INTRODUCTION

This update summarizes articles and guidelines published in the last year that may impact general internists’ women’s health clinical practice (Table 1).

METHODS

We reviewed the contents of leading medical journals including: the New England Journal of Medicine, the Journal of the American Medical Association, Annals of Internal Medicine, Archives of Internal Medicine, British Medical Journal, Lancet, Obstetrics and Gynecology, American Journal of Obstetrics and Gynecology, Journal of General Internal Medicine, PLOS Medicine, American Journal of Public Health, Circulation, Diabetes, and Diabetes Care between March 1, 2010 and February 28, 2011. We also reviewed updates to the Cochrane database of systematic reviews, Guideline Clearinghouse, and the articles highlighted by the ACP Journal Club, Journal Watch, and Journal Watch Women’s Health. Finally, we performed a MEDLINE search using the medical subject heading, “sex factors.” Those abstracts rated in the top third of importance by any author were read closely and rated by all authors. A process of individual ratings and multiple discussions was then used to reach consensus about the most important articles published in the last year.

ISSUES FOR REPRODUCTIVE AGED WOMEN

Abdool Karim Q, Abdool Karim S, Frohlich J. Effectiveness and safety of tenofovir gel, an antiretroviral microbicide, for the prevention of HIV infection in women. Science. 2010;329(5996):1168–74.

What Was Known? Women are disproportionately affected by the HIV epidemic, yet have few ways to protect themselves from sexually transmitted infections. Over the last 20 years, six microbicides have been studied; none provide meaningful protection against HIV infection.

What this Study Adds This double-blind, randomized, placebo-controlled trial followed 889 HIV-negative, South African women who applied a 1% tenofovir vaginal gel 12 h before and 12 h after intercourse. Over 30 months of follow-up, the gel was found to have an acceptable safety profile and reduced HIV incidence by 39%.

How Should I Change my Practice? The US FDA has previously approved oral tenofovir and recently “fast tracked” the 1% gel, creating hope it may be available to our patients in 2012.

Lukes AS, Moore KA, Muse KN, et al. Tranexamic acid treatment for heavy menstrual bleeding: a randomized controlled trial. Obstet Gynecol. 2010;116(4):865–75.

What Was Known? Tranexamic acid inhibits the activation of plasminogen, increasing clot formation. Injectable tranexamic acid was FDA approved in 1986; an oral formulation has been safely used in Europe for the last 40 years and was FDA approved in 2009.1

What this Study Adds. In this RCT (n=196), women treated with 3.9 g/day of tranexamic acid for 5 days with each menses had significant and personally meaningful reductions in menstrual blood loss at 1 month that were consistent to 6 months. In addition, women reported decreased social and physical limitations with menses.

How Should I Change my Practice? Oral transexamic acid is a new therapeutic alternative for providing short-term relief for menorrhagia, though a prior study found a levonorgestrel IUD was more effective than transexamic acid in providing long-term relief.2

Foster D, Hulett D, Bradshberry M, Darney P, Policar M. Number of oral contraceptive pill packages dispensed and
Table 1. Important Women’s Health Guidelines in 2010–2011: New or Updated

| Topic                  | Issuing organization | Updated recommendations |
|------------------------|----------------------|-------------------------|
| Contraceptive guidelines | CDC†                 | http://www.cdc.gov/mmwr/pdf/rr/rr590528.pdf | New includes guidance after bariatric surgery, organ transplantation, and with inflammatory bowel disease |
| STD treatment guidelines | CDC†                 | http://www.cdc.gov/std/treatment/ | Beware resistant gonorrhea |
| CHD prevention         | AHA‡                 | Encourage expedited partner therapy |
| Hormone therapy        | NAMS§                | No increased risk of CHD if combined hormone therapy is initiated in women aged 50–59 or within 10 years of menopause |
| Osteoporosis screening | USPSTF∥              | Screen women aged 65 and older Screen women whose fracture risk is equal to or greater than a 65-year-old white woman who has no additional risk factors |
| Vitamin D              | Institute of Medicine | RDA 600 IU for women aged 9–70 RDA 800 IU for women over age 70 Upper limit 4,000 IU per day 25-OH Vitamin D level of 20 ng/ml is the goal |

*Centers for Disease Control and Prevention  
†American Heart Association  
‡North American Menopause Society  
§United States Preventive Services Task Force

subsequent unintended pregnancies. Obstet Gynecol. 2011;117 (3):566–72.

What Was Known? Oral contraceptive pills (OCPs) are the most commonly used form of reversible contraception in the US, and receiving a year’s worth of pills upon initiating OCP use does not reduce preventive services such as Pap or chlamydia testing.

What this Study Adds. Women who were given 12 packs of OCPs were less likely to experience an unplanned pregnancy (OR=0.70, 95% CI 0.57–0.87) or abortion (OR=0.54, 95% CI 0.32–0.93) than those given 3 or less, even after controlling for age, race/ethnicity, and prior pill use.

How Should I Change my Practice? Provide a year’s worth of pills to women interested in using an OCP regardless of other plans to provide preventive services.

Munk-Olsen T, Laursen T, Pedersen C, et al. Induced first-trimester abortion and risk of mental disorder. NEJM. 2011;364 (4):332–9.

What Was Known? First trimester induced abortion is one of the most commonly US performed surgical procedures.

What this study adds: In this study from multiple Danish registries, the investigators found that first trimester abortion did not affect rates of psychiatric contact (p=0.19) in the 9 months before or 12 months after the procedure (n=84,620). However, rates of psychiatric contact were increased after childbirth (<0.001) for 6 to 9 months; rates of psychiatric contact for affective disorder were highest in the first month postpartum (RR=3.79, 95% CI 2.86–5.02), but remained elevated 7–9 months postpartum (RR=1.69, 1.31–2.17). Similarly, rates of psychiatric contact for neurotic, stress-related, or somatoform disorders were highest 1 month postpartum and remained elevated for 7–9 months postpartum (RR=1.33 (1.11–1.60).

How should I change my practice? Women can be reassured that first trimester abortion does not increase the risk of mental disorder. Routine screening for postpartum mental disorder may be useful.

DISEASES OF EARLY ADULTHOOD INTO MID-LIFE

Borzekowski D, Schenk S, Wilson J, Peebles R. e-Ana and e-Mia: A content analysis of pro-eating disorder web sites. Am J Public Health. 2010;100(8):1526–34.

What Was Known? Eating disorders, such as anorexia (Ana) and bulimia (Mia), are highly prevalent. Adolescents who frequent pro-eating disorder websites are less satisfied with their bodies and practice disordered eating longer than peers not using these sites. Little is systematically known about the content of these sites.

What this Study Adds. This is the first large systematic review of pro-eating disorder websites. Investigators used Yahoo and Google search engines to find 180 active websites, forums, journals, or blogs focusing on pro-eating disorder information. Two reviewers independently coded the content using 64 variables organized into six themes (site logistics, site accessibility, “thinspiration” materials, recovery, perceived themes, and perceived harm). Ninety-one percent of the websites were publicly accessible and 79% were interactive, wherein users could post content or use tools (e.g., diet calculators). Pro-anorexia content was provided on 84% of websites, pro-bulimia on 64%, thinspiration images or prose to inspire weight loss on 85%, and overt suggestions for engaging in eating-disordered behavior on 83%. Only 38% of websites offered recovery advice or a link to such information. The
investigators rated the websites on perceived harm using a 5-point Likert scale, in which 83 sites were ranked medium harm (score 2 or 3, content somewhat dangerous) and 39 sites were ranked high harm (score 4 or 5, content could lead to immediate and life-threatening problems). Common themes on pro-eating disorder websites were control, success, perfection, solidarity, and a cult-like mentality of anorexia/bulimia being a gift, not a lifestyle choice for “wannabe” dieters. Photographs of emaciated models and celebrities, creeds, and “Thin Commandments” are examples of the content used to inspire and motivate disordered eating behaviors.

**How Should I Change my Practice?** Practitioners should be aware of the detrimental messages on pro-eating disorder websites. These sites are dynamic communities that often serve as venues for expression, encouragement, and support for disordered eating behaviors.

Giuliano A, Hunt K, Ballman K, et al. Axillary dissection vs. no axillary dissection in women with invasive breast cancer and sentinel node metastasis. JAMA. 2011;305(6):569–75.

**What Was Known?** Axillary lymph node dissection (ALND) has been the standard of care for breast cancer patients with sentinel lymph node (SLN) metastasis. ALND is associated with a high risk of complications, and its effect on long-term survival has not been definitively demonstrated.

**What this Study Adds.** This multicenter, noninferiority trial randomized 891 women with clinically confirmed T1 or T2 invasive breast cancer and one or two tumor-involved SLNs to SLN dissection (SLND) alone or ALND. Adjuvant systemic chemotherapy was administered per the discretion of the treating physician; chemotherapy rates were similar for both groups (97% SLND alone, 96% ALND). After a median follow-up of 6.3 years, there were no differences in 5-year overall survival rates (92.5% SLND alone, 91.8% ALND) or disease-free survival rates (83.9% SLND alone, 82.2% ALND) between the groups. Rates of wound infections, axillary seromas, and paresthesias were significantly higher in the ALND than SLND-alone group (70% versus 25%, respectively, P<0.001), as was lymphedema (P<0.001).

**How Should I Change my Practice?** Although the follow-up period in this study was short, most axillary tumor recurrence presents at 1–2 years. Therefore, internists should feel comfortable reinforcing to patients that ALND does not improve the survival of patients with ductal carcinoma and positive SNLs, and results in more frequent complications such as wound infections and lymphedema.

Schnatz P, Marakovitz K, O’Sullivan D. The association of breast arterial calcification and coronary heart disease. Obstet Gynecol. 2011;117(2):233–41.

**What Was Known?** Coronary heart disease (CHD) is the leading cause of death among women. While radiographic studies have found an increased prevalence of intimal calcifications in vessels (e.g., coronary and aorta) among patients with CHD, and breast arterial calcifications (BAC) are common on mammograms (9% overall and up to 50% in women over age 65), the clinical significance of these calcifications is unknown.

**What this Study Adds.** This 5-year prospective cohort study enrolled 1,995 women from four outpatient radiology facilities. Surveys were administered at baseline and at years 2, 4, and 5. Mammograms were reviewed by a blinded radiologist to determine the presence of BAC. At baseline, the mean age was 56.3 years, 95% were white, 61% post-menopausal, and 16.3% were BAC-positive. The BAC-positive group was more than the BAC-negative group (69 vs 54 years, p<0.001) and had a higher prevalence of most CHD risk factors (hypertension, hypercholesterolemia, diabetes, and postmenopausal). Over the 5-year study period, the prevalence of CHD was 20.8% in the BAC-positive group compared to 5.4% in the BAC-negative group (P<0.001). In the cohort without CHD at baseline, the BAC-positive group was more likely to develop CHD (6.3% vs 2.3% respectively, P=0.003). The presence of BAC was a better 5-year predictor of incident CHD than hypertension, hypercholesterolemia, or family history of CHD, independent of age and other confounders.

**How Should I Change my Practice?** The presence of BAC on mammography is likely a marker for CHD and should be routinely reported. Clinicians evaluating patients with BAC-positive mammograms should consider factoring it into their decision making regarding CHD prevention.

**MANAGEMENT OF THE MENOPAUSAL WOMAN**

Freeman E, Guthrie K, Caan B, et al. Efficacy of escitalopram for hot flashes in healthy menopausal women: a randomized controlled trial. JAMA. 2011;305(3):267–74.

**What was Known?** Hormone replacement therapy is the most effective treatment for relieving menopausal vasomotor symptoms, but its use is limited by increased rates of stroke and venous thromboembolic disease. Non-hormonal therapies, including antidepressants and neuroleptics, reduce hot flashes by 40–84%. Few studies have examined whether the effectiveness of non-hormonal therapies is influenced by race, menopausal status, or a personal history of anxiety or depression.

**What this Study Adds** Women (n=250) with moderately severe hot flashes were randomized to escitalopram 10 mg daily or placebo for 8 weeks. Almost half of the study participants were African-American, and 14–23% reported mild depression or anxiety at baseline. After 8 weeks of treatment, patients in the escitalopram group experienced 1.40 fewer hot flashes per day and a reduction in severity than those in the placebo group. Although there was a substantial placebo effect, more women in the escitalopram group experienced at least a 50% reduction...
in hot flash frequency (55% vs 36%, p=0.009). Race, menopausal status, and a history of depression or anxiety did not modify the treatment effect, and adverse events were few. Patient satisfaction and desire to continue with treatment was higher with escitalopram than placebo (70% vs 43%, p <0.001, and 64% vs 42%, p=0.005, respectively).

**How Should I Change my Practice?** Escitalopram may be a good treatment option for symptomatic women who want to avoid systemic hormone therapy, and will be effective even in those without anxiety or depression.

Renoux C, Dell'aniello S, Garbe E, Suissa S. Transdermal and oral hormone replacement therapy and the risk of stroke: a nested case-control study. BMJ. 2010;340:c2519.

**What Was Known?** Oral hormone therapy for relief of menopausal symptoms increases the risk for ischemic stroke. Transdermal hormone therapy, while equally efficacious, avoids the hepatic first-pass effect and is less likely than oral therapy to increase cardiovascular risk markers. To date, no studies have compared the effects of transdermal and oral hormone therapy on stroke risk.

**What this Study Adds** Using a large British medical record database and a nested case-control study design, the investigators examined the rates of stroke among older women who were exposed to hormone therapy. The route (oral vs transdermal) and dose (high vs low dose estrogen) of hormone therapy were compared among 15,710 cases and 59,958 controls. After adjustment for multiple cardiovascular risk factors, women who used transdermal estrogen or estrogen/progesterone hormone therapy had no increased risk of stroke compared with non-users (rate ratio 0.95, 95% CI 0.75–1.20), although this was only true with doses of estrogen less than 50 μg. Conversely, treatment with oral estrogen significantly increased the risk of stroke by 35%, an effect that was mitigated only slightly if oral estrogen was combined with progesterone (rate ratio 1.24, 95% CI 1.08–1.41). Stroke risk with oral hormone therapy did not vary significantly when stratified according to low or high estrogen doses (rate ratio 1.25, 95% CI 1.12–1.40 vs 1.48, 95% CI 1.16–1.90, respectively). In a direct comparison, treatment with transdermal hormone therapy was associated with a lower risk of stroke than oral therapy (rate ratio 0.74, 95% CI 0.58–0.95).

**How Should I Change my Practice?** This study suggests that transdermal hormone therapy is safer than oral therapy. Although case control study designs have inherent biases, there is biological plausibility for these results based on prior studies. Providers should consider selecting transdermal routes of hormone administration to minimize the risks associated with systemic therapy.

Chlebowski R, Anderson G, Gass M, et al. Estrogen plus progestin and breast cancer incidence and mortality in postmenopausal women. JAMA. 2010;304(15):1684–92.

**What was Known?** In the Women’s Health Initiative (WHI) placebo-controlled trial, women who were treated with combined estrogen and progestin (E/P) therapy had an increased risk of invasive breast cancer. Moreover, these breast cancers tended to be larger and at a more advanced stage at the time of detection. Although the risk of breast cancer declined significantly after discontinuation of E/P, the long-term effects of E/P on breast cancer incidence, characteristics at diagnosis, and breast cancer mortality are unclear.

**What this Study Adds** Although the WHI was stopped early in 2002, all subjects were followed for an additional 3 years, until the original trial completion date (“post-intervention phase”). Eighty-three percent of women consented to participate in the “extension phase,” and data regarding breast cancer incidence and mortality were collected prospectively until 2009. In total, after a mean follow-up of 11.0 years, 678 cases of breast cancer were identified. Treatment with E/P, as compared to placebo, was associated with a significantly increased incidence of invasive breast cancer (HR 1.25, 95% CI 1.07–1.46). Women in the treatment group, as compared to those receiving placebo, were more likely to have positive lymph nodes at the time of diagnosis and were more likely to die from breast cancer (25 deaths vs 12 deaths, HR 1.96, 95% CI 1.00–4.04). Overall mortality after a diagnosis of breast cancer was higher in the E/P group than in the placebo group (51 deaths vs 31 deaths, HR 1.57, 95% CI 1.01–2.48).

**How Should I Change my Practice?** Treatment with E/P increases the incidence of invasive breast cancer, and the risk persists after discontinuation of therapy. Moreover, E/P increased breast cancer mortality and overall mortality, although the number of deaths in both groups was relatively small. In light of these new data, practitioners should continue to carefully weigh the risks and benefits of E/P therapy and counsel patients appropriately.

**OSTEOPOROSIS AND BONE HEALTH**

Bolland M, Avenell A, Baron J, et al. Effect of calcium supplements on risk of myocardial infarction and cardiovascular events: meta-analysis. BMJ. 2010;341:c3691.

**What was Known?** While widely recommended, calcium has been shown to be necessary but not sufficient for reducing osteoporosis risk. A previous randomized controlled trial in healthy older women with prespecified CVD outcomes showed possible increases in MI and cardiovascular events in women who took calcium.

**What this Study Adds** This meta-analysis included 15 RCTs that had at least 100 participants aged 40 and older and had a study duration of at least 1 year. Among a total of 15 eligible
25-OH vitamin D levels have been revised. A serum 25-OH is now the goal. Black D, Kelly M, Genant H, et al. Bisphosphonates and fractures of the subtrochanteric or diaphyseal femur. NEJM. 2010;362(19):1761–71.

What Was Known? Several case series have reported an increased risk of atypical femoral shaft fractures (fractures of the subtrochanteric or diaphyseal femur) with alendronate. However, it has not previously been possible to estimate population prevalence or identify risk factors.

What this Study Adds Study investigators performed secondary analyses of three large randomized trials of bisphosphonates. These trials included the Fracture Intervention Trial (FIT), the FIT Long-Term Extension (FLEX) trial and the Health Outcomes and Reduced Incidence with Zolendronic Acid Once Yearly (HORIZON) Pivotal Fracture Trial (FFT). They had detailed records of fractures and radiographs to ascertain exact fracture locations and typical or atypical characteristics of the fracture. There were a total of 284 fractures among 14,195 women. Of the 284 fractures, 12 fractures in 10 patients were subtrochanteric or diaphyseal fractures, resulting in combined rate of 2.3 per 10,000 patient years. Compared with placebo, the relative hazard for alendronate in FIT was 1.03 (95% CI, 0.06, 16.46); the relative hazard for zolendronic acid use in HORIZON-PFT was 1.5 (0.25, 9.00), and the relative hazard for continued alendronate use in FLEX was 1.33 (0.12, 14.67). Fracture of the subtrochanteric or diaphyseal femur was very rare even in women on bisphosphonates for up to 10 years. Although all the point estimates showed an increased risk, the confidence intervals were wide because of the small number of events.

How Should I Change my Practice? High dose annual supplementation of vitamin D is not recommended for fall and fracture prevention. Whether more frequent or lower doses of vitamin D may prevent falls or fractures remains unknown. Based on the results of new studies, the Institute of Medicine has recently updated their guidelines for vitamin D intake. The new guidelines for vitamin D are 600 IU (daily) for women aged 9–70, and 800 IU for women over age 70 with an upper recommended limit of 4,000 IU per day. In addition, the target 25-OH vitamin D levels have been revised. A serum 25-OH vitamin D level of 20 ng/ml is now the goal.

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