Economic Analysis: Randomized, Placebo-Controlled Clinical Trial of Dutasteride in Men at High Risk for Prostate Cancer

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Abstract

Objective: Given the economic burden of prostate cancer (PCa), a PCa risk-reduction medication would be desirable. A within-trial economic analysis of the Reduction by Dutasteride of Prostate Cancer Events (REDUCE) study was performed.

Methods: REDUCE, a 4-year, randomized, double-blind, placebo-controlled, parallel-group clinical trial, compared efficacy and safety of dutasteride 0.5 mg daily and placebo to reduce the risk of PCa in men at increased risk. Resource use was prospectively collected; costs from standard costing sources were applied. Utilities were obtained from published literature. Relative risks and Wilcoxon rank sums were used to examine differences between treatments.

Results: Placebo patients were at significantly higher risk ($P < 0.05$) for concomitant medication use; and health care visits related to surgical procedures, unscheduled biopsies, acute urinary retention, urinary tract infections, or macroscopic hematuria. Total costs were significantly lower ($P < 0.001$) in dutasteride patients ($\$1,300; 95\%$ confidence interval: $\$806, \$1,795$). Incremental cost per quality-adjusted life-year (QALY) was $\$26,516$; cost per PCa case avoided was $\$19$.

Conclusions: During the 4-year trial period, men at increased risk for PCa receiving dutasteride incurred fewer health care costs than men receiving placebo, which helped offset dutasteride costs. Dutasteride was good value for money.

Keywords: Benign prostatic hyperplasia; Prostate cancer; Resource use; Cost-effectiveness analysis; Economics

Abbreviations: 5ARI: 5-alpha reductase inhibitors; AUR: Acute urinary retention; BPH: Benign prostatic hyperplasia; CI: Confidence interval; ICER: Incremental cost-effectiveness ratios; PCa: Prostate cancer; PSA: Prostate-specific antigen; QALY: Quality-adjusted life-year; REDUCE: Reduction by Dutasteride of Prostate Cancer Events (clinical trial); US: United States; UTI: Urinary tract infection

Introduction

Prostate cancer (PCa) is one of the most common cancers in American men and the second leading cause of cancer death [1]. An estimated 1 in 6 men will be diagnosed with PCa during their lifetime, and 1 in 36 men will die of PCa [1]. Medical expenditures for PCa were estimated to be $11.85 billion in 2010 and projected to reach $16.34 billion in 2020 (in 2010 dollars) [2].

Given this burden, medication to reduce the risk of PCa could provide economic and quality-of-life benefits. In recent years, the use of 5-alpha reductase inhibitors (5ARIs) has been studied for PCa risk reduction. Specifically, finasteride has been shown to reduce the incidence of PCa by 24.8% (95% confidence interval [CI]: 18.6%-30.6%; $P < 0.001$) in men with serum prostate-specific antigen (PSA) < 3.0 ng/mL in the Prostate Cancer Prevention Trial [3]. In a recent clinical trial, the Reduction by Dutasteride of Prostate Cancer Events (REDUCE), dutasteride (0.5 mg/day) was shown to reduce the risk of PCa by 22.8% (95% CI: 15.2%-29.8%; $P < 0.001$) compared with placebo in men with a negative biopsy and PSA > 2.5 ng/mL at baseline [4].

Previous analyses have examined the potential cost-effectiveness of a chemoprevention agent for prostate cancer. Because of incomplete information at the time, these analyses used decision analytic techniques, such as Markov models, to compile data from multiple sources and extrapolate the potential impact on costs and outcomes that might be seen over a man’s remaining lifetime [5-11]. Within the REDUCE study, data on resource use were collected along side of the clinical data. This study was an economic analysis of resource use data from the REDUCE clinical trial, comparing dutasteride with placebo in men at increased risk for PCa in the first 4 years of drug administration.

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Materials and Methods

REDUCE Study Design

The REDUCE clinical trial was a 4-year, phase 3, international, multicenter, randomized, double-blind, placebo-controlled, parallel group study to evaluate the safety and efficacy of oral, once-daily 0.5-mg dutasteride in reducing the risk of biopsy-detecable PCa in men aged 50 to 60 years with PSA of 2.5 to 10 ng/mL or in men aged 60 years or older with PSA of 3.0 to 10 ng/mL, and with a single negative prostate biopsy in the prior 6 months and prostate volume of 80 cc or less [4]. The trial randomized 8,231 men to receive dutasteride or placebo. Patients underwent transrectal ultrasound-guided biopsies at 2 and 4 years. The primary endpoint was biopsy-detecable PCa. The REDUCE clinical trial was approved by the institutional review boards at each research site, and all participants provided written informed consent [4]. This analysis was performed using de-identified data (i.e., all researchers were blinded to any identifying information).

Resource Use

Resource use data were collected during the trial within case report forms. For our analysis, we included only resource use related to PCa, benign prostatic hyperplasia (BPH), or other conditions believed to be affected by the use of a 5ARI. Resource use included concomitant medications and health care visits associated with surgical and nonsurgical procedures, unscheduled biopsies, urinary tract infections (UTIs), acute urinary retention (AUR), and macroscopic hematuria and hematospermia episodes unrelated to trial-specified biopsies.

Concomitant medications included those related to the conditions previously mentioned—chemotherapies, hormone therapies, antibiotic therapy for treating UTI, and alpha-blocker use for treating BPH—as well as concomitant medications used to treat significant adverse events due to dutasteride use (e.g., impotence, decreased libido, and ejaculation disorders). A clinician was consulted on concomitant medications included in our analysis. All procedures and concomitant medication classes considered are presented in the online appendix.

Costs

Direct medical costs were obtained from the payer perspective. All costs were reported in 2010 United States (US) dollars. Details of how per-patient direct costs were calculated are explained in the online appendix.

Inpatient Costs

Inpatient costs per day for surgical and nonsurgical procedures, UTI, AUR, and macroscopic hematuria and hematospermia were obtained from the Healthcare Cost and Utilization Project Nationwide Inpatient Sample [12]. Charges obtained from this database were converted to costs using published costs-to-charge ratios [13].

Outpatient Costs

Outpatient visits, outpatient procedures, unscheduled biopsies, and drug administrations were assigned a Current Procedural Terminology code from the Current Procedural Terminology and Healthcare Common Procedure Coding System [14]. All codes were verified by a clinician. Costs were obtained from the Resource-Based Relative Value Scale [15].

Concomitant Medications Costs

Wholesale drug acquisition costs were obtained from the 2010 Red Book for Windows [16]. Generic drugs and/or lowest-cost drugs were used when branded medications were not specified. For hormone therapies not available in the US, we used the average US costs of hormone therapies within the REDUCE trial.

The objective of the resource use and cost analysis was to understand the impact on other medical costs when using dutasteride. As a result, the cost of dutasteride was excluded from this analysis. However, the cost of dutasteride was included in the cost-effectiveness analysis. Dutasteride’s wholesale acquisition cost for the cost-effectiveness analysis was $3.62 per day [16].

Utility and Quality of Life

To calculate quality-adjusted life-years (QALYs) and perform cost-utility analyses, we obtained utility weights from the published literature [17-19] and applied these to the men within the REDUCE trial according to health status. (Data required to calculate utilities were not collected within REDUCE.) Specifically, over the course of a man’s involvement in the trial, he was designated as being in one of the following health states: healthy with no PCa or BPH, BPH but no PCa, high-grade or low-grade PCa, or death. A man’s QALYs were decremented if adverse events, such as ejaculatory dysfunction and impotence, UTIs, AURs, and BPH-related surgeries, occurred. In particular, men on dutasteride incurred a utility decrement due to the sexual dysfunction AEs. Since utilities for high-grade and low-grade tumors consider AEs that may occur due to PCa treatment, utility decrements for AEs due to PCa treatment were not considered. Men with BPH who were taking dutasteride or an alpha-blocker were assumed to incur an improvement in symptoms. The utilities and utility decrements obtained from the published literature are presented in Table 1.

Statistical Analysis

The analysis population within this study, called the biopsy population, consisted of patients in the efficacy population (as defined in [4]) who had at least one post-baseline biopsy. A resource use event was included in the economic analysis if it happened during the trial period. Inclusion criteria and date imputation rules are presented in the online appendix.

Resource use was analyzed as categorical variables (i.e., whether a patient used a specific resource or not). The number and proportion of patients using the resource and 95% CIs were summarized for each treatment group. Statistical comparisons were made between treatment groups (i.e., placebo vs. dutasteride) using 95% CIs of relative risks. A resource use with a lower-bound 95% CI of relative risk greater than 1 means that the risk for having a resource use is significantly higher for patients in the placebo group than for patients in the dutasteride group.

Costs were analyzed as continuous variables; we estimated means and standard deviations among each resource use category for each treatment group since arithmetic mean cost is more appropriate [20]. We also calculated mean differences (i.e., placebo vs. dutasteride) and 95% CIs. However, because resource use distribution and therefore cost distribution can be heavily skewed, we preferred to use a nonparametric test, Wilcoxon rank sum, to test the null hypothesis that the cost distributions in the two treatment groups were the same. The cost distribution is significantly lower (or higher) in the dutasteride group than in the placebo group if the observed rank sum in the dutasteride group is less (or greater) than the expected rank sum. A test result was declared statistically significant if the P value was less than 0.05. No adjustments were made for multiple comparisons since this correction
would increase the frequency of type II errors (i.e., false negatives) [21].
Analyses were performed using SAS, version 9.1 (SAS Institute, Inc.,
Cary, North Carolina).

Cost-effectiveness Ratios

Cost-effectiveness and cost-utility analyses were performed. The
incremental cost-effectiveness ratios (ICERs) of incremental cost per
PCa case avoided [5] and incremental cost per QALY gained were
calculated. ICERs were calculated as the ratio of the difference in
average total costs between the two treatment groups and the difference
in average effectiveness between the two treatment groups. Dutasteride
was considered cost-effective if the incremental cost per QALY gained
was $50 000 or less [22-24].

Sensitivity Analyses

We performed sensitivity analyses by running the resource use and
cost analyses in other population subgroups defined by geographic
region and baseline PSA. These analyses were performed to understand
the effect that methodological assumptions had on the results. All
populations are described in Table 2, together with the respective
patient numbers, by treatment group.

To consider uncertainty due to sampling variation surrounding the
estimate of the ICERs, we ran 1 000 bootstrap replications to calculate
the median and bootstrapped 95% percentiles of the ICERs [25]. A
scatter plot and cost-effectiveness analysis curve were constructed [26].

Results

The biopsy population consisted of 3 305 patients in the dutasteride
group and 3 424 patients in the placebo group (Table 2). Date
imputation across treatment groups was similar, where 25% of patients
had imputed dates for concomitant medications and 4% of patients
had imputed dates for nonconcomitant medication resources. PCa
occurred in 599 patients in the dutasteride group and 782 patients in
the placebo group.

Resource Use

Table 3 presents the results of the resource use analysis. Overall, the
largest resource use categories were concomitant medication use (27%)
and unscheduled biopsies (12%).
Compared with patients in the dutasteride group, patients in the placebo group were at significantly higher risk for incurring use of concomitant medication (including alpha-blocker use); and health care visits for treating surgical procedures, unscheduled biopsies, UTIs, AURs, or macroscopic hematuria (Table 3).

Costs
Concomitant medication (40%) and surgical (42%) costs made up the majority of the total costs incurred (Figure 1). Surgical costs were the major contributor to total costs, although they occurred in only 4% of patients during the trial period. These proportions were similar between treatment groups.

Total costs, excluding the study drug cost, were significantly lower in the dutasteride group (mean difference of $1,300; 95% CI: $806, $1,795) (Table 4). Patients in the dutasteride group incurred significantly lower costs for concomitant medication use (including alpha-blocker use); and health care visits for treating surgical procedures, unscheduled biopsies, UTIs, AURs, or macroscopic hematuria. According to the Wilcoxon rank sum test, patients in the dutasteride group incurred significantly lower costs for concomitant medications and macroscopic hematuria despite mean total costs being higher for this group than for the placebo group. This counterintuitive result is possible due to skewness of cost distribution.

Cost-effectiveness
Over the 4-year period of the trial, the mean acquisition cost of dutasteride among men in the dutasteride group was $4,733. Thus, the acquisition cost of dutasteride was not fully offset by reductions in other medical costs. However, men in the dutasteride group did accrue 0.13 more QALYs than men in the placebo group (3.26 vs. 3.13). As a result, the use of dutasteride to reduce the risk of PCa was cost-effective, with an incremental cost per QALY of $2,516 (Table 5). The incremental cost per PCa avoided was calculated as $19 (Table 5).

Sensitivity Analysis
Differences in resource use within the subgroup populations were found to be similar to the differences observed for the biopsy population. The difference in mean total costs was found to be

Table 3: Number and Proportion of Resource Events or Procedures by Treatment Group.

<table>
<thead>
<tr>
<th>Resource</th>
<th>Dutasteride (N = 3,305)</th>
<th>Placebo (N = 3,424)</th>
<th>Relative Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Concomitant Medications</strong></td>
<td>841 (25, 27)</td>
<td>994 (29, 31)</td>
<td>1.14 (1.05, 1.23)</td>
</tr>
<tr>
<td><strong>Alpha-blockers</strong></td>
<td>713 (22, 23)</td>
<td>862 (25, 27)</td>
<td>1.17 (1.07, 1.27)</td>
</tr>
<tr>
<td><strong>Health Care Visits</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surgical procedures</td>
<td>60 (2, 2)</td>
<td>188 (5, 6)</td>
<td>3.02 (2.27, 4.03)</td>
</tr>
<tr>
<td>Nonsurgical procedures</td>
<td>5 (0, 0)</td>
<td>12 (0, 1)</td>
<td>2.32 (0.82, 6.57)</td>
</tr>
<tr>
<td>Unscheduled biopsies</td>
<td>322 (10, 11)</td>
<td>470 (14, 15)</td>
<td>1.41 (1.23, 1.61)</td>
</tr>
<tr>
<td>UTI</td>
<td>186 (6, 6)</td>
<td>299 (9, 10)</td>
<td>1.55 (1.30, 1.85)</td>
</tr>
<tr>
<td>AUR</td>
<td>53 (2, 2)</td>
<td>230 (7, 8)</td>
<td>4.19 (3.12, 5.62)</td>
</tr>
<tr>
<td>Macroscopic hematuria</td>
<td>114 (3, 4)</td>
<td>171 (5, 6)</td>
<td>1.45 (1.15, 1.83)</td>
</tr>
<tr>
<td>Macroscopic hematospermia</td>
<td>43 (1, 2)</td>
<td>35 (1, 1)</td>
<td>0.79 (0.50, 1.22)</td>
</tr>
</tbody>
</table>

AUR = acute urinary retention; CI = confidence interval; UTI = urinary tract infection.
*Table presents the proportions of patients within each treatment group who experience at least one episode of each resource.

Figure 1: Breakdown of Total Costs.
The incremental cost per QALY among the different population subgroups was consistent with a low of $22 391 in the baseline PSA < 4.9 ng/mL population ($887), the North American ($41 911 to $330.94) and the North American ($24 914.72 to $42 019.95) populations. We observed in the trial, we found that, compared with patients in the dutasteride group, patients in the placebo group were at significantly higher risk for incurring resource use pertaining to concomitant medications (including alpha-blockers) and health care visits for surgical procedures, unscheduled biopsies, UTI, AUR, or macroscopic hematuria. Total costs, excluding study drug cost, were significantly higher in the placebo group than in the dutasteride group of the biopsy population and in all examined population subgroups. We observed that the use of dutasteride to reduce the risk of PCa was cost-effective compared with placebo. In addition, the cost to avoid a case of PCa was greater in the baseline PSA ≥ 6.8 ng/mL population ($1 953) than the difference seen in the biopsy population (Figure 2), whereas lower differences were found in the baseline PSA < 4.9 ng/mL ($887), the baseline PSA ≥ 4.9 and < 6.8 ng/mL ($990), and the North American ($118) populations (Figure 2). The lower difference in mean total cost between the placebo group and the dutasteride group in the North American population was due to the occurrence of a lower proportion of hospitalizations and shorter lengths of stay for surgical procedures. Using the Wilcoxon rank sum test, the total costs for the dutasteride group remained significantly lower than the total costs for the placebo group in all sensitivity analysis population subgroups.

The incremental cost per QALY among the different population subgroups was consistent with a low of $22 391 in the baseline PSA ≥ 6.8 ng/mL population and a high of $51 780 in the North American population. Similar results occurred for the incremental cost per PCa case avoided: a low of $47 in the baseline PSA ≥ 6.8 ng/mL population and a high of $119 in the North American population. (Note the denominators for the population subgroups were summed to comprise the denominator for the biopsy population. Thus, the incremental costs per PCa case avoided are larger for population subgroups than for the biopsy population.) Table 5 shows the ICERs.

In examining uncertainty due to sampling variation, the baseline and bootstrapped results were similar. The scatter plot and cost-effectiveness acceptability curve for the biopsy population are presented in Figure 3; dutasteride was cost-effective 100% of the time since all bootstrapped ICERs fell to the east and south of the $50 000 threshold line. Table 5 displays bootstrapped median ICERs and 95% percentiles for each population.

### Discussion

We performed an economic analysis of dutasteride compared with placebo in men at increased risk for PCa within the biopsy population of the REDUCE clinical trial. Specifically, we analyzed resource utilization as recorded within the REDUCE trial and performed cost and cost-effectiveness analyses. As a result of the resource use observed in the trial, we found that, compared with patients in the dutasteride group, patients in the placebo group were at significantly higher risk for incurring resource use pertaining to concomitant medications (including alpha-blockers) and health care visits for surgical procedures, unscheduled biopsies, UTI, AUR, or macroscopic hematuria. Total costs, excluding study drug cost, were significantly higher in the placebo group than in the dutasteride group of the biopsy population and in all examined population subgroups. We observed that the use of dutasteride to reduce the risk of PCa was cost-effective compared with placebo. In addition, the cost to avoid a case of PCa was minimal. These results were observed over the 4-year trial period. As a result, they cannot be generalized beyond this period of time.
This analysis differs from previous analyses in that all previous analyses examined the impact of 5ARIs using decision analytic modeling techniques. Those techniques were used because information on the direct impact of 5ARI use on resource use was not available. In other words, decision analytic techniques were used to extrapolate the impact of 5ARI use through the compilation of data from a variety of sources because of the absence of complete data. To fully understand the impact that chemoprevention with a 5ARI would have on resource use, we would ideally prefer to collect these data as part of a real-world, prospectively designed study. However, this type of study is costly in terms of time and money. Since resource use was collected as part of the large clinical trial in this study, we felt it was important to examine the impact of dutasteride on unscheduled resource use as seen within the clinical trial and on potential costs associated with the use of these resources. Although these data were collected within a controlled setting, analyzing the unplanned resource use provides us with some sense of potential impact on these data.

In the biopsy population, the use of dutasteride was shown to reduce the proportion of patients on alpha-blocker therapy by approximately 14%. However, we observed only small decreases in alpha-blocker costs in patients on dutasteride because patients in the dutasteride group taking alpha-blockers had a greater number of days on alpha-blocker therapy (mean = 1 078 days) than patients in the placebo group who were taking alpha-blockers (mean = 911 days). Changing the imputation method may change our results to be more in favor of dutasteride.
This study has several limitations, particularly when making comparisons between treatments. First, this is an analysis of resource use data collected as part of a clinical trial. Patients within clinical trials are in a controlled setting and are monitored closely. We endeavored to exclude resource use that was trial driven, but we acknowledge that these data may not fully reflect resource use and costs as seen in the real world. For example, it is possible that unscheduled biopsies may be underestimated within the trial because scheduled biopsies occurred within the trial. Thus, investigators within the trial might have delayed performing biopsies they might have performed in a real-world clinical setting because they knew the patient was scheduled for a biopsy in a few months. In addition, resource use after completion of the trial was not considered.

Another limitation of this analysis is that dutasteride is currently indicated to treat BPH. Therefore, it might be argued that most of the benefits accrued for patients in the cost-effectiveness analysis are due to treating BPH. As a result, a limitation of the analysis to examine the cost-effectiveness of dutasteride as a chemoprevention agent is that it includes the impact that dutasteride has on preventing BPH-related events such as AURs and BPH-related surgeries. In other words, these analyses should consider only the impact on PCA-related costs. However, the incremental cost per PCA case avoided is well below the cost of treating PCA. In addition, it is important to realize that men at increased risk for PCA most likely will have BPH symptoms or be at increased risk for BPH. Thus, it is relevant to look at gains from both within the analysis.

In men who went on to have PCA, it is possible that complete resources to treat their PCAs were not captured. In fact, 95% of men with PCAs had less than 1 year of follow-up data within the trial setting. Thus, the full impact of potentially offsetting PCA costs was not captured within this analysis. We performed a subanalysis in which patients on each treatment who had less than 1 year follow-up after PCA diagnosis had their costs replaced with the average costs for patients on the same treatment who had 1 or more years of follow-up after diagnosis. We found an additional improvement in the cost-effectiveness, with an incremental cost per QALY of $16 342 and an incremental cost per PCA case of $7. Thus, the baseline results of this analysis may be considered conservative compared with what we might expect to see in actual clinical practice.

This was an analysis of pooled, multicountry data. We made no adjustments for differences in resource use patterns between countries. In addition, we applied US costs to the globally collected data. In evaluating the cost and resource use differences from a US perspective, we must use caution. However, with these limitations in mind, we performed sensitivity analyses including a sensitivity analysis in which the resource use and cost were examined in a population of patients in North America. Within this subanalysis specific to North America, a higher incremental cost per QALY was observed for the North American region than for the biopsy population. Overall, we observed a lower proportion of hospitalizations and shorter lengths of stay for surgical procedures in the North American Region than in the biopsy population. Thus, the use of dutasteride as a chemoprevention agent was not as cost-effective in the North American population as in the biopsy population.

In this analysis, we did not cost and analyze all concomitant medications that were reported in the REDUCE trial. Instead, we selected specific classes of concomitant medications that we believe would impact the overall costs of health care if dutasteride were taken to reduce the risk of PCA. This pragmatic approach was chosen to better reflect the true impact of treatments on relevant resource utilization. As a validation, medication classes that were included in the analysis were verified by a urologist as being most likely to be affected by the use of a 5ARI to reduce the risk of PCA.

Our analysis was intended to be exploratory and descriptive in nature. Because cost distributions are skewed, caution is advised when interpreting the parametrically estimated CIs, and they should be viewed as purely descriptive. Generally, it is safer to trust nonparametric conclusions because the inconsistency between parametric and nonparametric results suggests nonnormal distributions. However, one should be aware that the Wilcoxon test compares the cost distributions and not the arithmetic means. Given the exploratory nature of the
analysis, we did not make multiple comparison adjustments; thus, $P$
values should be interpreted with caution. Future work to provide a
more comprehensive picture of the impact that the use of dutasteride
would have on health care costs and resource use could include the use
of a study of routine clinical practice data in this area.

Overall, in the absence of a prospective study to examine the
complete economic impact of the use of a SARI as a prostate
chemoprevention agent in men at increased risk for prostate cancer, we
performed an analysis of the unscheduled resource use collected within
the REDUCE clinical trial. Although increased use of a dutasteride
would occur within the population, decreased use of concomitant
medications and various healthcare procedures may be observed. As
a result, a reduction in costs of non-SARI medications and medical
resources may result. In addition, although some increases in the cost-
effectiveness were seen when analyzing the sensitivity populations, we
may observe that the use of dutasteride may be good value for money
across all populations analyzed.

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Conflicts of Interest

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that has received research funding for this and other studies from GlaxoSmithKline and
pharmaceutical companies that market drugs for use as a chemoprevention
for prostate cancer and other medical conditions. Dr. Black is an employee of
GlaxoSmithKline, a pharmaceutical company that manufactures dutasteride.
Dr. Gerald Andriole is an employee at Washington University School of Medicine
that has received research funding for this and other studies from GlaxoSmithKline
employees of RTI Health Solutions, an independent contract research organization


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