Management of Menopausal Symptoms

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Most menopausal women experience vasomotor symptoms with bothersome symptoms often lasting longer than one decade. Hormone therapy (HT) represents the most effective treatment for these symptoms with oral and transdermal estrogen formulations having comparable efficacy. Findings from the Women’s Health Initiative and other recent randomized clinical trials have helped to clarify the benefits and risks of combination estrogen–progestin and estrogen-alone therapy. Absolute risks observed with HT tended to be small, especially in younger women. Neither regimen increased all-cause mortality rates. Given the lower rates of adverse events on HT among women close to menopause onset and at lower baseline risk of cardiovascular disease, risk stratification and personalized risk assessment appear to represent a sound strategy for optimizing the benefit–risk profile and safety of HT. Systemic HT should not be arbitrarily stopped at age 65 years; instead treatment duration should be individualized based on patients’ risk profiles and personal preferences. Genitourinary syndrome of menopause represents a common condition that adversely affects the quality of life of many menopausal women. Without treatment, symptoms worsen over time. Low-dose vaginal estrogen represents highly effective treatment for this condition. Because custom-compounded hormones have not been tested for efficacy or safety, U.S. Food and Drug Administration (FDA)–approved HT is preferred. A low-dose formulation of paroxetine mesylate currently represents the only nonhormonal medication FDA-approved to treat vasomotor symptoms. Gynecologists and other clinicians who remain abreast of data addressing the benefit–risk profile of hormonal and nonhormonal treatments can help menopausal women make sound choices regarding management of menopausal symptoms.

VOL. 126, NO. 4, OCTOBER 2015

OBSTETRICS & GYNECOLOGY

VOL. 126, NO. 4, OCTOBER 2015

OBSTETRICS & GYNECOLOGY

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Financial Disclosure

Dr. Kaunitz has served on the advisory boards for Actavis, Bayer, and Teva. His institution receives financial support for clinical trials from Bayer and TherapeuticsMD. Dr. Manson did not report any potential conflicts of interest.

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ISSN: 0029-7844/15

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disruption, and palpitations. During a hot flush, skin temperature rises as a result of peripheral vasodilation, particularly in the fingers and toes. Along with peripheral vasodilation and sweating, heart rate increases of 7–15 beats per minute occur. It may take 30 minutes or longer for the skin temperature to return to baseline.²

The pathophysiology of vasomotor symptoms remains poorly characterized. However, these symptoms are known to be caused by estrogen withdrawal, result in a drop in core body temperature,⁴ and are linked with pulsatile pituitary luteinizing hormone secretion as well as decreased endorphin concentrations in the hypothalamus. Vasomotor symptoms are associated with an increased release of norepinephrine, serotonin, and other neuroregulators. These neurotransmitters lower a set point in the hypothalamic thermoregulatory nucleus, triggering inappropriate heat loss.⁵⁻⁶

The large prospective cohort Study of Women’s Health Across the Nation found that vasomotor symptoms are most likely to be reported by women during the late perimenopausal transition (overall prevalence approximately 70%) and are more prevalent among African American women, those with higher body mass indices, less education, lower income, mood disorders, and women who are cigarette smokers.⁷ Surgical or induced menopause is associated with a greater prevalence and severity of vasomotor symptoms. The prevalence of reported vasomotor symptoms varies substantially among women from different cultures; for instance, Asian women are less likely to report such symptoms.⁸ This observation may reflect reporting bias related to cultural norms regarding what is acceptable to report.⁹

**Duration of Vasomotor Symptoms**

Understanding the duration of bothersome vasomotor symptoms informs clinical management. The population-based prospective Penn Ovarian Aging Study enrolled older reproductive-aged women with regular menstrual cycles and followed them over 16 years. The mean age of participants was 42.2 years at baseline and 51.5 years at the time of the final menstrual period. A key finding was that the median duration of hot flushes reported by women to be moderate to severe was 10.2 years. Among women whose symptoms began around the time of onset of the perimenopausal transition, the median duration of symptoms was 11.6 years. Similar to the Study of Women’s Health Across the Nation report, the Penn Ovarian Aging Study found that the prevalence of vasomotor symptoms was greater among African American and obese women.¹⁰⁻¹¹

**Behavioral Treatment of Vasomotor Symptoms**

Common-sense measures to address vasomotor symptoms include portable fans; lowering the ambient temperature; wearing layered clothing; avoiding tobacco, alcohol, caffeine, and spicy foods, as well as ingesting cool drinks.¹¹ A randomized controlled trial (RCT) of 187 symptomatic menopausal women found that clinical hypnosis was associated with a 74.2% reduction in hot flushes compared with a 17.1% reduction in women randomized to structured attention control (P < .001).¹² The effectiveness of acupuncture,¹³ exercise,¹⁴ yoga,¹⁵ paced respiration,¹⁶ relaxation training,¹⁷ and mindfulness-based stress reduction¹⁸ has not been established.

**Over-the-Counter Products**

Over-the-counter herbal formulations, including soy, black cohosh, and other botanicals, have been widely used to treat vasomotor symptoms. In a 1-year RCT of 341 symptomatic perimenopausal or postmenopausal women, these herbal agents were not found to be more effective than placebo, whereas the efficacy of conjugated equine estrogens at 0.625 mg was demonstrated.¹⁹ A meta-analysis of 30 placebo-controlled trials of phytoestrogens (which include genistein, daidzein, found in soybeans and red clover) concluded that these agents were not effective in treating vasomotor symptoms.²⁰ A recent randomized trial of omega-3 fatty acid supplements showed no efficacy for vasomotor symptom management.²¹

**HORMONAL PRESCRIPTION MEDICATIONS**

**Systemic Hormone Therapy**

Numerous RCTs have demonstrated that estrogen represents the most effective treatment for menopausal vasomotor symptoms and related issues including impaired sleep, irritability, and decreased quality of life.²² Higher doses are associated with enhanced efficacy. Oral and transdermal estrogen formulations have comparable efficacy in treating menopausal symptoms.²³ With the exception of the 0.14-mg ultralow-dose estradiol (E2) patch, all systemic estrogen formulations are approved for treatment of vasomotor symptoms.²⁴

In women with an intact uterus, treatment with estrogen alone is associated with an elevated risk of endometrial neoplasia with dose and duration of treatment directly related to the magnitude of this risk.²⁴ When adequate progestogen is combined with estrogen, risk of endometrial neoplasia is not higher than in untreated women. In the Women’s Health Initiative clinical trial, at a mean of 5.6 years of follow-up,
use of continuous oral conjugated equine estrogens at 0.625 mg plus medroxyprogesterone acetate at 2.5 mg daily was associated with a risk of endometrial cancer similar to placebo (hazard ratio [HR] 0.81, 95% confidence interval [CI] 0.48–1.36), and risk was significantly reduced after 13 years of cumulative follow-up. A large, long-term Finnish observational study noted that although continuous concomitant progestin therapy is associated with a risk of endometrial neoplasia lower than seen in women not using hormone therapy (HT), sequential progestin therapy (eg, 14 days each month) was associated with an elevated risk. Long-cycle progestin therapy (eg, 14 days every 3 months) was associated with a further elevation in risk of endometrial cancer. Neither route nor type of progestin appeared to affect the risk of endometrial neoplasia, although additional research is needed on nonmedroxyprogesterone formulations. Based on these observations, combination estrogen–progestin therapy is generally used when systemic HT is prescribed to women with an intact uterus.

Progestational formulations are also effective in treating vasomotor symptoms, but not as effective as estrogen. A Cochrane Review found that combination estrogen–progestin HT is more effective than estrogen alone in treating vasomotor symptoms. No long-term studies address the safety of progestogen-only treatment of menopausal symptoms. Use of progestogen-only therapy to treat vasomotor symptoms is further discussed later (see “Use of Hormone Therapy in Special Patient Populations”).

In 2013, a formulation combining 0.45 mg conjugated equine estrogens with the estrogen agonist–antagonist bazedoxifene (20 mg) was approved by the U.S. Food and Drug Administration (FDA) for the treatment of vasomotor symptoms and the prevention of osteoporosis among women with an intact uterus. In a 12-month trial, rates of endometrial hyperplasia and amenorrhea among women allocated to this combination formulation were similar to those randomized to placebo.

Although clinical trials of conjugated equine estrogens–bazedoxifene have not suggested that this formulation increases the risk of breast cancer, trials have not been large or long enough to adequately assess this association. In contrast with estrogen–progestin therapy, which increases mammographic breast density and breast tenderness, bazedoxifene has not been found to increase breast density or tenderness.

In the United States, the most commonly used oral estrogens for systemic treatment of menopausal symptoms are E2 and conjugated equine estrogens. Standard doses of oral E2 and conjugated equine estrogens are 1.0 mg and 0.625 mg, respectively. Higher and lower doses of these formulations are also available.

Transdermal E2 formulations are also available. Skin patches represent the most commonly used transdermal E2 formulations. Patches releasing 0.0375 mg and 0.05 mg E2 daily are considered standard dose. Patches with higher and lower doses are also available. Ultralow-dose patches, which release 0.025 mg or 0.014 mg E2 daily, are also available. In addition, transdermal E2 gels, emulsion, and spray are available in a variety of doses. A 3-month vaginal E2-releasing ring can also be used for systemic estrogen therapy: the ring releasing 0.05 mg E2 daily is considered the standard dose.

Although estrogen injections as well as estrogen pellets have been used to treat menopausal symptoms, serum E2 levels associated with use of these formulations have not been well-characterized. Accordingly, use of oral or transdermal estrogen formulations is recommended. Estrogen pellets are not FDA-approved (see “Custom-Compounded Bioidentical Hormone Therapy”).

Common adverse effects of estrogen include breast tenderness, bloating, and uterine bleeding. Although lowering the dose may reduce these adverse effects, this may also result in reduced relief of vasomotor symptoms. Some clinicians encourage menopausal women initiating systemic HT to start with a standard dose, whereas others recommend starting therapy using a lower dose and titrate upward. Either way, it is appropriate to use the lowest-effective dose for the shortest period of time that allows the individual patient to achieve her goals. The indications and contraindications for the use of menopausal HT and related medications are summarized in Box 1.

Few menopausal women using systemic HT benefit from monitoring serum estrogen levels. However, when postmenopausal patients report inadequate symptom relief despite the use of escalating doses of estrogen, monitoring serum E2 levels can be useful. Compared with the fluctuating serum levels associated with oral estrogen, the use of transdermal E2 results in a relatively steady level of serum E2 and facilitates the monitoring of serum estrogen levels. In general, serum E2 levels 40–100 pg/mL represent a reasonable target range when monitoring serum estrogen levels in women using transdermal E2. Our clinical experience suggests that in young highly symptomatic postmenopausal women, E2 levels below this range may not result in adequate symptom relief. Medical and iatrogenic conditions that can cause heat intolerance or the sensation of flushing.
and can masquerade as hot flushes are listed in Box 2.

In the United States, the progestational agents most commonly prescribed to menopausal women with intact uteri using systemic HT are medroxyprogesterone acetate (available in 2.5-mg, 5-mg, and 10-mg doses) and micronized progesterone (available in 100-mg and 200-mg doses). Because micronized progesterone is formulated with peanut oil, it should not be used by women with peanut allergies. With standard-dose estrogen, an appropriate dose of medroxyprogesterone acetate is 2.5 mg daily for continuous therapy, or 5 mg daily with sequential therapy. An appropriate dose of micronized progesterone is 100 mg daily for continuous therapy or 200 mg nightly for sequential therapy. Because micronized progesterone at either dose can cause hypnotic effects, this medication should be taken at bedtime.

**Box 1. Menopausal Hormone Therapy and Related Medications: Indications and Contraindications**

**Systemic Estrogen Therapy (oral, transdermal, and high-dose vaginal formulations)**

- **Indications**: management of moderate-to-severe vasomotor symptoms (prevention of osteoporosis among women at high risk of osteoporotic fracture who are unable to tolerate standard preventive medications)
- **Contraindications**:
  - **Absolute**: unexplained vaginal bleeding; liver dysfunction or disease; history of deep venous thrombosis or pulmonary embolism; known blood clotting disorder or thrombophilia (transdermal may be an option in some women at elevated risk for venous thrombosis); untreated hypertension; history of breast, endometrial cancer, or other estrogen-dependent neoplasia; known hypersensitivity to hormone therapy, or history of CHD, stroke, or TIA. Concomitant progestational therapy should be prescribed when a uterus is present.
  - **Relative**: high triglycerides (higher than 400 mg/dL, 4.5 mmol/L) or gallbladder disease (oral estrogen should be avoided but transdermal estrogen may be an option); elevated risk of breast cancer (5-year breast cancer risk greater than 5% by NCI or IBIS assessment: see http://www.cancer.gov/bcrisktool/).

**Conjugated estrogen and bazedoxifene**

- **Indications**: same indications as previously stated: an additional FDA-approved option for women with concerns about breast tenderness, breast density, or uterine bleeding with conventional therapy
- **Contraindications**: absolute and relative: same as above

**Low-Dose Vaginal Estrogen**

- **Indications**: treatment of genitourinary symptoms of menopause, including vaginal dryness and dyspareunia
- **Contraindications** (absolute): unexplained vaginal bleeding; known or suspected breast cancer or endometrial cancer, or other estrogen-dependent neoplasia. For some women with a personal history of breast cancer, use might be considered after consultation with the patient’s oncologist; however, particular caution is appropriate among women taking aromatase inhibitors. Although concomitant progestational therapy is not recommended, endometrial evaluation should be performed if any vaginal spotting or bleeding occurs.

**Ospemifene**

- **Indications**: treatment of genitourinary symptoms of menopause, including vaginal dryness and dyspareunia, among women preferring an oral treatment

**Box 2. Medical and Iatrogenic Conditions That Can Masquerade as Hot Flushes**

- Anxiety disorders
- Autoimmune disorders
- Carcinoid syndromes
- Diabetic autonomic dysfunction or hypoglycemia
- Epilepsy
- Infection
- Insulinoma or pancreatic tumors
- Leukemia or lymphoma
- Mast-cell disorders
- New-onset hypertension
- Thyroid disease
- Tuberculosis
- Use of selective reuptake inhibitors or serotonin norepinephrine reuptake inhibitors

In some menopausal women, dysphoria may result from use of progestins. Changing to a different progestin, lowering the progestin dose, prescribing progestin sequentially (including long-cycle), or initiating antidepressant therapy may be useful in such patients. Although off label, use of a progestin-releasing intrauterine device provides endometrial suppression while exposing the central nervous system to lower levels of progestin than those achieved with oral or transdermal administration.


### Systemic Hormone Therapy Safety

Findings from the Women’s Health Initiative and other recent randomized clinical trials have helped to clarify the benefits and risks of combination estrogen–progestin and estrogen-alone therapy. A 2013 Women’s Health Initiative report included a comprehensive overview of findings from the two HT trials with extended post-intervention follow-up (median 13 years of cumulative follow-up) and with results stratified by age. Importantly, absolute risks of adverse events related to HT were much lower for younger women (ages 50–59 years) than for older women, as shown in Figure 1.

Results for chronic disease outcomes in the Women’s Health Initiative HT trials are summarized for the overall cohort during the intervention phase and at 13 years of cumulative follow-up in Table 1. Although HT was associated with a complex pattern

#### Table 1: Chronic Disease Outcomes in the Women’s Health Initiative HT Trials

<table>
<thead>
<tr>
<th>Condition</th>
<th>Aged 50–59 years</th>
<th>Aged 60–69 years</th>
<th>Aged 70–79 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronary heart disease</td>
<td>Active 23/17</td>
<td>Active 37/37</td>
<td>Active 62/63</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>Placebo 15/27</td>
<td>Placebo 45/36</td>
<td>Placebo 56/41</td>
</tr>
<tr>
<td>Stroke</td>
<td>Active 10/10</td>
<td>Active 23/23</td>
<td>Active 63/50</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>Active 11/15</td>
<td>Active 19/11</td>
<td>Active 10/12</td>
</tr>
<tr>
<td>Colorectal cancer</td>
<td>Active 4/5</td>
<td>Active 13/5</td>
<td>Active 15/28</td>
</tr>
<tr>
<td>Hip fracture</td>
<td>Active 1/3</td>
<td>Active 9/12</td>
<td>Active 34/48</td>
</tr>
<tr>
<td>All-cause mortality</td>
<td>Active 21/31</td>
<td>Active 51/47</td>
<td>Active 106/102</td>
</tr>
</tbody>
</table>

#### Table 2: Chronic Disease Outcomes in the Women’s Health Initiative HT Trials

<table>
<thead>
<tr>
<th>Condition</th>
<th>Aged 50–59 years</th>
<th>Aged 60–69 years</th>
<th>Aged 70–79 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronary heart disease</td>
<td>Active 17/28</td>
<td>Active 61/63</td>
<td>Active 97/90</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>Placebo 24/29</td>
<td>Placebo 28/39</td>
<td>Placebo 32/37</td>
</tr>
<tr>
<td>Stroke</td>
<td>Active 16/17</td>
<td>Active 51/33</td>
<td>Active 32/37</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>Active 10/9</td>
<td>Active 17/10</td>
<td>Active 14/16</td>
</tr>
<tr>
<td>Colorectal cancer</td>
<td>Active 7/10</td>
<td>Active 16/19</td>
<td>Active 33/15</td>
</tr>
<tr>
<td>Hip fracture</td>
<td>Active 4/1</td>
<td>Active 5/12</td>
<td>Active 38/38</td>
</tr>
<tr>
<td>All-cause mortality</td>
<td>Active 29/40</td>
<td>Active 78/77</td>
<td>Active 156/129</td>
</tr>
</tbody>
</table>

Fig. 1. Women’s Health Initiative hormone therapy trials: absolute risks (cases per 10,000 person-years) for outcomes in the intervention phases of the estrogen–progestin and estrogen-alone trials by age group. CEE, conjugated equine estrogens; MPA, medroxyprogesterone acetate. Modified from Manson JE, Chlebowski RT, Stefanick ML, Aragaki AK, Rossouw JE, Prentice RL, et al. Menopausal hormone therapy and health outcomes during the intervention and extended post-stopping phases of the Women’s Health Initiative randomized trials. JAMA 2013;310:1353–68. Copyright © 2013 American Medical Association. All rights reserved.

of benefits and risks in both trials, absolute risks (increase or decrease in number of cases per 10,000 women treated per year) tended to be small. The HR for coronary heart disease during the intervention phase was 1.18 (95% CI 0.95–1.45) in the conjugated equine estrogens plus medroxyprogesterone acetate trial and 0.94 (0.78–1.16) in the conjugated equine estrogens-alone trial; neither demonstrate a statistical significance. In both trials, women in the treatment arm had reduced risks of vasomotor symptoms, hip fractures, and diabetes and increased risks of stroke, venous thromboembolism, and gallbladder disease compared with women receiving placebo. Results for breast cancer were divergent in the two trials with increased risk in the conjugated equine estrogens plus medroxyprogesterone acetate trial and borderline reduced risk in the conjugated equine estrogens-alone trial during the intervention period. Rates of dementia (in women aged 65 years or older) were also increased in the conjugated equine estrogens plus medroxyprogesterone acetate trial, but not in the conjugated equine estrogens-alone trial.

During the poststopping phase, most risks and benefits dissipated, although some elevation in breast cancer risk persisted in the conjugated equine estrogens plus medroxyprogesterone acetate trial (cumulative HR over 13 years 1.28, 95% CI 1.11–1.48), whereas a significantly reduced risk of breast cancer (HR 0.79, 95% CI 0.65–0.97) emerged over this time period in the conjugated equine estrogens-alone trial. To put the elevated risk of breast cancer observed among women randomized to conjugated equine estrogens plus medroxyprogesterone acetate in perspective, the attributable risk with combination HT is less than one

### Table 1. Health Outcomes (Hazard Ratios [95% Confidence Intervals]) of Menopausal Hormone Therapy in the Women’s Health Initiative Trials*

| Outcome | Conjugated Equine Estrogens (0.625 mg/d) Plus Medroxyprogesterone Acetate (2.5 mg/d) | | | | | |
|---------|-----------------------------------------------------------------------------------|---|---|---|---|
|         | Intervention† | P | Cumulative Follow-up‡ | P |
| Cardiovascular disease | | | | | |
| Coronary heart disease | 1.18 (0.95–1.45) | .13 | 1.09 (0.96–1.24) | .19 |
| Stroke | 1.37 (1.07–1.76) | .01 | 1.16 (1.00–1.35) | .06 |
| Coronary artery bypass graft or percutaneous coronary intervention | 0.95 (0.78–1.16) | .64 | 1.04 (0.92–1.18) | .50 |
| Deep vein thrombosis | 1.87 (1.37–2.54) | <.001 | 1.24 (1.01–1.53) | .04 |
| Pulmonary embolism | 1.98 (1.36–2.87) | <.001 | 1.26 (1.11–1.48) | <.001 |
| Cardiovascular deaths | 1.05 (0.76–1.45) | .77 | 0.97 (0.83–1.14) | .73 |
| Cancer | | | | | |
| Invasive breast cancer | 1.24 (1.01–1.53) | .04 | 1.28 (1.11–1.48) | <.001 |
| Colorectal cancer | 0.62 (0.43–0.89) | .009 | 0.80 (0.63–1.01) | .06 |
| Endometrial cancer | 0.83 (0.49–1.40) | .49 | 0.67 (0.49–0.91) | .01 |
| All cancer types§ | 1.02 (0.91–1.15) | .69 | 1.04 (0.96–1.12) | .33 |
| Other endpoints | | | | | |
| Hip fracture | 0.67 (0.47–0.95) | .03 | 0.81 (0.68–0.97) | .02 |
| All-cause mortality | 0.97 (0.81–1.16) | .76 | 0.99 (0.91–1.08) | .87 |
| Vasomotor symptoms¶ | 0.36 (0.27–0.49) | <.001 | NR | |
| Diabetes§ | 0.81 (0.70–0.94) | .005 | 1.02 (0.93–1.12) | .66 |
| Gallbladder disease | 1.57 (1.36–1.80) | <.001 | NR | |
| Global index** | 1.12 (1.02–1.24) | .02 | 1.06 (1.00–1.13) | .05 |

NA, not applicable; NR, not reported.
† Median follow-up was 5.6 years.
‡ Median follow-up was 13.0 years.
§ Median follow-up was 7.2 years.
¶ In symptomatic women aged 50–54 years.
# Self-reported in women who did not report the condition at baseline.
** The global index of risks and benefits includes the primary endpoints of coronary heart disease and invasive breast cancer plus other outcomes including stroke, pulmonary embolism, endometrial cancer (for conjugated equine estrogen plus medroxyprogesterone acetate group), colorectal cancer, hip fracture, and all-cause mortality.

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additional case of breast cancer diagnosed per 1,000 users of combination HT annually. Regarding risk of breast cancer, another way to put this elevated HR in perspective is to recognize that a HR of 1.28 is slightly higher than that seen with one daily glass of wine and less than that noted with two daily glasses.\textsuperscript{35}

In the conjugated equine estrogens plus medroxyprogesterone acetate trial, a nonsignificantly elevated risk of ovarian cancer also was identified over 13 years of cumulative follow-up (94 cases; HR 1.24, 95% CI 0.83–1.87), whereas statistical power was too low to assess this outcome in the conjugated equine estrogens-alone trial. In general, results for chronic disease outcomes tended to be more favorable for conjugated equine estrogens-alone than for conjugated equine estrogens plus medroxyprogesterone acetate. Neither regimen affected all-cause mortality rates.\textsuperscript{26}

In addition to lower absolute risks of adverse events on HT in the younger compared with older women, age and time since menopause appeared to influence several of the HRs observed in the trials. In the conjugated equine estrogens-alone trial, women aged 50–59 years had more favorable results for all-cause mortality, myocardial infarction, colorectal cancer, and the global index (nominal P values for trend by age in the HRs were <.05 for each). In the conjugated equine estrogens plus medroxyprogesterone acetate trial, an elevated risk of myocardial infarction with treatment was apparent only in women at least a decade past menopause onset.\textsuperscript{26} Both HT regimens, however, were associated with increased risks of stroke, venous thrombosis, and gallbladder disease with no clear differences in HRs by age for these outcomes. For conjugated equine estrogens plus medroxyprogesterone acetate, an increased risk of breast cancer was an additional adverse effect in all age groups. However, the lower absolute risks of adverse events in younger women (Fig. 1), together with the generally more favorable HRs for many outcomes in the younger women, resulted in much lower rates of adverse events attributable to HT in the younger compared with older age groups.

Regarding coronary heart disease, the potential influence of age or time since menopause on the vascular response to HT has received considerable attention.\textsuperscript{22,36–39} It has been postulated that estrogen may slow early stages of atherosclerosis but has plaque-destabilizing and other adverse effects on advanced atherosclerotic lesions.\textsuperscript{36,39} In the Women’s Health Initiative, the findings that women with elevated low-density lipoprotein cholesterol concentrations, other dyslipidemias, or metabolic syndrome, but not those without these risk factors, had increased risks of coronary events while using HT\textsuperscript{40–43} are consistent with this hypothesis. Additional support for the “critical window” hypothesis...
derivates from a randomized trial of women aged younger than 50 years at enrollment, in which 10 years of combination estrogen–progestogen was associated with reduction of a composite endpoint that included cardiovascular disease and preliminary findings from the Early versus Late Intervention Trial with Estradiol showing an effect of estrogen in slowing atherosclerosis progression among women in early, but not late, menopause. However, the former trial was designed to assess osteoporosis and has been criticized for a non-blinded design, a composite outcome that included heart failure, and low statistical power. Also, a recent primary prevention trial of newly menopausal women (The Kronos Early Estrogen Prevention Study) did not demonstrate a difference in progression of atherosclerosis by noninvasive imaging after 4 years of therapy. Thus, the “critical window” hypothesis, although supported by extensive evidence, remains controversial.

Given the lower rates of adverse events on HT among women close to menopause onset and at lower baseline risk of cardiovascular disease, the use of risk stratification (categorizing individuals by their clinical characteristics and risk factor status) and personalized risk assessment has been proposed to improve the benefit–risk profile and safety of HT. One approach to decision-making from the NAMS is presented in an algorithm in Figure 2 and is available in a free mobile application (app) called MenoPro (dual mode for clinicians and patients, downloadable on iPhone, iPad, and Android devices). This approach includes assessment of whether the patient has moderate-to-severe menopausal symptoms, incorporation of the patient’s preferences regarding hormonal compared with nonhormonal therapy, and evaluation of the presence of contra-indications to HT as well as the patient’s time since menopause onset and baseline risks of atherosclerotic cardiovascular disease (using an internal risk calculator that incorporates information on the patient’s age, smoking status, systolic blood pressure level, antihypertensive therapy, presence or absence of diabetes, and total and high-density lipoprotein cholesterol measurements; also available at http://www.imedicalapps.com/2014/04/ascvd-risk-estimator-app/).

The algorithm and app also address the patient’s risk of breast cancer, osteoporosis, and other conditions relevant to clinical decision making about HT initiation as well as duration of treatment. Information about treatment options, including regimens and doses, is also provided.

Use of Estrogen Therapy in Women With Prior Hysterectomy

In the 1990s, more than 90% of women in their 50s who had undergone hysterectomy used estrogen therapy. In contrast, subsequent to the initial publication of the Women’s Health Initiative findings in 2002, this proportion has fallen to less than one third. Investigators used modeling to estimate the health effects of the decline in use of estrogen therapy posthysterectomy, concluding that this decline may have increased mortality in this subgroup of U.S. women. When counselling symptomatic women in their 50s (or younger) after hysterectomy, clinicians should clarify that for most such women, benefits of estrogen therapy are likely to outweigh risks.

Systemic Hormone Therapy: Dose Tapering, Discontinuation, and Extended Duration Use

Although common in clinical practice, extended (eg, more than 5 years) use of systemic HT is controversial because little clinical trial data address this issue. Although some have suggested that systemic HT should not be prescribed for more than 2 or 3 years, given that the mean duration of bothersome symptoms is more than one decade, this advice will not address the need for symptomatic treatment in many menopausal women.

The following recommendations are drawn from our clinical experience: Several years after initiating treatment using standard dose HT, it is reasonable to encourage the patient to try a lower dose with the reassurance that should bothersome symptoms occur, she can resume the prior higher dose without having to return for an office visit. This approach can continue until the patient is using a low or ultralow dose of HT (eg, oral E2 at 0.5 mg, conjugated equine estrogens at 0.3 mg, or transdermal E2 at 0.014 mg–0.025 mg).

After remaining free of symptoms on low-dose HT, many women may choose to try stopping HT. However, others, particularly those at elevated risk for osteoporosis (eg, lean women and those with a maternal or paternal history of osteoporosis) as well as those who perceive an enhanced overall quality of life with use of systemic HT may choose to continue low-dose therapy.

In 2012, the American Geriatric Society updated its list of medications that should be used with caution. This has resulted in physicians prescribing systemic HT to women older than age 65 years receiving notices denying reimbursement for such therapy. The American College of Obstetricians and Gynecologists (the College) recommends against arbitrarily discontinuing systemic HT at age 65 years, but
instead encouraging that treatment decisions be "...individualized based on each woman’s individual risk–benefit ratio and clinical presentation." Although randomized trials have not compared risk of thromboembolic disease with oral and transdermal HT formulations, observational studies suggest that the risk of thrombosis is lower with transdermal HT. In addition, in contrast with oral HT, transdermal HT does not appear to increase the risk of thrombosis in women with thrombophilic conditions.

Transdermal estrogen formulations represent an appropriate choice for older menopausal women, particularly those who are overweight or obese. Likewise, extended-duration use of HT represents a more attractive option when estrogen-only therapy can be used.

**Custom-Compounded Bioidentical Hormone Therapy**

Although most often used to describe custom-made HT formulations compounded according to a clinician’s prescription, the term bioidentical hormone refers to hormones identical to those made by the ovaries. Recently, use of compounded HT has increased among U.S. women with as many as 2.5 million U.S. women using these formulations, a trend propelled by advocacy from celebrities.

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**Fig. 2.** Algorithm for menopausal symptom management developed in collaboration with the North American Menopause Society and available in a free mobile app called MenoPro (dual mode for clinicians and patients). GSM, genitourinary syndrome of menopause; HT, hormone therapy; CVD, cardiovascular disease; SSRIs, selective serotonin reuptake inhibitors; SNRIs, serotonin-norepinephrine reuptake inhibitors; ACC, American College of Cardiology; AHA, American Heart Association. *Contraindications are listed in Box 1. Modified from Manson JE, Ames JM, Shapiro M, Gass ML, Shifren JL, Stuenkel CA, et al. Algorithm and mobile app for menopausal symptom management and hormonal/non-hormonal therapy decision making: a clinical decision-support tool from the North American Menopause Society. Menopause 2015;22:247–53.

than three fourths of women using compounded HT are not aware these formulations are not evaluated or approved by the FDA.\textsuperscript{55} Prescription of compounded HT often follows salivary hormone testing, which does not reliably assess serum hormone levels.\textsuperscript{57} Most women using compounded HT are not aware that FDA-approved oral, transdermal, and vaginal E2 as well as oral progesterone formulations are in fact bio-identical. Because custom-compounded hormones have not been tested for efficacy or safety by the FDA, and dose standardization and purity are uncertain, the College indicates that FDA-approved HT is preferred.\textsuperscript{23} Occasionally, when a specific FDA-approved formulation is not available, it may be appropriate to prescribe compounded HT.

**Use of Estrogen-Only Therapy in Women With an Intact Uterus**

Some menopausal women with an intact uterus wishing to use systemic HT would like to avoid concomitant use of a progestogen. One option for such women would be the use of the formulation combining oral conjugated equine estrogens with bazedoxifene. Another option, rarely used in the United States, is estrogen-only therapy with an intact uterus. In a 2-year trial, British investigators randomized menopausal women with an intact uterus to an estrogen-only regimen, oral E2 valerate at 2 mg (the equivalent of approximately 1.5 mg oral micronized E2),\textsuperscript{58} or placebo. As with the longer-term Women’s Health Initiative trial, the British study found that 2 years of estrogen did not increase risk for breast cancer.\textsuperscript{59}

Concerns regarding endometrial cancer prevent clinicians from routinely recommending estrogen-only therapy for symptomatic menopausal women with an intact uterus. In the British trial, with 2 years of unopposed estrogen therapy, and a protocol that used endometrial biopsy with short-term cyclical progestin treatment in response to any reported bleeding, no increased risk of endometrial cancer was observed.\textsuperscript{59} Other placebo-controlled trials of menopausal women with an intact uterus who received 2–3 years of estrogen therapy (either E2 at 1 mg or conjugated equine estrogens at 0.625 mg) observed no cases of endometrial cancer in women randomized to estrogen therapy.\textsuperscript{60,61} However, the incidence of endometrial hyperplasia is high in women taking unopposed estrogen. In a 3-year trial, the incidence of simple, complex, and atypical hyperplasia was 27.7%, 22.7%, and 11.8%, respectively, among women randomized to estrogen only, whereas the incidence of each of these types of hyperplasia was less than 1% among those randomized to placebo.\textsuperscript{60} Disadvantages of prescribing estrogen-only therapy to women with an intact uterus include uterine bleeding, the need for endometrial surveillance with vaginal ultrasonography–endometrial biopsy, and the recognition that some women will likely undergo hysterectomy should hyperplasia with atypia be found. Given these issues and the fact that combination therapy currently represents the standard of care for treatment of symptomatic women with a uterus,\textsuperscript{62} few clinicians and women will choose estrogen-only therapy when a uterus is present. However, some symptomatic women seeking hormonal management may be more concerned about the elevated risk of breast cancer with combination estrogen–progesterin therapy than the potential for bleeding or endometrial hyperplasia with estrogen-only therapy. For such women who also decline or are not candidates for conjugated estrogens–bazedoxifene, short-term estrogen-only therapy with appropriate endometrial surveillance represents an appropriate option. Because endometrial cancer can occur in women after endometrial ablation,\textsuperscript{62} estrogen therapy should be accompanied by post-gestational endometrial suppression when women use HT after ablation.

**Testosterone**

Used concomitantly with estrogen therapy and HT, testosterone does not appear to enhance efficacy of treatment of vasomotor symptoms.\textsuperscript{63} Although testosterone can increase sexual desire in menopausal women,\textsuperscript{64} it appears to achieve efficacy only with supraphysiologic doses, which are likely to cause hyperandrogenic adverse effects.\textsuperscript{65}

**NONHORMONAL PRESCRIPTION MEDICATIONS**

Selective serotonin reuptake inhibitors and serotonin-norepinephrine reuptake inhibitors (more commonly used to treat depression) as well as anticonvulsant medications (more commonly used to treat neuropathic pain) are increasingly being used off-label in the nonhormonal management of vasomotor symptoms. In RCTs, paroxetine, escitalopram, citalopram, venlafaxine, desvenlafaxine, gabapentin, and pregabalin have been effective in treating vasomotor symptoms.\textsuperscript{66} Low-dose paroxetine mesylate currently represents the only nonhormonal formulation FDA-approved for the treatment of vasomotor symptoms.

The antidepressants best studied in the treatment of vasomotor symptoms agents are paroxetine mesylate and venlafaxine. Lower doses of these agents than those used to treat psychiatric disorders are often
effective in treating hot flushes. Although these medications may cause nausea, headache, and dizziness, these side effects are less common than when the same medications are used to treat psychiatric conditions; furthermore, these side effects often subside within several weeks of initiating treatment. In two RCTs (12 and 24 weeks in duration) of 1,184 postmenopausal women, a 7.5-mg formulation of paroxetine was found to be effective in treating vasomotor symptoms. In contrast with use of higher-dose paroxetine for psychiatric indications, initial dose escalation is not needed with paroxetine at 7.5 mg; likewise, patients discontinuing this medication do not need to taper off of it.

In an 8-week double-blind RCT of 339 perimenopausal and postmenopausal women, venlafaxine at 75 mg extended-release and low-dose oral E2 (0.5 mg) demonstrated similar efficacy in treating vasomotor symptoms with both agents found to be more effective than placebo.

Tamoxifen is a prodrug that must be metabolized by the hepatic enzyme Cyp2d6 to endoxifen to exert its effect as adjuvant therapy in the treatment of receptor-positive breast cancer. Paroxetine and fluoxetine represent potent inhibitors of the hepatic enzyme Cyp2d6 and therefore should not be used by women taking tamoxifen. In contrast, venlafaxine, desvenlafaxine, gabapentin, and pregabalin do not inhibit Cyp2d6 and therefore may be appropriate nonhormonal medications for the treatment of bothersome vasomotor symptoms in women taking tamoxifen.

In RCTs, gabapentin at daily doses ranging from 900 to 2,400 mg has been found more effective than placebo in treating hot flushes. The recommended dose is 900 mg per day in three divided doses with a starting dose of 300 mg per day. Likewise, pregabalin (a newer formulation similar to gabapentin) has been found to effectively treat vasomotor symptoms. The recommended dose is 75 mg twice daily with a starting dose of 50 mg per day. Both of these medications can produce dose-related dizziness and sedation. Bellerjal, clonidine, and methylldopa represent central-acting medications that have demonstrated some efficacy in treating hot flushes in clinical trials. However, because these medications produce substantial side effects, their use in treating hot flushes is in general not recommended unless other treatments are ineffective or poorly tolerated.

**Stellate Ganglion Block**

Stellate ganglion block has been assessed in the treatment of vasomotor symptoms. A small RCT of 40 postmenopausal women compared the effects of stellate ganglion block with a sham injection. The resulting overall frequency of vasomotor symptoms was similar in the active and sham groups. However, the frequency of moderate-to-severe hot flushes as well as that of objectively measured symptoms was reduced significantly more by the active than by the sham treatment. A larger, sham-controlled RCT of this modality would be appropriate.

**GENITOURINARY SYNDROME OF MENOPAUSE**

Although use of systemic HT has dramatically declined since the initial 2002 publication of findings from the Women’s Health Initiative, some three fourths of U.S. women aged 57–74 years are sexually active. These observations underscore the importance of recognizing and treating the genitourinary syndrome of menopause.

Also known as vulvovaginal atrophy, genitourinary syndrome of menopause refers to the physical changes of the vulva, vagina, and lower urinary tract that result from estrogen deficiency and represents a common and progressive condition that adversely affects health, sexuality, and quality of life of many menopausal women. The word “atrophy” may be perceived in a negative fashion by midlife women; likewise, the media and the public may not be comfortable using the term “vagina.” Furthermore, vulvovaginal atrophy does not encompass the lower urinary tract. Accordingly, a consensus panel (with representation from the College) was convened in 2013 and recommended “genitourinary syndrome of menopause” as preferred terminology.

Because high concentrations of estrogen-receptor α are present in the vaginal epithelium of premenopausal and postmenopausal women, endogenous E2 causes the epithelium to become a thick, well-vascularized rugated surface that provides adequate lubrication for most premenopausal women. Genitourinary syndrome of menopause results from low systemic estrogen levels, which most commonly are associated with spontaneous menopause, but can also occur as a result of induced menopause or transient low estrogen states including lactation and use of gonadotropin-releasing hormone agonists.

Genitourinary syndrome of menopause may cause the vagina to shorten and narrow and the introitus may contract, particularly in the absence of penetrative sexual activity. The vaginal epithelium becomes thin and sebaceous gland secretions diminish, causing lubrication from sexual stimulation to be delayed as well as decreased. The term atrophic vaginitis refers to inflammation that occurs with this syndrome.

Although approximately one third of midlife and older women report dryness and pain with
intercourse, only a minority seek help for genitourinary syndrome of menopause. In contrast with vasomotor symptoms, the genitourinary syndrome of menopause often progresses without treatment. Because many women with this condition may not report symptoms, clinicians should enquire about symptoms, including vaginal dryness and sexual discomfort, during well-women visits.

Genitourinary syndrome of menopause can be presumptively diagnosed based on a characteristic history alone. However, performing a pelvic examination helps exclude other vulvovaginal conditions that may present with similar symptoms, including chronic infections, contact dermatitis, lichen sclerosus, lichen planus, and malignancy. During inspection of the external genitalia, the clinician may note loss of the fat pad in the mons and labia majora as well as loss of labia minora tissue and pigmentation. The urethral meatus may become erythematous and prominent. If introital or vaginal narrowing is present, use of an ultrathin speculum minimizes patient discomfort. The vaginal epithelium may appear smooth (loss of rugation), shiny, and dry. Use of a spatula or cotton-tipped swab may cause bleeding (friability), and the fornices may become attenuated, resulting in a cervix flush with the vaginal apex. Laboratory assessment is not necessary to diagnose the genitourinary syndrome of menopause. If assessed, vaginal pH is characterized typically higher than 5.0 and microscopic wet prep may reveal more white blood cells than epithelial cells, immature epithelial cells (large nuclei), and reduced or absent lactobacilli. Performed in a clinical laboratory, the vaginal maturation index assesses vaginal estrogen status by evaluating a vaginal smear for percentage of superficial parabasal cells. Estrogen-deprived vaginal epithelium will shed a higher proportion of parabasal cells.

Nonhormonal Treatment of Genitourinary Syndrome of Menopause

Many women may report having few, if any, symptoms of genitourinary syndrome or may choose to not treat their symptoms. However, when this condition is identified, clinicians should advise women that without active management, it often worsens over time. In women without severe symptoms, recommending use of over-the-counter water-based or silicone-based vaginal lubricants for sexual activity and regular use of long-acting vaginal moisturizers is appropriate. Use of vaginal lubricants reduces friction-related discomfort that women with genitourinary syndrome of menopause experience during sexual activity. Oil-based lubricants should not be used with latex condoms. Consistent use of long-acting vaginal moisturizers can reduce the vaginal pH to premenopausal levels. Women with genitourinary syndrome of menopause should also be advised that regular sexual activity may help address symptoms and prevent progression. In some women, introital–vaginal contraction or vaginismus may prevent penetration. Use of graduated vaginal dilators, sometimes facilitated by physical therapists specializing in pelvic floor disorders, will often allow such women to resume or initiate comfortable sexual activity. In this setting, initiating low-dose vaginal estrogen therapy can accelerate progress.

Treating Genitourinary Syndrome of Menopause With Systemic Hormone Therapy

In general, systemic HT is effective in treating symptoms of genitourinary atrophy. However, 10–15% of women will experience vulvovaginal symptoms during use of systemic HT; such symptoms become more likely when lower than standard doses are used. In this setting, low-dose vaginal estrogen can be added to systemic HT. Alternatively, a woman using oral or transdermal systemic HT can switch to the 3-month systemic vaginal ring (Femring, which releases 0.05 mg/d or 0.10 mg/d of E2), which will address both vasomotor as well as vaginal symptoms. The systemic vaginal E2 ring (Femring) should not be confused with the 3-month local vaginal ring (Estring), which releases 7.5 micrograms E2 per day.

Low-Dose Vaginal Estrogen Therapy in the Treatment of Genitourinary Syndrome of Menopause

Low-dose vaginal estrogen represents highly effective treatment for symptomatic genitourinary syndrome of menopause. Systematic reviews have noted that tablets, the ring, and creams have comparable efficacy in treating vulvovaginal symptoms. Vaginal estrogen has also been found to reduce risk of recurrent urinary tract infections and overactive bladder symptoms in menopausal women. The low-dose vaginal E2 ring is approved to treat urinary urgency and dysuria. In contrast, systemic HT has been noted to increase urinary incontinence.

In the United States, two low-dose E2 creams, a ring, and a tablet are available. An advantage of the creams is that, in contrast with the ring and tablets, they can be applied not only intravaginally (using graduated plastic applicators supplied with each tube of cream), but also digitally to the vestibular tissues–introitus. However, some women find creams messy. The E2 cream includes 0.1 mg active ingredient per gram. Package labeling indicates 2–4 g should
initially be applied daily for 1 or 2 weeks then gradually reduced to half of the initial dosage for a similar period. The recommended maintenance dose is 1 g one to three times weekly.

The conjugated equine estrogens cream includes 0.625 mg active ingredient per gram. Package labeling indicates 0.5–2 g should be administered cyclically (daily for 21 days then off for 7 days). In practice, during maintenance therapy, most women use vaginal cream one to three times weekly on a continuous basis.

The 3-month low-dose E2 vaginal ring releases 7.5 micrograms E2 per day. An advantage of the ring is the convenience of placement and removal only once every 3 months. A disadvantage is that some women are not comfortable inserting or removing the ring. Many women and their partners choose to leave the ring in place during intercourse. If the ring is removed, it can be rinsed (avoiding soap and hot water) and replaced as soon as possible.

Vaginal tablets contain 10.3 micrograms E2 hemihydrate, equivalent to 10 micrograms E2. Using a single-use plastic applicator, one tablet is placed intravaginally nightly for 2 weeks and then twice weekly for maintenance therapy. An advantage of the tablets is that women find them less messy than creams; in addition, the applicator is thinner than the cream applicators.

The College and NAMS recommend that low-dose vaginal estrogen therapy may be used as long as needed, including indefinitely, and do not recommend routine use of concomitant progesterin therapy for endometrial protection during use of long-term vaginal estrogen therapy. However, clinicians should recognize that endometrial safety data for currently marketed formulations are based on 1-year trials with no trials assessing safety of longer-term administration available. In the absence of reported spotting or bleeding, routine endometrial surveillance during use of vaginal estrogen therapy is not recommended.

Current package labeling for low-dose vaginal estrogen, like for all systemic HT formulations, includes a boxed warning that refers to endometrial cancer, myocardial infarction, stroke, invasive breast cancer, pulmonary emboli, and dementia. This warning reflects safety concerns associated with findings from clinical trials of systemic HT, including the Women’s Health Initiative. In 2014, a group, which included members of NAMS and the College, published an editorial calling for a change in product labeling for low-dose vaginal estrogen, pointing out that the boxed warning is not evidence-based and hurts women by discouraging use of an effective local treatment. The recommendations to change the boxed warning are based on several observations: 1) blood steroid levels are substantially lower with low-dose vaginal estrogen formulations than those seen with systemic estrogen therapy, 2) neither clinical trials nor consistent observational data suggest an elevated risk of the conditions referred to in the boxed warning, and 3) the absence of evidence suggesting that the small changes in blood hormone levels observed during use of these low-dose formulations increase risk for endometrial or breast cancer, venous thromboembolism, coronary artery disease, or cerebrovascular disease. This publication proposes modified labeling language that would better reflect the safety profile of low-dose vaginal estrogen and potentially enhance safety by focusing on crucial information regarding these formulations that would benefit women and clinicians. At the time of this article’s publication, the FDA is reviewing and considering the proposal.

**Ospemifene**

Ospemifene is a systemic estrogen agonist–antagonist that improves genitourinary syndrome of menopause. A 1-year clinical trial did not identify endometrial safety concerns. However, hot flushes were reported by 7.2% of participants randomized to ospemifene at 60 mg compared with 2.0% of those randomized to placebo. In 2013, the FDA approved ospemifene 60-mg oral tablets for the treatment of dyspareunia in postmenopausal women.

As with systemic HT and low-dose vaginal estrogen therapy, package labeling for ospemifene cautions against using this medication in women with a personal history of breast cancer or thromboembolic disease (Box 1). Published data have not addressed the safety of ospemifene in these patient populations.

**Androgens**

Dehydroepiandrosterone, an androgen derivative, is available without a prescription as a dietary supplement. Trials of intravaginal dehydroepiandrosterone have demonstrated effectiveness in treating genitourinary syndrome of menopause. Clinical trial data addressing the use of vaginal testosterone are limited; longer as well as larger studies are needed to assess efficacy and safety.

**USE OF HORMONE THERAPY IN SPECIAL PATIENT POPULATIONS**

**Women With Early Menopause**

Women who experience spontaneous or induced menopause (caused by surgery, chemotherapy, or other factors) in their 40s or younger face more severe
vasomotor symptoms than women reaching meno-
pause at the mean age (51–52 years) and are at higher
risk for osteoporosis and possibly coronary heart and
neurodegenerative disease. Although little clinical
trial data inform management of women with early
menopause, use of systemic menopausal HT or oral
contraceptives is appropriate in the absence of contra-
indications.22 When systemic HT in used in this
patient population, higher-than-standard doses are
often appropriate.77

Women With a History of Breast Cancer
In premenopausal women treated for breast cancer,
induced menopause caused by chemotherapy often
leads to bothersome vasomotor symptoms. Although
observational studies have not found that use of
systemic or low-dose vaginal HT increases recurrence
risk in breast cancer survivors, this might reflect
selection bias in that healthier women with a better
baseline prognosis might be more likely to pursue
such treatment.78,79

In 1997, two clinical trials of HT in women with
a history of breast cancer were initiated in Sweden. In
the HABITS (Hormonal Replacement Therapy After
Breast Cancer—Is It Safe?) (n=447) and Stockholm
(n=378) studies,80,81 the progestin component in most
women randomized to estrogen–progestin therapy
consisted of continuous norethindrone acetate and
cyclical (sequential) medroxyprogesterone, respect-
ively. Because a statistically elevated risk of recurrent
breast cancer was noted in the HABITS trial, both
studies stopped administering study medications in
2003. At 4 and 10.8 years median follow-up, respec-
tively, no elevated breast cancer-specific or overall
mortality was noted in the HABITS or Stockholm
studies. Hormone therapy was associated with a per-
sistent elevated risk of recurrence in the HABITS
study; in contrast, in the Stockholm study, HT was
not associated with an elevated recurrence risk at the
time study medication was stopped, or at 10.8
years follow-up.80,81 Several factors limit the clinical
applicability of these two Swedish studies, including
that no standardized regimen of HT was used and that
many participants concomitantly used tamoxifen and
HT, a practice not accepted in the United States.

Given the uncertainty regarding the effect of
systemic HT on risk of recurrence, systemic HT in
general should not be used in breast cancer survivors;
nonhormonal options represent appropriate first-line
options for treatment of vasomotor symptoms in this
patient population.82 Symptomatic breast cancer sur-
vivors considering off-label use of systemic HT should be
counseled that such therapy may increase recurrence
risk. In addition, given the findings from the Women’s
Health Initiative detailed previously in this article,
clinicians considering prescribing off-label HT to treat
bothersome menopausal symptoms in patients with
breast cancer with an intact uterus may consider using
estrogen-only therapy (see section on “Use of Estrogen-
Only Therapy in Women With an Intact Uterus”).

Genitourinary syndrome of menopause is prev-
alent among women with breast cancer, and deci-
sions regarding use of low-dose vaginal estrogen are
surrounded with controversy. In contrast with
women using systemic HT, in women using the
low-dose vaginal ring or tablets, serum estrogen
levels remain in the postmenopausal range. Although
use of vaginal estrogen cream can increase
serum estrogen levels substantially, using very small
amounts can improve symptoms without raising
serum estrogen levels above those seen in untreated
menopausal women.73,75

A large observational study found that, in healthy
menopausal women, use of low-dose vaginal estrogen
was not associated with increased risk of breast
cancer.83 Likewise, a large observational study of
breast cancer survivors found that vaginal estrogen
was not associated with an elevated risk of breast can-
cer recurrence.79 Although serum levels of estrogen
remain within the normal menopausal range in
women using low-dose vaginal estrogen, clinicians
and women should be aware that modest increases
in serum estrogen levels may occur in this setting.
Even such minimal increases may be of concern in
women using aromatase inhibitors as adjuvant treat-
ment for breast cancer. Accordingly, clinicians consid-
ering prescription of low-dose vaginal estrogen to
breast cancer survivors with symptomatic genitouri-
mary syndrome of menopause should make this deci-
sion in concert with the patient and her oncologist.75,82
The safety of ospemifene has not been assessed in
breast cancer survivors, and package labeling for this
estrogen agonist–antagonist lists known or suspected
estrogen-dependent neoplasia as a contraindication.

Carriers of BRCA 1 or BRCA 2 Mutations With
No Personal History of Breast Cancer
As more BRCA mutation carriers are being identified,
and more carriers undergo risk-reducing salpingoo-
ophorectomy and bilateral mastectomy, clinicians
and mutation carriers more frequently face decisions
regarding use of systemic HT. As a result of concerns
regarding HT and breast cancer, some mutation car-
rriers may delay or avoid salpingo-oophorectomy,
a surgery that reduces not only future risk of ovarian
cancer, but also future risk for breast cancer.84

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Mutation carriers who have undergone bilateral risk-reducing mastectomy experience a very low baseline future risk for breast cancer.\textsuperscript{84} Accordingly, concerns regarding this disease should not play a major role in decision-making regarding use of systemic HT.

Among mutation carriers with intact breasts, several studies address risk of breast cancer with use of systemic HT. A 2005 cohort study followed \textit{BRCA} 1 and 2 carriers with intact breasts, 155 of whom had undergone risk-reducing salpingo-oophorectomy, for a mean follow-up of 3.6 years. Of these women, 60\% and 7\%, respectively, of those who had and had not undergone salpingo-oophorectomy used HT. This study found that bilateral salpingo-oophorectomy reduced future breast cancer risk by some 60\%, whether or not women used HT.\textsuperscript{85} In a 2008 case-control study of 472 menopausal \textit{BRCA} 1 carriers, half had been diagnosed with breast cancer (cases) and the other half (control participants) had not received this diagnosis. Prior use of HT was associated with a 43\% reduction in risk of breast cancer.\textsuperscript{86} Finally, a 2011 presentation described a cohort study in which 1,299 \textit{BRCA} 1 and 2 carriers with intact breasts who had undergone salpingo-oophorectomy were followed for a mean of 5.4 years postoperatively. Overall, use of HT was not associated with an increased risk of breast cancer. Among women with \textit{BRCA} 1 mutations, use of systemic HT was associated with a reduced risk of breast cancer.\textsuperscript{87} In aggregate, these studies provide reassurance that short-term use of systemic HT does not increase breast cancer risk in women with \textit{BRCA} 1 or 2 mutations and intact breasts.

**Women With a History of Venous Thromboembolism**

Package labeling lists a history of venous thromboembolism as a contraindication to use of all formulations of systemic HT. However, a French observational study found that in women with a history of venous thromboembolism (26\% of whom were known to have a thrombophlic condition), oral (but not transdermal) estrogen therapy increased the risk of recurrent thrombotic events.\textsuperscript{88} Although this study provides limited support for the safety of transdermal estrogen in women with a history of venous thromboembolism, it cannot be considered definitive given its small size.

**Women With a History of Endometriosis**

In women with symptomatic endometriosis who have completed childbearing, hysterectomy with bilateral salpingo-oophorectomy may be performed. Often, such women are in their 40s or younger and face severe menopausal symptoms. In menopausal women who have undergone surgery for endometriosis and elect hormonal treatment for menopausal symptoms, some authorities recommend progestin-only or combined estrogen–progestin formulations to prevent benign or neoplastic growth of endometriosis implants. The concomitant use of continuous progestin in such patients may be particularly important if endometriosis is symptomatic or large residual volumes of endometriotic tissue remained after surgery.\textsuperscript{89,90}

**DISCUSSION**

Hormone therapy represents the most effective treatment for menopausal vasomotor and related symptoms. Among appropriate candidates, estrogen–progestogen combination HT is recommended to treat bothersome symptoms in women with an intact uterus, whereas estrogen-alone therapy can be used for symptomatic women posthysterectomy. Findings from the Women’s Health Initiative and other recent randomized clinical trials have helped to clarify the benefits and risks of combination estrogen–progestin and estrogen-alone therapy. Although combination and estrogen-alone therapy was associated with a complex pattern of benefits and risks in both Women’s Health Initiative trials, absolute risks tended to be small. Given the lower rates of adverse events on HT among women close to menopause onset and at lower baseline risk of cardiovascular disease, the use of risk stratification and personalized risk assessment represents a sound strategy for optimizing the benefit–risk profile and safety of HT. Treatment duration should be individualized based on patients’ risk profiles and personal preferences. Genitourinary syndrome of menopause represents a common condition that reduces the quality of life of many menopausal women. Low-dose vaginal estrogen is highly effective in treating this condition. Because custom-compounded hormones have not been tested for efficacy or safety, FDA-approved HT is preferred. A low-dose formulation of paroxetine mesylate currently represents the only nonhormonal medication that is FDA-approved to treat vasomotor symptoms. Gynecologists and other clinicians who remain abreast of data addressing the benefit–risk profile of HT as well as nonhormonal treatments can help menopausal women make sound choices regarding management of menopausal symptoms.

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