>
<td>Loss of libido</td>
<td>9 (1.1)</td>
<td>19 (2.4)</td>
<td>1.3 (0.03 to 2.6)</td>
</tr>
<tr>
<td>Erectile dysfunction</td>
<td>42 (5.1)</td>
<td>71 (9.0)</td>
<td>3.9 (1.4 to 6.4)</td>
</tr>
<tr>
<td>Gynaecomastia</td>
<td>6 (0.7)</td>
<td>19 (2.4)</td>
<td>1.7 (0.5 to 2.9)</td>
</tr>
<tr>
<td>Decreased semen volume</td>
<td>1 (0.1)</td>
<td>12 (1.5)</td>
<td>1.4 (0.5 to 2.3)</td>
</tr>
</tbody>
</table>

*All adverse events reported to occur in >1% of subjects in either group.
<table>
<thead>
<tr>
<th>Event to be prevented</th>
<th>NNT (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composite end point*</td>
<td>6 (5 to 9)</td>
</tr>
<tr>
<td>IPSS score increase of ≥4 points (urination globally “worse”&quot;&quot;)</td>
<td>11 (8 to 20)</td>
</tr>
<tr>
<td>Acute urinary retention</td>
<td>16 (12 to 25)</td>
</tr>
<tr>
<td>BPH-related surgery</td>
<td>26 (18 to 44)</td>
</tr>
<tr>
<td>Acute urinary retention or BPH-related surgery</td>
<td>13 (10 to 18)</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td>26 (15 to 102)</td>
</tr>
</tbody>
</table>

BPH=benign prostatic hyperplasia. IPSS=International Prostate Symptom Score

*Composite end point=acute urinary retention, IPSS score increase of ≥4 points, BPH-related surgery, or urinary tract infection.
Figures

**Fig 1** Absolute rates and relative risk reduction for acute urinary retention, surgery related to benign prostatic hyperplasia, and clinical progression of benign prostatic hyperplasia among 1617 asymptomatic men with enlarged prostate glands who were randomised to treatment with dutasteride or placebo.

**Fig 2** Time to first event indicating progression of benign prostatic hyperplasia among 1617 asymptomatic men with enlarged prostate glands who were randomised to treatment with dutasteride or placebo.