Commentary

Menopausal hormone therapy after breast cancer

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See related research by Col et al., http://breast-cancer-research.com/content/7/4/R535

Abstract

The use of postmenopausal hormone therapy after breast cancer remains controversial. Evidence shows variation by study design, and even among three randomized controlled trials there is substantial heterogeneity of results. Two Swedish trials of comparable size show relative risks of recurrence of 3.3 and 0.82 on comparing women receiving postmenopausal hormone therapy with control women. The extent of use of tamoxifen and concomitant use of progestins in combination with estrogen, although raised as one possible explanation for this heterogeneity, are not supported by evidence from trials of high-dose progestins used after breast cancer. Caution is needed when considering the use of postmenopausal hormone therapy after breast cancer.

There is now resounding evidence that use of exogenous hormones by postmenopausal women increases risk for breast cancer [1-3]. Likewise, obesity among postmenopausal women with breast cancer is a major determinant of relapse and mortality [4,5], being directly related to poor outcomes. On the other hand, tamoxifen provides lasting reduction in risk for recurrence and mortality [6], an effect that is independent of adjuvant chemotherapy. Despite this evidence relating endogenous and exogenous hormones to breast cancer risk and outcomes, debate continues regarding the use of postmenopausal hormones for symptom relief among women with breast cancer. Col and colleagues [7], in this issue of Breast Cancer Research, report a systematic review in which they identified eight observational studies and two randomized controlled trials of postmenopausal hormone therapy. They note that the combined results for the two trials show increased risk for recurrence (relative risk \(RR = 3.4\), 95% confidence interval \([CI] = 1.6\) to 7.3), which contrasts with the findings of the observational results, which show a decrease in risk \((RR = 0.6, 95\% CI = 0.5–0.8)\). While noting heterogeneity between the results according to study design, Col and colleagues do not explore the data to describe the possible sources of heterogeneity [8,9] beyond study design.

As noted below, the quality of data reporting limits their ability in this regard.

Biologic evidence supports estrogenic exposures being related to incidence and poor outcome among women with breast cancer. Weight after diagnosis is a strong predictor of outcome [4], and weight gain after diagnosis is associated with increased recurrence and mortality [5]. There are direct positive relations between adiposity and circulating estrogens [10], and between adult weight gain and postmenopausal breast cancer risk [11], and circulating endogenous hormones represent the primary explanation for this positive relation between adiposity and postmenopausal breast cancer incidence [12]. Together, these data support an increase in incidence of breast cancer with use of postmenopausal hormone therapy. Should the effect of exogenous postmenopausal hormonal therapy differ after diagnosis, when adiposity and weight gain have adverse effects on recurrence and mortality?

To cloud further the current evidence, a third randomized trial [13], which has been published since the synthesis by Col and colleagues, found no increase in risk with use of postmenopausal hormone therapy \((RR = 0.8, 95\% CI = 0.4–1.9)\), based on 4.1 years of follow up in 434 women (Table 1). The authors raise the intriguing possibility that the choice of hormone regimen may modify the risk for recurrence and account for some of the heterogeneity observed. This trial is of comparable size and design to the HABITS (Hormone Replacement Therapy after Breast Cancer – Is It Safe?) trial, also conducted in Sweden [14]. Clearly, more detailed analysis of this small body of evidence might improve our understanding of the difference between unopposed estrogen and combination estrogen plus progestin therapy among women who have been treated for histologically confirmed breast cancer. Alternatively, we can consider other
progestin clearly increases the incidence of breast cancer, therapy (including MPA) [20]. Finally, given that estrogen plus survival advantage for higher versus lower doses of endocrine trials published through 1996 shows no overall improvement light on this possibility. Synthesis of data from randomized trials published through 1996 shows no overall improvement between the observational and randomized study findings in the systematic review conducted by Col and coworkers [7]. As Col and colleagues note, the observational studies often did not report results separately for women using combination hormone therapy and those using unopposed estrogens. It is noteworthy that the Stockholm trial attempted to minimize the use of progestogen, and the HABITS trial recommended either cyclical estrogen plus progestin for women with uterine bleeding within 2 years of randomization or continuous combined therapy for other women with a uterus. The Stockholm trial recommended 20 mg medroxyprogesterone acetate (MPA) for 14 days every 105 days for women over 55 years of age. The Stockholm trial also included a higher percentage of patients receiving cocon-mitant adjuvant tamoxifen therapy than did the HABITS trial (52% versus 21%), and a lower percentage of patients with node-positive breast cancer (16% versus 26%). Thus, a lower risk population may have been studied in the Stockholm trial, with lower exposure to MPA.

It has been hypothesized that MPA might act through glucocorticoid receptors to reduce metastatic potential [17]. Two trials that used high-dose MPA for 6 months in women with breast cancer and followed them for more than a decade [18,19] fail to support a reduction in recurrence. A systematic review of trials of MPA in combination with either tamoxifen or combination chemotherapy after breast cancer sheds some light on this possibility. Synthesis of data from randomized trials published through 1996 shows no overall improvement in median survival among women treated with MPA, and no survival advantage for higher versus lower doses of endocrine therapy (including MPA) [20]. Finally, given that estrogen plus progestin clearly increases the incidence of breast cancer, and that the Women’s Health Initiative showed that the distribution of tumor stage was comparable in the estrogen plus progestin arm and the placebo arm [21], treatment with MPA before diagnosis does not appear to reduce metastatic potential. These data combined then do not provide strong evidence that merely the dose or frequency of MPA administration could account for the heterogeneity among the Swedish trials. Furthermore, these data clearly do not support MPA as an agent to reduce recurrence and improve survival.

To conclude, given the divergent results from the randomized trials, we recommend caution. Symptom management remains a challenge in some postmenopausal women. The consensus statement suggesting that other established means of controlling symptoms or preventing osteoporosis should be utilized before considering estrogen and progestin therapy in these women remains prudent [22].

### Competing Interests
The author(s) declare that they have no competing interests.

### References