Effect of Soy Protein Isolate Supplementation on Biochemical Recurrence of Prostate Cancer After Radical Prostatectomy
A Randomized Trial

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IMPORTANCE Soy consumption has been suggested to reduce risk or recurrence of prostate cancer, but this has not been tested in a randomized trial with prostate cancer as the end point.

OBJECTIVE To determine whether daily consumption of a soy protein isolate supplement for 2 years reduces the rate of biochemical recurrence of prostate cancer after radical prostatectomy or delays such recurrence.

DESIGN, SETTING, AND PARTICIPANTS Randomized, double-blind trial conducted from July 1997 to May 2010 at 7 US centers comparing daily consumption of a soy protein supplement vs placebo in 177 men at high risk of recurrence after radical prostatectomy for prostate cancer. Supplement intervention was started within 4 months after surgery and continued for up to 2 years, with prostate-specific antigen (PSA) measurements made at 2-month intervals in the first year and every 3 months thereafter.

INTERVENTION Participants were randomized to receive a daily serving of a beverage powder containing 20 g of protein in the form of either soy protein isolate (n=87) or, as placebo, calcium caseinate (n=90).

MAIN OUTCOMES AND MEASURES Biochemical recurrence rate of prostate cancer (defined as development of a PSA level of \( \geq 0.07 \) ng/mL) over the first 2 years following randomization and time to recurrence.

RESULTS The trial was stopped early for lack of treatment effects at a planned interim analysis with 81 evaluable participants in the intervention group and 78 in the placebo group. Overall, 28.3% of participants developed biochemical recurrence within 2 years of entering the trial (close to the a priori predicted recurrence rate of 30%). Among these, 22 (27.2%) occurred in the intervention group and 23 (29.5%) in the placebo group. The resulting hazard ratio for active treatment was 0.96 (95% CI, 0.53-1.72; log-rank \( P = .89 \)). Adherence was greater than 90% and there were no apparent adverse events related to supplementation.

CONCLUSION AND RELEVANCE Daily consumption of a beverage powder supplement containing soy protein isolate for 2 years following radical prostatectomy did not reduce biochemical recurrence of prostate cancer in men at high risk of PSA failure.

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Prostate cancer is the most frequently diagnosed malignancy and the second most frequent cause of male cancer death in the United States and other Western countries but is far less frequent in Asian countries. Prostate cancer risk has been inversely associated with intake of soy and soy foods in observational studies, which may explain this geographic variation because soy consumption is low in the United States and high in Asian countries. Although it has been repeatedly proposed that soy may prevent prostate cancer development, this hypothesis has not been tested in randomized studies with cancer as the end point. A substantive fraction (48%-55%) of men diagnosed as having prostate cancer use dietary supplements including soy products, although the exact proportion is not known. However, no evidence exists that soy supplementation has any prostate cancer–related benefits for these men. Soy contains several constituents, including isoflavones, which possess anticancer activities in laboratory studies. Several animal studies also provide support for the hypothesis that soy consumption may protect against prostate cancer.

The majority of prostate cancers detected by prostate-specific antigen (PSA) screening are indolent; only those that have potential to progress to an aggressive, fatal phenotype are clinically (or biologically) significant. Thus, prevention approaches focusing on biologically significant cancers have the greatest potential to reduce prostate cancer–specific mortality. We investigated the effect of soy supplementation on biologically significant prostate cancer in a randomized trial in men who were at high risk of recurrence after radical prostatectomy. The objective was to determine whether daily consumption of a soy protein–based supplement for 2 years reduced the rate of recurrence or delayed recurrence after radical prostatectomy using PSA failure as the intermediate end point. To our knowledge, there have been no randomized clinical trials testing this hypothesis.

Methods

This study and its consent process were approved by the institutional review boards of all participating institutions. Flow of study participant screening, enrollment, and treatment is shown in Figure 1.

Eligibility

Patients were eligible if they had undergone radical prostatectomy for clinically localized (T1c or T2) prostate cancer less than 4 months before randomization, had a postoperative PSA value of less than 0.07 ng/mL confirmed by the assay used in this study, and fulfilled 1 or more of the following criteria for high risk: preoperative PSA of greater than 20.0 ng/mL, final Gleason score of 8 or greater, established positive surgical margins (but not apical margins), established extracapsular extension (but not in the bladder neck), seminal vesicle invasion, or micrometastases in any removed pelvic lymph nodes. To confirm eligibility, prostatectomy slides of each prospective participant were centrally reviewed by 1 of the 4 participating pathologists (J.M., M.X.K., V.M., and A.K.-B.) who collectively had established standardized diagnostic criteria as members of the Cooperative Prostate Cancer Tissue Resource.

Recruitment

Participants were enrolled at the New York University School of Medicine (NYU) (88%) and the Manhattan VA Medical Center (VA) (7%), both in New York City, as well as 5 other medical centers (5%) (Duke University, Durham, North Carolina; Moffitt Cancer Center, Tampa, Florida; Long Beach VA Medical Center, Long Beach, California; University of Chicago, Chicago, Illinois; and University of Illinois at Chicago); one private practitioner referred a single eligible patient. The pathology reports of all radical prostatectomy patients at the VA and NYU sites between July 1997 and November 2005 (VA) or May 2009 (NYU) were systematically screened. At the other sites, no systematic procedure to identify eligible patients was implemented, and enrollment relied on individual urologists. Recruitment at the University of Chicago and the University of Illinois at Chicago was continued until May 2010. Patients whose potential eligibility was confirmed by a study pathologist were first approached by participating urologists (by letter at NYU) asking them to contact study staff. If they agreed to participate and no exclusionary factors (Figure 1) were identified during an in-person or telephone interview, a baseline visit was conducted by a study coordinator to obtain written informed consent, demographic information (including self-identified race/ethnicity using categories listed in Table 1), and a blood sample to confirm postoperative PSA level. A significant baseline intake of soy (more than once per week) was also an exclusionary criterion; this was assessed using a standardized questionnaire.

Randomization and Blinding

After eligibility was confirmed, participants were randomized to the intervention or placebo groups (1:1) using the dynamic intervention allocation procedure of Begg and Iglewicz, stratified by (1) hospital/clinical site (NYU vs VA vs other sites); (2) number of high-risk characteristics (1 vs >1); and (3) self-defined race/ethnicity (African American vs non-African American). The randomization procedure was programmed using SAS software (SAS Institute Inc) and carried out by study staff. All study investigators and staff as well as all study participants remained blinded to the assigned treatments until after the interim analysis was conducted, the trial was stopped, and the intention-to-treat analysis had been completed.

Intervention

The intervention agent was soy protein isolate based and the placebo was a caseinate-based product, each incorporated into a beverage powder developed and manufactured for this clinical trial by Solae LLC. A daily 47-g serving of beverage powder contained protein in the form of either soy protein isolate (19.2 g as analyzed) or calcium caseinate (19.8 g). The soy protein dose represents 37% to 40% of the daily reference pro-
The soy protein beverage powder contained per 1 g of protein 3.67 mg of all forms of isoflavones (aglycones, glycosides, and glycoside esters) or 2.13 mg as aglycone equivalents, amounting to (in aglycone equivalents) 1.24 mg of genistein, 0.78 mg of daidzein, and 0.11 mg of glycitein. The beverage powders were sweetened with a mixture of sucrose and fructose to improve palatability. Artificial strawberry flavoring was added to mask the taste difference between the 2 powders, which were packaged identically and differed nutritionally only in the type of protein and the presence of isoflavones and other soy-specific constituents. The nutrient composition of the powders is listed in Table 2. NYU indicates New York University; PSA, prostate-specific antigen.

*aExclusionary medical factors included recent or current history of anemia, iron deficiency problems or subclinical iron deficiency at baseline, diabetes or insulin resistance requiring use of medication, thyroid disease, significant renal impairment, need for a sodium-restricted diet, substantive tendency to be constipated (grade ≥2 experienced regularly), a medical problem precluding the consumption of soy or casein such as allergies to soy or milk protein, and postoperative PSA of 0.07 ng/mL or higher. Fifty-four patients had diabetes, 8 had thyroid disease, 4 had anemia or low iron status, 2 had other malignancies, 1 had renal disease, and 1 had mental disease.

*bThe 4-month postsurgery deadline had passed for these patients before pathology review could occur.

*cPreviously undetected exclusionary medical factors included 4 patients who had diabetes, 1 who had thyroid disease, 1 who had anemia, 2 who had protein allergies, 1 who had recurrent constipation, 1 who had a restricted diet, and 1 who had mental disease.

*dEight eligible participants were referred to the study by participating clinical sites other than NYU and the Manhattan VA; 1 eligible participant was referred to the study by a urologist in private practice. All 9 participants were confirmed to be at high risk by pathology review.

*eThe median time between surgery and randomization was identical in both groups (14 weeks; 95% CI, 13.1-14.1; range, 7 or 8 to 18 weeks).
Participants were instructed to consume one 47-g packet of beverage powder mixed in approximately 10 oz of water or fruit juice each day.

**Participant Follow-up**
Participants were counseled by a study coordinator on avoiding intake of soy-based food and drink and limiting excessive dairy products while in the study. If postsurgical serum PSA level was confirmed to be less than 0.07 ng/mL, participants started the intervention within 1 or 2 weeks following randomization for 2 years and returned for follow-up visits at 2-month intervals during the first year and at 3-month intervals thereafter. At each follow-up visit, a blood sample was obtained for PSA measurement and occurrence of adverse events was assessed using the National Cancer Institute's Common Toxicity Criteria version 4.0. For some participants, a number of follow-up visits were conducted by telephone and blood samples were mailed overnight by express delivery to the trial office; this has been shown to be feasible without affecting PSA levels, which we confirmed. The intervention was stopped when biochemical recurrence or serious adverse effects developed.

**Adherence**
Adherence was assessed by measuring serum isoflavone levels at baseline and at least 2 time points while in the study by high-performance liquid chromatography with electrochemical detection using the procedure of Gamache and Acworth, with slight modifications as described by Franke et al. Self-reported adherence was also assessed using a daily calendar, and soy intake during the previous 2 to 3 months was evaluated at each follow-up visit using the aforementioned questionnaire.

**End Points**
The primary end points of this trial were the 2-year rate of biochemical cancer recurrence and time to recurrence for those in whom cancer recurred. Biochemical recurrence was de-
fined as development of a serum PSA level of 0.07 ng/mL or higher confirmed by 2 subsequent PSA values of 0.07 ng/mL or higher at least 1 month apart. Serum PSA levels were measured with an ultrasensitive automated immunoenzymometric assay (Tosoh Bioscience) using the 2 standard Tandem Hybritech monoclonal antibodies. We determined that the limit of quantitation of this assay was 0.03 ng/mL and the intra-assay and interassay variation were 4.0% to 6.2% and 8.4% to 13.9%, respectively.

Statistical Analysis
The low detection limit of the PSA assay allowed us to define biochemical recurrence as a sustained increase in PSA level above 0.07 ng/mL approximately 1 year earlier than if we had used a higher cutoff (≥0.1 or 0.2 ng/mL) as commonly used. On the basis of a literature review and an analysis of data from NYU, as detailed in the eTable in the Supplement, the projected 2-year biochemical recurrence rate in the placebo group was estimated to be 30%. With 252 evaluable participants (126 per group) the study had 80% power to detect a 50% reduction in biochemical recurrence rate with a 2-sided significance level of 0.05 and 1 planned interim analysis after observation of 45 recurrences using the O’Brien-Fleming spending function. The 50% reduction in recurrence was chosen because (1) the prostate cancer risk reduction by soy in animal studies has been in this range; (2) the prostate cancer rate differential between populations who habitually consume soy and those who do not is more than 2-fold; and (3) a risk reduction of this magnitude would be needed to have important public health effects. Baseline participant characteristics and adverse events were compared between the 2 groups using the t test or the Mann-Whitney test for continuous variables and the χ² test or Fisher exact test for categorical variables. Time to recurrence was calculated as the time (in weeks) between randomization and first occurrence of a sustained PSA level of 0.07 ng/mL or higher. Participants who ceased study participation and had a PSA value of less than 0.07 ng/mL at their last visit were censored at the time of their last PSA measurement. Because PSA measurements during follow-up were necessary for the assessment of the main study outcome (ie, PSA recurrence), participants who were found ineligible after randomization, withdrew before their baseline visit, or never returned after baseline were excluded from the main intention-to-treat analysis, which was therefore a modified intention-to-treat analysis. The Cox proportional hazards model was used to calculate the hazard ratios and 95% confidence intervals comparing the 2 treatment groups and the log-rank test to assess statistical significance. The proportional hazard assumption was checked by inclusion of the cross-product of log (follow-up time) by treatment. SAS software, versions 9.2 and 9.1.3, and GraphPad Prism software, version 4.0, were used for statistical analysis.

Results
A total of 177 eligible participants were randomized between July 1997 and May 2010. As planned, a blinded interim analysis was conducted once 45 participants had developed confirmed biochemical recurrence, which was reviewed by the

### Table 2. Supplement Nutritional Information

<table>
<thead>
<tr>
<th>Nutrient/Dietary Factor</th>
<th>Calculated Amount Per 47-g Daily Serving</th>
<th>Daily Value, %&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein, g</td>
<td>20&lt;sup&gt;b&lt;/sup&gt;</td>
<td>36</td>
</tr>
<tr>
<td>Total carbohydrate, g</td>
<td>21</td>
<td>16</td>
</tr>
<tr>
<td>Sugars, g</td>
<td>20</td>
<td>NA</td>
</tr>
<tr>
<td>Total fat, g</td>
<td>1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>NA</td>
</tr>
<tr>
<td>Energy, kcal</td>
<td>175</td>
<td>-9&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Sodium, mg</td>
<td>200&lt;sup&gt;b&lt;/sup&gt;</td>
<td>15</td>
</tr>
<tr>
<td>Calcium, mg</td>
<td>700&lt;sup&gt;b&lt;/sup&gt;</td>
<td>70</td>
</tr>
<tr>
<td>Phosphorus, mg</td>
<td>500</td>
<td>71</td>
</tr>
<tr>
<td>Magnesium, mg</td>
<td>40</td>
<td>9.5</td>
</tr>
<tr>
<td>Vitamin A, IU</td>
<td>500</td>
<td>17</td>
</tr>
<tr>
<td>Vitamin C, mg</td>
<td>2.4</td>
<td>2.7</td>
</tr>
<tr>
<td>Vitamin D, IU</td>
<td>100</td>
<td>17</td>
</tr>
<tr>
<td>Vitamin B&lt;sub&gt;6&lt;/sub&gt;, mg</td>
<td>0.12</td>
<td>7</td>
</tr>
<tr>
<td>Vitamin B&lt;sub&gt;12&lt;/sub&gt;, mg</td>
<td>0.9</td>
<td>38</td>
</tr>
<tr>
<td>Thiamin, mg</td>
<td>0.09</td>
<td>7.3</td>
</tr>
<tr>
<td>Folate, µg</td>
<td>60</td>
<td>15</td>
</tr>
<tr>
<td>Riboflavin, mg</td>
<td>0.4</td>
<td>31</td>
</tr>
<tr>
<td>Pantothenic acid, mg</td>
<td>0.8</td>
<td>16</td>
</tr>
<tr>
<td>Zinc, mg</td>
<td>0.9</td>
<td>8</td>
</tr>
<tr>
<td>Iron, mg</td>
<td>3.6</td>
<td>45</td>
</tr>
<tr>
<td>Total isoflavones, mg</td>
<td>41&lt;sup&gt;d&lt;/sup&gt;</td>
<td>NA</td>
</tr>
<tr>
<td>Genistein, mg</td>
<td>24&lt;sup&gt;d&lt;/sup&gt;</td>
<td>NA</td>
</tr>
</tbody>
</table>

Abbreviation: NA, not applicable.

<sup>a</sup> Percentage of dietary reference intake; the first number is based on the current Recommended Daily Allowances and Adequate Intakes based on the dietary reference intakes of the Nutrition Board of the Institute of Medicine (IOM)<sup>21</sup> and the second number is based on the current food labeling regulations of the US Food and Drug Administration (FDA) based on a 2000-kcal/d diet with 50 g protein.<sup>22</sup> Amounts of saturated fats, cholesterol, and dietary fiber were negligible.

<sup>b</sup> The actual measured amounts were 19.2 and 19.8 g protein, 112 and 0.68 g total fat, 196 and 184 mg sodium, and 707 and 707 mg calcium for soy and placebo products, respectively.

<sup>c</sup> Percentage of a 2000-kcal diet.

<sup>d</sup> Isoflavones (as aglycone equivalents) from soy not present in the placebo product.
study’s data safety and monitoring committee. At the recommendation of this committee, the trial was stopped at that time because of lack of evidence of treatment effect. The conditional power—i.e., the probability of observing a significant result at the end of the trial if enrollment had continued until the target sample size had been reached and all participants had been followed-up for two years—was extremely low at 0.0012, given the data observed at the interim analysis.

Of the 177 randomized participants, 18 (10.2%) did not contribute data and were thus not evaluable because they had ineligibly high baseline PSA (n = 3), withdrew before their baseline visit (n = 2), or never returned after baseline (n = 13) (Figure 1). These participants were not evaluable because they had no postrandomization PSA measurements. A total of 159 participants (81 in the intervention group and 78 in the placebo group) completed the baseline visit and at least 1 follow-up visit and were therefore evaluable for biochemical recurrence. There were no significant differences between the 2 treatment groups in any of the baseline characteristics of these evaluable participants (Table 1). Thirteen of the evaluable participants (8.2%) withdrew before recurrence or completion of 2 years of treatment. They were included in the intention-to-treat analysis with censoring at the time of withdrawal; 6 of these were in the intervention group (median number of weeks in study, 15 [range, 8-69 weeks]) and 7 were in the placebo group (median number of weeks in study, 29 [range, 10-94 weeks]) (P = .88 for difference in dropout incidence and P = .07 for difference in number of weeks). Most evaluable participants (n = 146 [92%]) completed 2 years of intervention or experienced recurrence earlier. Less than 2% of all PSA follow-up data were not obtained and in no case did this impair detection of biochemical recurrence. Three participants were classified as having recurrence based on only 1 PSA measurement of 0.07 ng/mL or higher at their last study visit because there was evidence from chart review that they subsequently had more than 1 PSA value higher than 0.1 ng/mL and/or hormone ablation or radiation treatment.

Twenty two (27.2%) of the 81 participants in the intervention group developed confirmed biochemical recurrence, whereas 23 (29.5%) of the 78 participants receiving placebo developed recurrence. The modified intention-to-treat analysis of the evaluable participants revealed no evidence of treatment effect with the 2 survival curves closely overlapping (Figure 2). The hazard ratio for the difference between the 2 groups was 0.96 (95% CI, 0.53-1.72; log-rank P = .89). Among participants who developed recurrence, the median time to recurrence was somewhat shorter in the intervention group (31.5 weeks) than in the placebo group (44 weeks), but this difference was not statistically significant (P = .62 by Mann-Whitney test).

Self-reported adherence was excellent, with 152 participants (96%) reporting having consumed more than 90% of the packets supplied. Only 7 evaluable participants reported serious nonadherence (consuming <50% of the packets), 3 in the intervention group and 4 in the placebo group. Analysis of serum genistein levels identified 1 additional case of nonadherence in the placebo group in a participant who consistently had genistein levels above 150 ng/mL. These 8 participants were considered definitively nonadherent. An additional 13 participants (7 in the intervention group and 6 in the placebo group) had serum genistein levels that were potentially inconsistent with their treatment assignment (>20 ng/mL in the placebo group and <20 ng/mL in the intervention group); these participants were considered possibly nonadherent. Median serum genistein levels were 0.0 ng/mL (interquartile range, 0.0-3.3 ng/mL), 94.7 ng/mL (interquartile range, 36.7-201.7 ng/mL), and 129.8 ng/mL (interquartile range, 64.6-208.6 ng/mL) at baseline and after 6 and 12 months, respectively, in the 33 of 69 fully adherent participants taking soy for whom we had complete data at these time points.

Eleven evaluable participants who were considered fully adherent stopped treatment at some point during the study but continued follow-up without treatment; they were included in the intention-to-treat analysis up to the point of last follow-up. Analysis censoring these participants at the time they ceased taking their assigned treatment and excluding participants who were considered definitively nonadherent (n = 8) did not change the results (hazard ratio, 0.89; 95% CI: 0.49-1.62; log-rank P = .70). Further eliminating the 13 participants...
who were possibly nonadherent from the analysis also did not change the outcome (hazard ratio, 0.97; 95% CI, 0.50-1.73; log-rank P = .81).

There were no differences in adverse events between the 2 groups (Table 3). The major reason for withdrawal from the study was difficulty with palatability or sweetness of the products. The second most frequent reason was lack of time or frequent traveling interfering with participation. Three participants (2 in the placebo group and 1 in the intervention group) stopped treatment because of symptoms that proved to be not related to treatment and 1 participant in the intervention group stopped because of recurrent grade 2 constipation considered treatment related; all 4 participants continued follow-up. No deaths occurred during the study.

### Discussion

This randomized clinical trial was stopped early because it was found that development of biochemical recurrence of prostate cancer after radical prostatectomy was not reduced or delayed by daily consumption of a 20-g soy protein isolate supplement. Intention-to-treat survival analysis yielded overlapping curves and a hazard ratio close to 1. An analysis limited to adherent participants did not change the results. To our knowledge, this is the first randomized clinical trial to test the efficacy of soy protein in reducing biochemical recurrence of prostate cancer. The soy treatment duration in this study was one of the longest reported to date, and this study demonstrated that soy protein isolate supplementation for 2 years is well tolerated and safe in men.

The only other interventional studies with soy relevant to prostate cancer used serum PSA levels as an end point and had mixed results. Soy protein or isoflavone supplementation did not significantly affect serum PSA levels in randomized trials with healthy men of different ages, men with prostate cancer under active surveillance, men with high-grade prostate intraepithelial neoplasia but no cancer on biopsy, and men with untreated prostate cancer. Consumption of isoflavones or soy as supplements to the diet slowed down progression in subsets of men with increasing PSA after surgical intervention or radiation in small studies with short intervention periods.

The lack of protective activity of soy against prostate cancer recurrence observed in this study was limited to men at above-average risk of recurrence within the first 2 years after surgery and to the soy protein dose tested. The findings of this study may therefore not be generalizable to prostate cancer patients at average risk of recurrence. It is also possible that the dose or duration of treatment used in this trial were not sufficient to affect aggressive prostate cancer. It is unlikely that the use of a casein-based supplement as placebo prevented detection of a protective effect of soy as, to our knowledge, no published evidence exists that milk or milk protein reduces risk of prostate cancer or improves clinical outcome. The modest representation of minority participants (self-identified nonwhite) limits the generalizability of the results to white men. The accrual to this study was slow and the possibility of accrual bias thus cannot be excluded but is unlikely because accrual by calendar year was evenly distributed between the 2 groups.

The design of the present study represents one possible approach to clinical studies for the assessment of the efficacy of chemoprevention agents for prostate cancer. Prostate cancer chemoprevention treatments are likely to be initiated in middle-aged men when as many as 50% already have small prostate cancers, of which only a small fraction have aggressive potential. Thus, focusing on prostate cancer patients at high risk of postsurgical recurring cancers, which are aggressive and biologically significant, has potential to identify treatments that reduce prostate cancer mortality. In addition, the study design is simple, does not interfere with clinical practice, and requires a much smaller sample size than prevention trials with healthy men at high or average risk of prostate cancer. Also, men with prostate cancer may be eager to participate in such trials (50% in this study), but effective recruitment requires strong support from participating urologists.

The findings of this study provide another example that associations in observational epidemiologic studies between purported preventive agents and clinical outcomes need con-

### Table 3. Most Frequent Adverse Events by Treatment Group

<table>
<thead>
<tr>
<th>Adverse Events</th>
<th>Intervention (n = 81)</th>
<th>Placebo (n = 78)</th>
<th>P Value for Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of adverse events per participant, mean (SD)</td>
<td>1.74 (1.84)</td>
<td>1.42 (1.53)</td>
<td>.40a</td>
</tr>
<tr>
<td>Participants with event, No. (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any adverse event</td>
<td>50 (61.7)</td>
<td>50 (64.1)</td>
<td>.87b</td>
</tr>
<tr>
<td>1 Adverse event</td>
<td>26 (32.1)</td>
<td>28 (35.9)</td>
<td>.88c</td>
</tr>
<tr>
<td>&gt;1 Adverse event</td>
<td>33 (40.7)</td>
<td>30 (38.4)</td>
<td></td>
</tr>
<tr>
<td>Gastrointestinal issues, all</td>
<td>13 (16.1)</td>
<td>8 (10.3)</td>
<td>.35b</td>
</tr>
<tr>
<td>Potentially treatment related</td>
<td>6 (7.4)</td>
<td>6 (7.7)</td>
<td>.99b</td>
</tr>
<tr>
<td>Not treatment related</td>
<td>7 (8.6)</td>
<td>2 (2.6)</td>
<td>.19b</td>
</tr>
<tr>
<td>Urinary tract issues (surgery related)</td>
<td>13 (16.1)</td>
<td>8 (10.2)</td>
<td>.35b</td>
</tr>
<tr>
<td>Initiation of high cholesterol treatment</td>
<td>9 (11.1)</td>
<td>10 (12.8)</td>
<td>.81b</td>
</tr>
<tr>
<td>Initiation of hypertension treatment</td>
<td>9 (11.1)</td>
<td>7 (9.0)</td>
<td>.79b</td>
</tr>
<tr>
<td>Musculoskeletal pain, not treatment related</td>
<td>11 (13.6)</td>
<td>4 (5.1)</td>
<td>.10b</td>
</tr>
</tbody>
</table>

* By Mann-Whitney test.  
* By Fisher exact test.  
* By χ2 test comparing no adverse events with 1 or more than 1.  
* Gastrointestinal problems that could be treatment related included constipation, bloating, irregular bowel movements, diverticulosis, diverticulitis, and heartburn.


