Adjuvant Ovarian Suppression in Premenopausal Breast Cancer


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*A complete list of investigators in the Suppression of Ovarian Function Trial (SOFT) and the International Breast Cancer Study Group is provided in the Supplementary Appendix, available at NEJM.org.

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BACKGROUND
Suppression of ovarian estrogen production reduces the recurrence of hormone-receptor–positive early breast cancer in premenopausal women, but its value when added to tamoxifen is uncertain.

METHODS
We randomly assigned 3066 premenopausal women, stratified according to prior receipt or nonreceipt of chemotherapy, to receive 5 years of tamoxifen, tamoxifen plus ovarian suppression, or exemestane plus ovarian suppression. The primary analysis tested the hypothesis that tamoxifen plus ovarian suppression would improve disease-free survival, as compared with tamoxifen alone. In the primary analysis, 46.7% of the patients had not received chemotherapy previously, and 53.3% had received chemotherapy and remained premenopausal.

RESULTS
After a median follow-up of 67 months, the estimated disease-free survival rate at 5 years was 86.6% in the tamoxifen–ovarian suppression group and 84.7% in the tamoxifen group (hazard ratio for disease recurrence, second invasive cancer, or death, 0.83; 95% confidence interval [CI], 0.66 to 1.04; P = 0.10). Multivariable allowance for prognostic factors suggested a greater treatment effect with tamoxifen plus ovarian suppression than with tamoxifen alone (hazard ratio, 0.78; 95% CI, 0.62 to 0.98). Most recurrences occurred in patients who had received prior chemotherapy, among whom the rate of freedom from breast cancer at 5 years was 82.5% in the tamoxifen–ovarian suppression group and 78.0% in the tamoxifen group (hazard ratio for recurrence, 0.78; 95% CI, 0.60 to 1.02). At 5 years, the rate of freedom from breast cancer was 85.7% in the exemestane–ovarian suppression group (hazard ratio for recurrence vs. tamoxifen, 0.65; 95% CI, 0.49 to 0.87).

CONCLUSIONS
Adding ovarian suppression to tamoxifen did not provide a significant benefit in the overall study population. However, for women who were at sufficient risk for recurrence to warrant adjuvant chemotherapy and who remained premenopausal, the addition of ovarian suppression improved disease outcomes. Further improvement was seen with the use of exemestane plus ovarian suppression. (Funded by Pfizer and others; SOFT ClinicalTrials.gov number, NCT00066690.)
ADJUVANT ENDOCRINE THERAPY WITH tamoxifen has been recommended for premenopausal women with hormone-receptor–positive breast cancer (positive for estrogen receptor, progesterone receptor, or both) during the past 15 years.\(^1\)\(^2\) The value of therapeutic suppression of ovarian estrogen production in premenopausal women who receive tamoxifen is uncertain.\(^3\) The American Society of Clinical Oncology endorsed guidelines recommending that ovarian ablation or suppression (hereafter, ovarian suppression) not be added routinely to adjuvant therapy in premenopausal women.\(^4\) Chemotherapy-induced ovarian suppression (amenorrhea) is correlated with a reduced risk of relapse\(^5\)\(^6\) but is less likely to be achieved in very young women. International consensus guidelines for breast-cancer management in young women suggested that the addition of a gonadotropin-releasing hormone (GnRH) agonist to tamoxifen be discussed on an individualized basis.\(^8\)

In 2003, the International Breast Cancer Study Group (IBCSG) initiated two randomized, phase 3 trials, the Suppression of Ovarian Function Trial (SOFT) and the Tamoxifen and Exemestane Trial (TEXT), involving premenopausal women with hormone-receptor–positive early breast cancer. SOFT was designed to determine the value of adding ovarian suppression to tamoxifen and to determine the role of adjuvant therapy with the aromatase inhibitor exemestane plus ovarian suppression in premenopausal women. Here we report the results of the planned primary analysis in SOFT\(^9\) comparing adjuvant tamoxifen plus ovarian suppression with tamoxifen alone after a median follow-up of 67 months.

METHODS

PATIENTS

The trial was designed to evaluate adjuvant endocrine therapy in women who remained premenopausal after the completion of adjuvant or neo-adjuvant chemotherapy and in premenopausal women for whom adjuvant tamoxifen alone was considered suitable treatment. Eligibility criteria included documented premenopausal status, operable breast cancer, and tumor that expressed estrogen or progesterone receptors in at least 10% of the cells (see the Supplementary Appendix, available with the full text of this article at NEJM.org).

Patients had to have undergone either a total mastectomy with subsequent optional radiotherapy or breast-conserving surgery with subsequent radiotherapy. Either axillary dissection or a sentinel-node biopsy was required. Patients who had not received chemotherapy underwent randomization within 12 weeks after definitive surgery. Patients who received chemotherapy before randomization and remained premenopausal were enrolled within 8 months after completing chemotherapy, once a premenopausal estradiol level was confirmed by a local laboratory. Patients were allowed to receive adjuvant oral endocrine therapy before randomization.

STUDY DESIGN

Women were randomly assigned to receive oral tamoxifen at a dose of 20 mg daily, tamoxifen plus ovarian suppression, or oral exemestane (Aromasin, Pfizer) at a dose of 25 mg daily plus ovarian suppression. Treatment was for 5 years from the date of randomization, according to the study protocol, available at NEJM.org. Ovarian suppression was achieved by choice of triptorelin (Decapeptyl Depot [triptorelin acetate], Ipsen; or Trelstar Depot [triptorelin pamoate], Debio) at a dose of 3.75 mg administered by means of intramuscular injection every 28 days, bilateral oophorectomy, or bilateral ovarian irradiation. Patients receiving triptorelin could subsequently opt to undergo oophorectomy or irradiation. Randomization was performed by means of the IBCSG Internet-based system and was stratified according to prior chemotherapy (yes vs. no), lymph-node status (positive vs. negative), and intended initial method of ovarian suppression, if assigned. The assessments of the patients and the recording of adverse events followed a regular schedule (see the Supplementary Appendix).

The primary end point was disease-free survival, defined as the time from randomization to the first appearance of one of the following: recurrence of invasive breast cancer (local, regional, or distant), invasive contralateral breast cancer, second (nonbreast) invasive cancer, or death without recurrence or second cancer. Secondary end points included the interval without breast cancer, defined as the time from randomization to the recurrence of invasive breast cancer (local, regional, or distant) or invasive contralateral breast cancer; the interval from randomization to the recurrence of breast cancer at a distant site; and overall survival, defined as the time from randomization to death from any cause.

The ethics committee at each participating...
center approved the study protocol, and all the patients provided written informed consent. The IBCSG was responsible for the trial design, data collection, and analysis. Pfizer and Ipsen, the respective manufacturers of exemestane and trip-torelin, donated the study drugs; neither manufacturer imposed restrictions with respect to the trial data. The manuscript was written solely by the authors, who vouch for the data and analyses reported and for the fidelity of the study to the protocol. The steering committee (which included employees of Pfizer and Ipsen) reviewed the manuscript and made the decision to submit it for publication.

STATISTICAL ANALYSIS
The original statistical analysis plan for SOFT was to assess disease-free survival between the treatment groups with three pairwise comparisons. The design assumed the enrollment of predominantly very young women who remained premenopausal after chemotherapy and would have an expected disease-free survival rate at 5 years of 67% when treated with tamoxifen, on the basis of outcomes for patients younger than 35 years of age in previous trials. The enrolled patients were older and had lower-risk characteristics than anticipated, and the rate of disease-free survival was higher than expected. A protocol amendment to the analysis plan was adopted in 2011, designating the test of the superiority of tamoxifen plus ovarian suppression over tamoxifen alone as the primary analysis for SOFT. We calculated that with an estimated 186 events of disease recurrence, second invasive cancer, or death in the two treatment groups after a median follow-up of 5 years, the study would have at least 80%, 69%, and 52% power to detect reductions in risk of 33.5%, 30%, and 25%, respectively, with tamoxifen plus ovarian suppression versus tamoxifen alone, at a two-sided alpha level of 0.05. The comparison of exemestane plus ovarian suppression with tamoxifen alone became a secondary objective, and the comparison of exemestane plus ovarian suppression with tamoxifen plus ovarian suppression was analyzed by means of a combined analysis with the TEXT data.

Analyses were performed according to the

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Figure 1. Randomization and Primary Analysis Populations.
The flow diagram shows the intention-to-treat population of 2033 patients included in the primary analysis (shaded) of tamoxifen plus ovarian suppression, as compared with tamoxifen alone, and the analogous population of patients assigned to receive exemestane plus ovarian suppression. Additional details are provided in Figure S1 in the Supplementary Appendix.
intention-to-treat principle. Kaplan–Meier estimates of time-to-event end points were calculated. Hypothesis tests compared the two groups with the use of log-rank tests, stratified according to prior use of chemotherapy (yes vs. no) and lymph-node status (positive vs. negative). Stratified Cox proportional-hazards regression was used to estimate hazard ratios and 95% confidence intervals. In prespecified secondary analyses, heterogeneity of the treatment effect according to subgroup was investigated by tests of treatment-by-covariate interaction, and an adjusted hazard ratio for the treatment effect was estimated.

### STUDY POPULATION

From December 2003 through January 2011, we randomly assigned 1021 premenopausal women to tamoxifen, 1024 to tamoxifen plus ovarian suppression, and 1021 to exemestane plus ovarian suppression. After exclusions, 2033 women were included in the intention-to-treat population for the primary analysis comparing tamoxifen plus ovarian suppression with tamoxifen alone (Fig. 1, and Fig. S1 in the Supplementary Appendix). The median age of the patients was 43 years (Table 1). A total of 46.7% of the pa-

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No Chemotherapy (N = 949)</th>
<th>Prior Chemotherapy (N = 1084)</th>
<th>Overall (N = 2033)</th>
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<tr>
<td>Age at randomization</td>
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<tr>
<td>Median  — yr</td>
<td>46</td>
<td>40</td>
<td>43</td>
</tr>
<tr>
<td>Distribution — no. (%)</td>
<td></td>
<td></td>
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<tr>
<td>&lt;35 yr</td>
<td>14 (1.5)</td>
<td>219 (20.2)</td>
<td>233 (11.5)</td>
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<td>35–39 yr</td>
<td>78 (8.2)</td>
<td>309 (28.5)</td>
<td>387 (19.0)</td>
</tr>
<tr>
<td>40–49 yr</td>
<td>702 (74.0)</td>
<td>522 (48.2)</td>
<td>1224 (60.2)</td>
</tr>
<tr>
<td>≥50 yr</td>
<td>155 (16.3)</td>
<td>34 (3.1)</td>
<td>189 (9.3)</td>
</tr>
<tr>
<td>Lymph-node status — no. (%)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>861 (90.7)</td>
<td>463 (42.7)</td>
<td>1324 (65.1)</td>
</tr>
<tr>
<td>Positive</td>
<td>88 (9.3)</td>
<td>621 (57.3)</td>
<td>709 (34.9)</td>
</tr>
<tr>
<td>Tumor size — no. (%)†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤2 cm</td>
<td>806 (84.9)</td>
<td>526 (48.5)</td>
<td>1332 (65.5)</td>
</tr>
<tr>
<td>&gt;2 cm</td>
<td>136 (14.3)</td>
<td>513 (47.3)</td>
<td>649 (31.9)</td>
</tr>
<tr>
<td>Tumor grade — no. (%)‡</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>389 (41.0)</td>
<td>151 (13.9)</td>
<td>540 (26.6)</td>
</tr>
<tr>
<td>2</td>
<td>483 (50.9)</td>
<td>523 (48.2)</td>
<td>1006 (49.5)</td>
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<tr>
<td>3</td>
<td>65 (6.8)</td>
<td>374 (34.5)</td>
<td>439 (21.6)</td>
</tr>
<tr>
<td>HER2-positive — no. (%)§</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>40 (4.2)</td>
<td>196 (18.1)</td>
<td>236 (11.6)</td>
</tr>
<tr>
<td>Interval from surgery to randomization — mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>1.8</td>
<td>8.0</td>
<td>3.2</td>
</tr>
<tr>
<td>Interquartile range</td>
<td>1.2–2.4</td>
<td>5.8–10.3</td>
<td>1.7–8.33</td>
</tr>
<tr>
<td>Endocrine therapy before randomization — no. (%)§</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>47 (5.0)</td>
<td>475 (43.8)</td>
<td>522 (25.7)</td>
</tr>
</tbody>
</table>

* A more complete summary is provided in Table S1 in the Supplementary Appendix. Characteristics were well balanced according to treatment assignment (Table S2 in the Supplementary Appendix). The statistical analysis plan did not specify hypothesis testing regarding comparisons between these groups. HER2 denotes human epidermal growth factor receptor 2. † Data were missing for 7 patients who did not receive chemotherapy and for 45 who had received chemotherapy previously. ‡ Data were missing for 12 patients who did not receive chemotherapy and for 36 who had received chemotherapy previously. § Oral endocrine therapy before randomization was allowed while premenopausal status was established or reestablished.
tients had not received chemotherapy previously, and 53.3% received chemotherapy before randomization and remained premenopausal. Node-positive disease was present in 34.9% of the patients.

**Efficacy**

At a median follow-up of 67 months, 299 patients (14.7%) had recurrent disease or a second invasive cancer or had died. The rate of disease-free survival at 5 years was 86.6% (95% confidence interval [CI], 84.2 to 88.7) among patients assigned to receive tamoxifen plus ovarian suppression, as compared with 84.7% (95% CI, 82.2 to 86.9) among those assigned to receive tamoxifen alone (hazard ratio for recurrence, second invasive cancer, or death, 0.83; 95% CI, 0.66 to 1.04; P=0.10) (Fig. 2A). A total of 58.2% of the first events involved distant sites, and 12.0% were second (nonbreast) malignant conditions (Table S3 in the Supplementary Appendix). Planned subgroup analyses did not reveal heterogeneity of treatment effect across most subgroups (Fig. S2 in the Supplementary Appendix). However, the subgroup of patients with human epidermal growth factor receptor 2–positive tumors appeared to have a greater benefit with tamoxifen plus ovarian suppression than with tamoxifen alone. In the multivariable Cox proportional-hazards model, with adjustment for covariates, tamoxifen plus ovarian suppression significantly reduced the hazard of recurrence, a second invasive cancer, or death, as compared with tamoxifen alone (hazard ratio, 0.78; 95% CI, 0.62 to 0.98; P=0.03) (Table S4 in the Supplementary Appendix).

At 5 years, 88.4% (95% CI, 86.1 to 90.3) of the patients assigned to receive tamoxifen plus ovarian suppression remained free from breast cancer, as compared with 86.4% (95% CI, 84.0 to 88.5) of those assigned to receive tamoxifen alone (hazard ratio for recurrence, 0.81; 95% CI, 0.63 to 1.03; P=0.09) (Fig. 3A). After adjustment for covariates in the multivariable Cox proportional-hazards model, tamoxifen plus ovarian suppression reduced the hazard of breast-cancer recurrence, as compared with tamoxifen alone (hazard ratio, 0.75; 95% CI, 0.59 to 0.96; P=0.02). Among patients assigned to receive exemestane plus ovarian suppression, 90.9% (95% CI, 88.9 to 92.6) remained free from breast cancer at 5 years.

Recurrence of breast cancer at a distant site was reported in 185 patients (9.1%), with no significant difference between those assigned to tamoxifen plus ovarian suppression and those assigned to tamoxifen alone (hazard ratio for recurrence, 0.88; 95% CI, 0.66 to 1.18; P=0.40) (Fig. 3B). Death was reported in 106 patients (5.2%); 4 patients died without a breast-cancer recurrence or a second invasive cancer. Overall survival at 5 years was 96.7% (95% CI, 95.2 to 97.7) among patients assigned to tamoxifen plus ovarian suppression and 95.1% (95% CI, 93.4 to 96.3) among those assigned to tamoxifen alone (hazard ratio for death, 0.74; 95% CI, 0.51 to 1.09; P=0.13) (Fig. 2B).

Among patients who did not receive chemotherapy, more than 95% remained free from breast cancer at 5 years in each group (Fig. 3C), with few distant recurrences (Fig. 3D), and 32.9% of first events (a second invasive cancer or death) were not related to breast cancer (Table S3 in the Supplementary Appendix). Most recurrences of breast cancer were in patients who remained premenopausal after receiving chemother-
therapy (Fig. 3E). In this cohort, the rate of freedom from breast cancer at 5 years was 82.5% (95% CI, 78.8 to 85.6) among those assigned to receive tamoxifen plus ovarian suppression and 78.0% (95% CI, 74.0 to 81.5) among those assigned to receive tamoxifen alone (hazard ratio for recurrence, 0.78; 95% CI, 0.60 to 1.02). In the chemotherapy cohort, among patients assigned
to exemestane plus ovarian suppression, 85.7% (95% CI, 82.3 to 88.5) remained free from breast cancer at 5 years (Fig. 3E).

Most recurrences of breast cancer at a distant site occurred in the patients who had received chemotherapy previously. The rates of freedom from distant recurrence at 5 years in this cohort were as follows: 83.6% among patients assigned to tamoxifen alone, 84.8% among those assigned to tamoxifen plus ovarian suppression, and 87.8% among those assigned to exemestane plus ovarian suppression (Fig. 3F).

More than 90% of the deaths occurred in patients who had received chemotherapy previously. Overall survival at 5 years in the chemotherapy cohort was 94.5% (95% CI, 92.0 to 96.2) among patients assigned to tamoxifen plus ovarian suppression, as compared with 90.9% (95% CI, 87.9 to 93.2) among those assigned to tamoxifen alone (hazard ratio for death, 0.64; 95% CI, 0.42 to 0.96) (Fig. 2B, and Fig. S5 in the Supplementary Appendix).

A total of 350 women younger than 35 years of age participated in the trial, 233 of whom were included in the primary analysis. Among these women, the rate of freedom from breast cancer at 5 years was 67.7% (95% CI, 57.3 to 76.0) for patients assigned to tamoxifen alone, 78.9% (95% CI, 69.8 to 85.5) for those assigned to tamoxifen plus ovarian suppression, and 83.4% (95% CI, 74.9 to 89.3) for those assigned to exemestane plus ovarian suppression. In this very young subgroup, 94.0% of the patients had received chemotherapy previously.

**TREATMENT AND ADVERSE EVENTS**

At a median follow-up of 67 months, 25.8% of the patients were continuing to receive some or all of the protocol-assigned treatment. Tamoxifen was discontinued early, with or without the substitution of alternative endocrine therapy, in 16.7% of the tamoxifen–ovarian suppression group and 21.7% of the tamoxifen group (Table S5 in the Supplementary Appendix). Rates of nonadherence with ovarian suppression were 5.0%, 9.2%, 14.9%, 18.3%, and 21.9% at 0.5, 1, 2, 3, and 4 years after randomization, respectively. Among patients assigned to ovarian suppression, it was achieved entirely through administration of the GnRH agonist triptorelin in 80.7% of the patients.

Targeted adverse events of grade 3 or higher were reported in 31.3% of the patients assigned to receive tamoxifen plus ovarian suppression, as compared with 23.7% of those assigned to receive tamoxifen alone (Table 2, and Table S6 in the Supplementary Appendix). Hot flushes, sweating, decreased libido, vaginal dryness, insomnia, depression, musculoskeletal symptoms, hypertension, and glucose intolerance (diabetes) were reported more frequently in the tamoxifen–ovarian suppression group than in the tamoxifen group. Osteoporosis as defined by a T score of less than −2.5 was reported in 5.8% of the patients assigned to tamoxifen plus ovarian suppression and in 3.5% of those assigned to tamoxifen alone.

**DISCUSSION**

The results of SOFT show that, considering the entire population of patients who underwent randomization, the addition of ovarian suppression to adjuvant tamoxifen did not significantly improve disease-free survival. However, SOFT investigated ovarian suppression in two distinct patient cohorts. The first cohort included 949 premenopausal women for whom adjuvant tamoxifen without chemotherapy was considered to be suitable treatment. These patients were predominantly older than 40 years of age, had small, node-negative tumors of low to intermediate grade, and had excellent outcomes with tamoxifen alone after a median follow-up of 67 months. The findings in this cohort do not currently inform us about the clinical relevance of ovarian suppression because one third of the first events were not related to breast cancer, freedom from recurrence exceeded 95% at 5 years, and addi-
Adjuvant Ovarian Suppression in Breast Cancer

A Freedom from Breast Cancer

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 1018 140 86.4 0.81 (0.63–1.03)
Tamoxifen–OS 1015 120 88.4 0.81 (0.63–1.03)
Exemestane–OS 1014 94 90.9 0.64 (0.49–0.83)

Years since Randomization

No at Risk
Tamoxifen 1018 956 900 855 728 533 314
Tamoxifen–OS 1015 970 932 886 752 568 356
Exemestane–OS 1014 957 912 869 766 550 342

B Freedom from Distant Recurrence

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 1018 96 90.7 0.88 (0.66–1.18)
Tamoxifen–OS 1015 89 91.3 0.88 (0.66–1.18)
Exemestane–OS 1014 70 93.0 0.71 (0.52–0.96)

Years since Randomization

No at Risk
Tamoxifen 1018 966 915 875 755 559 333
Tamoxifen–OS 1015 977 943 901 772 582 363
Exemestane–OS 1014 962 920 882 783 562 352

C No Chemotherapy, Freedom from Breast Cancer

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 476 24 95.8 0.95 (0.54–1.69)
Tamoxifen–OS 473 23 95.1 0.95 (0.54–1.69)
Exemestane–OS 470 14 97.1 0.59 (0.31–1.14)

Years since Randomization

No at Risk
Tamoxifen 476 461 445 429 377 277 169
Tamoxifen–OS 473 454 447 429 373 285 179
Exemestane–OS 470 443 425 414 374 278 176

D No Chemotherapy, Freedom from Distant Recurrence

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 476 6 98.6 0.71 (0.52–0.96)
Tamoxifen–OS 473 7 98.7 0.71 (0.52–0.96)
Exemestane–OS 470 3 99.3 0.52 (0.13–2.07)

Years since Randomization

No at Risk
Tamoxifen 476 465 449 436 386 284 176
Tamoxifen–OS 473 458 453 437 385 293 184
Exemestane–OS 470 444 429 419 381 283 180

E Prior Chemotherapy, Freedom from Breast Cancer

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 542 116 78.0 0.78 (0.60–1.02)
Tamoxifen–OS 542 97 82.5 0.78 (0.60–1.02)
Exemestane–OS 544 80 85.7 0.65 (0.49–0.87)

Years since Randomization

No at Risk
Tamoxifen 542 494 455 426 352 255 144
Tamoxifen–OS 542 516 485 456 378 283 176
Exemestane–OS 544 514 487 455 391 273 166

F Prior Chemotherapy, Freedom from Distant Recurrence

No of Patients 5-Yr Rate Hazard Ratio (95% CI)
Tamoxifen 542 90 83.6 0.87 (0.64–1.17)
Tamoxifen–OS 542 82 84.8 0.87 (0.64–1.17)
Exemestane–OS 544 67 87.8 0.72 (0.52–0.98)

Years since Randomization

No at Risk
Tamoxifen 542 501 466 439 369 274 156
Tamoxifen–OS 542 519 490 463 386 289 178
Exemestane–OS 544 518 491 463 401 280 172
Table 2. Key Targeted Adverse Events Reported during Follow-up, According to Treatment Assignment.*

<table>
<thead>
<tr>
<th>Adverse Event</th>
<th>Tamoxifen (N = 1006)</th>
<th>Tamoxifen plus Ovarian Suppression (N = 1005)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Any Event</td>
<td>Grade 3 or 4 Event</td>
</tr>
<tr>
<td></td>
<td>no. of patients</td>
<td>% (95% CI)</td>
</tr>
<tr>
<td>Hot flushes</td>
<td>803</td>
<td>79.8 (77.2–82.3)</td>
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<td></td>
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<tr>
<td>Depression</td>
<td>469</td>
<td>46.6 (43.5–49.8)</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>sweating</td>
<td>486</td>
<td>48.3 (45.2–51.4)</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>Insomnia</td>
<td>466</td>
<td>46.3 (43.2–49.5)</td>
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<td></td>
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<tr>
<td>Hypertension</td>
<td>173</td>
<td>17.2 (14.9–19.7)</td>
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<td></td>
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<tr>
<td>Musculoskeletal symptoms</td>
<td>694</td>
<td>69.0 (66.0–71.8)</td>
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<tr>
<td>Osteoporosis</td>
<td>124</td>
<td>12.3 (10.4–14.5)</td>
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<tr>
<td>Vaginal dryness</td>
<td>421</td>
<td>41.8 (38.8–45.0)</td>
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<tr>
<td>Decreased libido</td>
<td>427</td>
<td>42.4 (39.4–45.6)</td>
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<tr>
<td>Glucose intolerance†</td>
<td>18</td>
<td>1.8 (1.1–2.8)</td>
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<tr>
<td>Any targeted adverse event‡</td>
<td>959</td>
<td>95.3 (93.8–96.5)</td>
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* Data are for the 2011 patients in the safety population who received a protocol-assigned treatment (except for 3 patients who withdrew consent within 1 month after randomization and had no adverse-event data submitted). Targeted adverse events (22 events; see Table S6 in the Supplementary Appendix) and other adverse events of grade 3 or higher were categorized according to the Common Terminology Criteria for Adverse Events, version 3.0.11 A dash indicates that grade 3 or 4 was not a possible grade for the specified adverse event. There was one targeted adverse event of grade 5 (cardiac ischemia or infarction in a patient randomly assigned to tamoxifen).

† Glucose intolerance (diabetes) was added as a targeted adverse event in 2011 and therefore may be underreported.

‡ The category of any targeted adverse event includes the 22 targeted adverse events summarized in Table S6 in the Supplementary Appendix.
rence, as compared with tamoxifen alone. After a protocol amendment, the comparison of exemestane plus ovarian suppression with tamoxifen plus ovarian suppression, on the basis of a combined analysis with TEXT, showed a 28% reduction in the relative risk of breast-cancer recurrence, a second invasive cancer, or death and a 34% reduction in the relative risk of breast-cancer recurrence in the exemestane–ovarian suppression group (P<0.001).10

Overall, tamoxifen plus ovarian suppression resulted in an absolute improvement of 2.0 percentage points, as compared with tamoxifen alone, in the proportion of patients without recurrent breast cancer at 5 years. In the higher-risk cohort of patients who remained premenopausal after chemotherapy, tamoxifen plus ovarian suppression resulted in an absolute improvement of 4.5 percentage points, as compared with tamoxifen alone, in the proportion of patients without recurrent breast cancer at 5 years; with exemestane plus ovarian suppression, the absolute improvement was 7.7 percentage points, as compared with tamoxifen alone. These observed relative and absolute benefits at 5 years from ovarian suppression plus tamoxifen or ovarian suppression plus exemestane, as compared with tamoxifen alone, compare favorably with the practice-changing results of randomized trials of adjuvant aromatase inhibitors versus tamoxifen in postmenopausal women.16

Patients who receive a diagnosis of hormone-receptor–positive breast cancer when they are younger than 35 years of age are a subgroup considered to be at higher risk for adverse outcomes than are older premenopausal women, on the basis of retrospective analyses of data from IBCSG and U.S. Intergroup trials.17,18 The results observed in this subgroup in SOFT add to the evidence that ovarian suppression plays an important role in younger premenopausal patients.13–15 Among the women younger than 35 years of age, breast cancer recurred within 5 years in approximately one third of the patients assigned to receive tamoxifen alone but in one sixth of those assigned to receive exemestane plus ovarian suppression.

Any benefit from ovarian suppression must be weighed against the adverse effects. Adding ovarian suppression to tamoxifen resulted in increased adverse events — most notably, menopausal symptoms, depression, and adverse events with possible long-term health implications such as hypertension, diabetes, and osteoporosis. When exemestane is combined with ovarian suppression, adverse sexual, musculoskeletal, and bone-density effects are more frequent than with tamoxifen plus ovarian suppression.10

Longer follow-up is required, because SOFT is currently underpowered, and the overall survival analysis is premature after 5% of patients have died. The combined analysis from TEXT and SOFT showed that adjuvant endocrine therapy with ovarian suppression plus exemestane is significantly more effective than ovarian suppression plus tamoxifen.10 The current analysis indicates that in a cohort selected for chemotherapy and persistent premenopausal status, ovarian suppression plus tamoxifen improves outcomes, as compared with tamoxifen alone, with the most striking differences observed among younger patients.

We conclude that adding ovarian suppression to tamoxifen did not provide a significant benefit in the overall population of premenopausal women in this trial. However, in the cohort of women who had a sufficient risk of recurrence to warrant adjuvant chemotherapy and who had premenopausal estradiol levels despite chemotherapy, ovarian suppression in addition to tamoxifen reduced the risk of breast-cancer recurrence, as compared with tamoxifen alone. Ovarian suppression combined with an aromatase inhibitor further reduced the risk of recurrence, as compared with tamoxifen-based therapy, in this higher-risk premenopausal cohort.

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APPENDIX

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REFERENCES


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