Summary of Updated Recommendations for Primary Prevention of Cardiovascular Disease in Women

JACC State-of-the-Art Review

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ABSTRACT

Cardiovascular disease (CVD) remains the leading cause of morbidity and mortality for women in the United States and worldwide. There has been no American College of Cardiology (ACC)/American Heart Association guideline update specifically for the prevention of CVD in women since 2011. Since then, the body of sex-specific data has grown, in addition to updated hypertension, cholesterol, diabetes, atrial fibrillation, and primary prevention guidelines. The ACC CVD in Women Committee undertook a review of the recent guidelines and major studies to summarize recommendations pertinent to women. In this update, the authors address special topics, particularly the risk factors and treatments that have led to some controversies and confusion. Specifically, sex-related risk factors, hypertension, diabetes, hyperlipidemia, anticoagulation for atrial fibrillation, use of aspirin, perimenopausal hormone therapy, and psychosocial issues are highlighted. (J Am Coll Cardiol 2020;75:2602–18) © 2020 by the American College of Cardiology Foundation.

Cardiovascular disease (CVD) remains the leading cause of morbidity and mortality for women in the United States and worldwide (1). Overall, 1 in 3 women die from CVD, and 45% of women over age 20 years have some form of CVD (1).

There has been no American College of Cardiology (ACC)/American Heart Association (AHA) guideline update specifically for the prevention of CVD in women since 2011. The last statement to address this was the Effectiveness-Based Guidelines for the Prevention of Cardiovascular Disease in Women—2011.
CVD RISK FACTORS UNIQUE TO WOMEN

PREGNANCY-ASSOCIATED CONDITIONS THAT INCREASE FUTURE RISK OF CVD. Adverse pregnancy outcomes (APO) occur in 10% to 20% of all pregnancies and are associated with a 1.8- to 4.0-fold risk of future CVD (7,8). Risk of CVD is higher with more severe forms of APO or more than 1 pregnancy complicated by an APO (9). Studies of vascular abnormalities in women with APO suggest placental dysfunction, and abnormal endothelial function may be a common pathway and early indicator of later cardiometabolic risk (10). The American College of Obstetrics and Gynecology recommends that women with APO and/or cardiovascular risk factors undergo cardiovascular risk screening within 3 months postpartum (11) (Figure 1, Table 1).

HYPERTENSIVE DISORDERS OF PREGNANCY. Hypertensive disorders of pregnancy are associated with development of incident hypertension after delivery and overall CVD. A meta-analysis of 3,488,160 women, including 198,252 with pre-eclampsia reported that after 10 to 15 years, women with pre-eclampsia had a 3.7-fold risk of hypertension, 2.2-fold risk of ischemic heart disease, 1.8-fold risk of stroke, and 1.5-fold risk of overall mortality (8). Pre-eclampsia was included as a “risk-enhancer” in the updated 2018 cholesterol guideline (4) and in the 2019 ACC/AHA Guideline on the Primary Prevention of Cardiovascular Disease (5). In addition, all hypertensive disorders of pregnancy are associated with increased risk of chronic hypertension (12,13) as early as the first year after delivery (13), twice the risk of CVD-related hospitalizations within 3 years of delivery (14), and development of other classic CVD risk factors such as diabetes and hyperlipidemia (15). A 2019 study of the United Kingdom Biobank cohort found that hypertension during pregnancy was associated with increased risk of coronary disease (hazard ratio [HR]: 1.8; 95% confidence interval [CI]: 1.3 to 2.6; p < 0.001) as well as increased risk of heart failure and valvular heart disease (16). The American College of Obstetrics and Gynecology currently recommends initiation of low-dose aspirin in women with at least 1 high risk factor (history of pre-eclampsia, multifetal pregnancy, chronic hypertension, diabetes mellitus I or II, chronic kidney disease, or autoimmune disorder) or at least 2 moderate risk factors (nulliparity, obesity, family history of pre-eclampsia, socioeconomic factors, age >35 years, or personal history factors) to reduce the risk of pre-eclampsia (17).

GESTATIONAL DIABETES MELLITUS. Women with a history of gestational diabetes mellitus are at increased risk of future CVD, including a 1.4- to 20-fold increased risk of type 2 diabetes mellitus, 2-fold risk of hypertension, 2-fold risk of stroke, and 2.8-fold risk of ischemic heart disease (18).
**PRE-TERM BIRTH.** Pre-term birth is defined as delivery prior to 37 weeks gestation; idiopathic pre-term birth is associated with a 2-fold increased risk of CVD and deaths caused by coronary heart disease (19) even when adjusted for pre-pregnancy lifestyle and CVD risk factors (20). Risk of CVD is higher with more pre-term births and earlier pre-term birth (prior to 34 weeks).

**PREGNANCY LOSS.** Women with prior pregnancy loss (miscarriage and stillbirth) are at approximately 2-fold increased risk of myocardial infarction (MI), cerebral infarction, and renovascular hypertension (21). In a meta-analysis of 10 studies, miscarriage was associated with a 1.45-fold increased risk of CVD, and more than 1 miscarriage was associated with a 2-fold risk of CVD (22).

**Intrauterine growth restriction.** Intrauterine growth restriction (IUGR) is defined as an estimated fetal weight <10th percentile for the gestational age, often related to suboptimal uterine-placental perfusion (23). Several maternal factors are associated with fetal growth restriction, including hypertensive disorders and diabetes (23). Women with prior IUGR pregnancies are at increased risk for hyperlipidemia, hypertriglyceridemia, and insulin resistance (24). Additionally, echocardiographic cardiac changes have been observed in women during normotensive IUGR pregnancies, including higher prevalence of diastolic dysfunction and less cardiac reserve compared with control subjects (25). Low-dose aspirin started in early pregnancy may prevent IUGR in certain patients (26,27).

**RISK PREDICTION MODELS.** Although adverse pregnancy outcomes are associated with later risk of CVD, the addition of pregnancy complications to standard cardiovascular risk prediction models have not significantly enhanced the predictive capabilities (28,29). Because adverse pregnancy outcomes are
also associated with other conventional cardiovascular risk factors that are included in the standard risk models, the additive impact of pregnancy complications becomes less significant, particularly with increasing age. A history of adverse pregnancy outcomes may be most useful in younger women, prior to the development of conventional risk factors, and important for counseling of women about risk prevention.

**PREMATURE MENOPAUSE.** Premature menopause (age <40 years) was considered a risk-enhancing factor in the 2018 cholesterol guideline (4). Menopause increases CVD risk because of the physiological responses to estrogen withdrawal, including changes in body fat distribution, reduced glucose tolerance, abnormal lipids, higher blood pressure (BP), increased sympathetic tone, endothelial dysfunction, and vascular inflammation (30). A 2019 pooled analysis from 15 observational studies including 301,438 women reported increased risk of nonfatal CVD in women with premature menopause (HR: 1.55; 95% CI: 1.38 to 1.73; p < 0.0001), early menopause (age 40 to 44 years; HR: 1.30; 95% CI: 1.22 to 1.39; p < 0.0001), and relatively early menopause (age 45 to 49 years; HR: 1.12; 95% CI: 1.07 to 1.18; p < 0.0001) (31). Recent data from the United Kingdom Biobank cohort reported premature menopause (before age 40 years) was associated with increased risk of CVD (HR: 1.36; 95% CI: 1.19 to 1.56; p < 0.001) after adjustment for conventional risk factors (32). The interaction between CVD and menopause is complex, and it may be that women at increased risk for CVD experience menopause at an earlier age.

**POLYCYSTIC OVARIAN SYNDROME.** Polycystic Ovarian Syndrome (PCOS) is a common endocrine disorder that affects young women, and is characterized by ovulatory dysfunction (oligomenorrhea or amenorrhea), hyperandrogenism, infertility, and insulin resistance (33). Whether PCOS by itself confers high CVD risk, or whether the associated

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**FIGURE 1** Recommendations for Cardiovascular Risk Screening After Adverse Pregnancy Outcomes

<table>
<thead>
<tr>
<th>Conditions:</th>
</tr>
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<tbody>
<tr>
<td>Hypertensive disorders of pregnancy (chronic hypertension, gestational hypertension, preeclampsia, eclampsia, HELLP syndrome)</td>
</tr>
<tr>
<td>Gestational diabetes mellitus</td>
</tr>
<tr>
<td>IUGR (intrauterine growth retardation)</td>
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<tr>
<td>Preterm birth (idiopathic/spontaneous)</td>
</tr>
<tr>
<td>Placental abruption</td>
</tr>
<tr>
<td>Obesity/excessive pregnancy weight gain/post-partum weight retention</td>
</tr>
<tr>
<td>Sleep disorders; moderate-to-severe obstructive sleep apnea</td>
</tr>
<tr>
<td>Maternal age older than 40 years</td>
</tr>
</tbody>
</table>

**Cardiovascular risk screening within 3 months post-partum**

**Medical History**
- Smoking history
- Physical activity
- Breastfeeding
- PMH of hypertension, diabetes, CVD
- First degree family history of CVD, HTN, DM

**Physical Examination**
- Resting blood pressure and heart rate
- Body mass index and waist circumference

**Laboratory testing**
- Lipid profile
- Diabetes screening
- Urine protein:creatinine ratio

Adverse pregnancy conditions that require further cardiovascular screening within 3 months post-partum based on medical history, physical examination, and laboratory. CVD = cardiovascular disease; DM = diabetes mellitus; HELLP = hemolysis, elevated liver enzyme, low platelet count; HTN = hypertension; IUGR = intrauterine growth restriction; PMH = past medical history.
cardiometabolic features are the reason for increased risk is unclear (34). Women with PCOS are at an increased risk for development of metabolic syndrome features of abdominal obesity, diabetes, dyslipidemia, and hypertension (35). These factors contribute to endothelial dysfunction, which is a marker of CVD risk, and several studies have shown endothelial function abnormalities and subclinical atherosclerosis in PCOS (36). Ethnic variation in PCOS has also been reported, with East Asian women with PCOS having the highest prevalence of metabolic syndrome, despite a lower body mass index and less hyperandrogenic features (37). In addition to treatment of menstrual irregularities with oral contraceptives, metformin is recommended for patients who have cardiometabolic features such as abdominal obesity and insulin resistance (38). Although the 2018 cholesterol guideline did not include PCOS as a risk enhancer (4), the international guidelines for PCOS recommend that all women with PCOS should be screened for CVD risk, including close monitoring for weight changes every 6 to 12 months, at least annual BP check, fasting lipid panel, screen for glycemic control, and assessments for smoking and physical activity (39). Psychological factors, such as anxiety, depression, and eating disorders, are prevalent in PCOS, and guidelines recommend that health professionals take into consideration cultural sensitivities and weight-related stigma in women when addressing lifestyle-based interventions (38).

AUTOIMMUNE DISEASE. Women are more likely to have underlying autoimmune and inflammatory conditions that contribute to increased CVD risk, beyond the traditional CVD risk factors. Conditions such as systemic lupus erythematosus (SLE) and rheumatoid arthritis (RA) are highly prevalent in women and are associated with accelerated atherosclerosis as well as coronary microvascular dysfunction (2,40). SLE is more prevalent in Asians, African Americans, African Caribbeans, Hispanic Americans compared with Caucasians. Black women are 2 to 4 times more likely to have SLE than white women (41). Ischemic heart disease is the number 1 cause of mortality in SLE. One study reported that young women with SLE (ages 35 to 44 years) were over 50 times more likely to have an MI compared with those of similar age in the Framingham Offspring study (42). There is a 50% increased risk of CVD mortality in RA compared with the general population (43). Furthermore, there is emerging data that the duration of time in RA flares is associated with increased risk of CVD events (44). A lipid paradox described in 1 study demonstrated that elevated erythrocyte sedimentation rate and low cholesterol levels were associated with CVD risk in RA patients (45). Higher levels of inflammation are associated with major adverse outcomes despite low normal cholesterol levels in other studies (46,47). Treatment with anti-inflammatory agents such as statins, interleukin-1β receptor antagonist canakinumab, and colchicine improves CVD outcomes in various cohorts (47-49). The ACC/AHA risk score derived from pooled cohort equation to estimate atherosclerotic CVD risk does not incorporate these unique risk factors for women; however, the 2018 cholesterol guideline lists these factors as risk enhancing factors that should be taken into consideration when assessing a patient’s CVD risk (4).

**TABLE 1** Complications During Pregnancy That Are Associated With Increased Cardiovascular Disease Risk

<table>
<thead>
<tr>
<th>Adverse Pregnancy Outcomes</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertensive disorders of pregnancy</td>
<td>This category includes gestational hypertension, chronic hypertension, and pre-eclampsia</td>
</tr>
<tr>
<td>Gestational hypertension</td>
<td>New-onset hypertension (DBP ≥140 mm Hg or DBP ≥90 mm Hg) after 20 weeks gestation</td>
</tr>
<tr>
<td>Pre-eclampsia</td>
<td>New-onset hypertension (SBP ≥140 mm Hg or DBP ≥90 mm Hg) after 20 weeks gestation with proteinuria or evidence of end-organ dysfunction</td>
</tr>
<tr>
<td>Chronic (pre-existing) hypertension</td>
<td>Hypertension present prior to 20 weeks gestation</td>
</tr>
<tr>
<td>Gestational diabetes</td>
<td>Glucose intolerance with onset or first recognition during pregnancy</td>
</tr>
<tr>
<td>Pre-term birth</td>
<td>Delivery before 37 weeks gestation</td>
</tr>
<tr>
<td>Early pre-term birth</td>
<td>Delivery before 34 weeks gestation</td>
</tr>
<tr>
<td>Pregnancy loss</td>
<td>Miscarriage or stillbirth</td>
</tr>
<tr>
<td>Intrauterine growth restriction</td>
<td>Fetal birthweight less than expected for the gestational age, &lt;10th percentile</td>
</tr>
</tbody>
</table>

DBP = diastolic blood pressure; SBP = systolic blood pressure.

**HYPERTENSION.** The 2017 ACC/AHA guidelines for the prevention, detection, evaluation, and management of high BP provides limited sex-specific guidance in the management of hypertension and focuses primarily on hypertension during pregnancy (3), which has been extensively reviewed in a paper discussing hypertension across a woman’s lifespan (50). However, there are certain unique aspects in women of the prevention, epidemiology, evaluation, and management of hypertension. Common risk factors for hypertension in women include: obesity, physical inactivity, increased salt intake, diabetes, and more than moderate alcohol consumption (i.e., >1 alcoholic drink/day). The combination of these risk factors is associated with a higher risk of hypertension, and...
obesity has the highest impact on the incidence of hypertension among women (51). Due to the up-regulation of renin-angiotensin receptors after menopause, salt restriction is beneficial in reducing the risk of hypertension. Indeed, reducing salt intake has been shown to reduce systolic BP in women with and without hypertension (52). The 2017 ACC/AHA guidelines recommend to ideally limit sodium intake to <1,500 mg/day or at least aim for a 1,000 mg/day reduction, and to enhance the intake of potassium from foods to at least 3,500 to 5,000 mg/day (3); however, there is no specific recommendations based on sex (Figure 2).

Attention needs to be given to the possible presence of secondary causes of hypertension among pre-menopausal women. In particular, women account for >90% of cases of fibromuscular dysplasia, a condition that affects 3.3% of the general population (53). Combined hormonal contraceptive can also result in an increase in BP, particularly among women with a pre-existing diagnosis of hypertension. Pre-menopausal women requiring antihypertensive therapy also require counseling on potential medication teratogenicity, particularly if receiving angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, or aldosterone receptor antagonists (54).

The prevalence of hypertension among pre-menopausal women tends to be lower than men of similar age; however, hypertension becomes more prevalent in women after menopause (1).

In a recent sex-specific longitudinal BP analysis of >32,000 patients, women were found to have steeper increases in BP than men, which began as early as the third decade of life and persisted with aging, even after adjusting for the cardiometabolic risk factors. This is contrary to the beliefs that vascular diseases lag 10 years or more among women compared with men (55). Because BP represents a simple accessible measure of vascular aging and is a significant contributor to future cardiovascular events, these findings could help explain some of the differences in CVD presentations among women versus men, such as diastolic heart failure (55,56).

The 2017 ACC/AHA guideline recommends out-of-office monitoring of BP for confirmation and management of hypertension irrespective of sex (3). Notably, studies suggested that post-menopausal women are likely to experience a nondipping nighttime BP pattern.
finding that CVD mortality is significantly increased for people diagnosed with type 2 DM before the age of 40 years (67).

Diabetes increases the risk of having an MI or stroke by 2-fold (66). In the presence of type 2 DM, the absolute rate difference between the sexes is significantly diminished, although not fully eliminated (68,69). The cardioprotection that occurs in premenopausal women is thus reduced significantly with diabetes. A recent systematic review and meta-analysis of over 5 million patients found that the pooled relative risk for CVD mortality in patients with DM was 2.42 in women (95% CI: 2.10 to 2.78) and 1.86 (95% CI: 1.70 to 2.03) in men (70). There also appears to be greater excess risk of CVD mortality in women with DM compared with men (relative risk: 1.30; 95% CI: 1.13 to 1.49; p < 0.001) in pooled multiple adjusted analysis, although there was significant heterogeneity between the studies (70). Recently, the Atherosclerosis Risk In Communities study found DM was a stronger risk factor for CVD as well as CVD mortality among African-American women than among African-American men (71). These findings are similar to what has been seen in white male and female patients (71). In addition to atherosclerotic events, having DM increases the incidence of congestive heart failure. In the UK biobank study of 468,941 patients followed for 9.0 years, women with type 2 DM had significantly higher rates of incidence of heart failure (HR: 1.73; 95% CI: 1.34 to 2.24; p < 0.0001) as well as heart failure mortality (HR: 1.92; 95% CI: 1.25 to 2.94; p < 0.003) compared with men (72). Last, DM increases the risk of cancer mortality by 26% in women (95% CI: 1.16 to 1.36) and by 29% in men (95% CI: 1.18 to 1.42) (70). There was no sex difference in the association between diabetes and cancer mortality for diabetic patients (70).

There appears to be some sex-specific effects of pharmacotherapy for DM. For example, it has been reported that glucagon-like peptide-1 receptor agonists have better glycemic control among men than women; however, women had more weight loss (73). Thiazolidinediones appear to have better glycemic reduction in obese women, whereas nonobese men responded better with sulfonylureas (74,75). Reassuringly, the EMPA-REG (Empagliflozin, Cardiovascular Outcomes, and Mortality in Type 2 Diabetes) study, which showed reduction in cardiovascular mortality in diabetic patients treated with the sodium glucose cotransporter 2 inhibitor empagliflozin, showed no significant sex differences in benefit with the drug (76).

Given the increased cardiovascular risk, all patients with DM require aggressive risk factor
reduction. However, studies have consistently shown that women are underdiagnosed and undertreated compared with men (77,78). Women with DM have poorer BP, lipid, and DM control compared with their male counterparts (66).

Table 2 lists BP, lipid, antiplatelet, and hemoglobin A1c goal for diabetic patients without established CVD. Although there are some differing targets among different societies (3,4,6,79-81) regarding BP target, the societies are consistent with aggressive lipid control for diabetic patients. There are no sex differences between the treatment and target recommendations. All societies agree that asymptomatic patients not be routinely screened for CAD.

**BLOOD CHOLESTEROL MANAGEMENT IN WOMEN**

Despite contemporary advancements in cholesterol-lowering therapy, women are less likely to receive guidelines-recommended statin therapy compared with men. They are also more likely to decline initial treatment and less likely to continue prescribed statin therapy (82). The 2018 AHA/ACC multisociety guideline on the management of blood cholesterol and the 2019 ACC/AHA Guideline on the Primary Prevention of Cardiovascular Disease emphasize the importance of lipid management for reduction of atherosclerotic cardiovascular disease (ASCVD) risk and include some sex-specific risk-enhancing factors to help further identify women at increased ASCVD risk (4,6). In addition to lifestyle interventions with diet, exercise, and weight loss, the guidelines recommend statin therapy as the mainstay treatment in 4 groups of patients:

1. Clinical ASCVD;
2. Severe hypercholesterolemia (low-density lipoprotein [LDL] cholesterol >190 mg/dl);
3. Diabetes mellitus in adults (age 40 to 75 years);
4. Primary prevention in adults age 40 to 75 years at high risk (≥20%) and some adults at intermediate risk (≥7.5% to <20%) or borderline risk (5% to <7.5%) based on the presence of risk enhancers, the presence of an elevated coronary artery calcium score if measured, and clinician-patient risk discussion.

The benefit of statin therapy has been widely accepted for reduction of CVD events for secondary prevention in both sexes; however, the role of statin therapy for primary prevention in women has been debated over the past decade. This controversy stemmed in part from a lack of robust data on the efficacy of statins for primary prevention in women, as under-representation of women in randomized controlled trials left studies underpowered to
adequately analyze outcomes by sex. In addition, early meta-analyses of statin therapy for primary prevention yielded conflicting data, with some studies showing no significant reduction in mortality or cardiovascular events in women (83). Since the Effectiveness-Based Guidelines for the Prevention of Cardiovascular Disease in Women—2011 Update (2), 2 larger meta-analyses including over 40,000 women have demonstrated a similar benefit of statin therapy in women and men for both primary and secondary prevention, and this benefit was seen in both sexes across all levels of risk in primary prevention studies (84,85). Although no significant sex differences in adverse effects were identified in these meta-analyses, few statin trials reported adverse drug reactions by sex. Despite a paucity of randomized trial data, international consensus statements recognized female sex as a risk factor for statin-associated muscle symptoms (86,87). In patients with statin-associated muscle symptoms, careful review of concomitant medications and detailed history should be taken to understand factors that may contribute to statin side effects (86,87). Change in statin (hydrophilic vs. lipophilic) as well as intermittent statin dosing can be used to help overcome some of the muscle symptoms associated with statins (88).

There are currently no sex-specific guidelines for the management of blood cholesterol with statin therapy. Statins reduce cardiovascular events and all-cause mortality regardless of sex, and should be considered at recommended doses in women who meet criteria for 1 of the 4 guideline-recommended patient populations (Figure 3).

Reasons for sex differences in quality metrics on the patient and physician level need to be further investigated to ensure optimal primary and secondary prevention in women, given that sex differences in statin prescribing patterns and adherence continue to exist.

**WOMEN WITH DYSLIPIDEMIA AND PREGNANCY**

The guidelines recommend that premenopausal women on statin therapy need to stop the statin 1 to 2 months before attempting pregnancy (4). If the pregnancy is unplanned, the statin should be discontinued as soon as the pregnancy is known (4). Optimal management of cholesterol with healthy lifestyle habits should be discussed first in pregnant women with dyslipidemias (4). Bile acid sequestrants are approved for use during pregnancy.
FUTURE DIRECTIONS, STATINS, AND PREGNANCY. The safety of pravastatin has been under study for the prevention of pre-eclampsia in high-risk pregnant women (89). Statins are known to have pleiotropic effects, which may diminish inflammation and oxidative stress, increase angiogenesis, inhibit the coagulation cascade, and protect the endothelium (90). Human clinical trials are now currently in progress to determine whether a hydrophilic statin (90). Human clinical trials are now currently in progress to determine whether a hydrophilic statin may be used to prevent pre-eclampsia in high-risk women.

NONSTATIN THERAPY IN WOMEN

Ezetimibe reduces cholesterol absorption in the small intestine and is a modest but effective lipid-lowering agent for both men and women. In particular, for women who experience statin-induced myalgias, ezetimibe is a nonstatin alternative for patients who are considered intolerant to statin therapy (defined as intolerant to 2 or more statins and failed alternate dosing therapy) or require additional LDL lowering in addition to maximum-tolerated statin. Monotherapy with ezetimibe will provide an 18% LDL reduction and add on therapy provides a 25% reduction (4). The IMPROVE IT (Ezetimibe added to Statin after Acute Coronary Syndrome) trial, which validated the effectiveness of ezetimibe in combination with simvastatin was conducted in a secondary prevention setting among post-acute coronary syndrome patients, average age over 60 and were predominantly men. Therefore, the effectiveness of ezetimibe in women (in particular midlife women) in the primary prevention setting is less understood (91).

Proprotein convertase subtilisin/kexin type 9 (PCSK9) are monoclonal antibodies with 2 U.S. Food and Drug Administration approved injectables currently available on the market. Cardiovascular outcome studies of PCSK9 inhibitors using alirocumab (ODYSSEY Outcomes [Alirocumab and Cardiovascular Outcome after Acute Coronary Syndrome]) and evolocumab (FOURIER [Evolocumab and Clinical Outcomes in Patients with Cardiovascular Disease]) demonstrated that PCSK9 inhibition added to maximum-tolerated statin significantly reduced LDL cholesterol levels and the rate of major adverse cardiovascular events (92,93). Both studies had smaller numbers of women who participated in their clinical trials; however, subgroup analysis found no treatment heterogeneity by sex (92,93). The ODYSSEY-C (Open Label Study of Long-Term Evaluation Against LDL-C Trial) evaluated longer-term effects of evolocumab during open-label hypercholesterolemia treatment for up to 5 years in over 1,000 patients who tolerated evolocumab up to 4 years (94). Women accounted for 53% of the cohort and demonstrated excellent tolerability to evolocumab with an annual 1.4% discontinuation rate. Although there has not been a primary prevention trial of PCSK9 inhibitors, they seem to be well tolerated and effective at lowering LDL in both men and women.

ASPIRIN THERAPY. Among women with established ASCVD, the role of aspirin is well-established; aspirin reduces subsequent vascular events by approximately 25% (95). Aspirin reduces the risk of atherothrombosis by irreversibly inhibiting platelet function, but this same mechanism comes at a trade-off of increased risk of bleeding, especially in the gastrointestinal tract. In primary prevention, the role of aspirin has been controversial and net benefit less certain for most healthy women. This is because in primary prevention, the absolute risk of vascular events is lower than in secondary prevention, but the complication rates (bleeding) are comparable.

The 2005 WHS (Women’s Health Study), the largest aspirin primary prevention trial, evaluated low-dose aspirin (100 mg every other day) versus placebo in nearly 40,000 women ≥45 years that were free of ASCVD at baseline. The WHS found that low-dose aspirin reduced the risk of stroke over a 10-year follow-up without reducing the risk of MI; however, among women age ≥65 years, aspirin significantly reduced risk of major cardiovascular events including both ischemic stroke and MI (96). Longer (15-year) follow-up suggested that low-dose aspirin was ineffective or harmful for most healthy women, but there may be benefit for women over age 65 years when considering both colorectal cancer and ASCVD events (97).

However, 3 more recent randomized clinical trials, ASCEND (Effects of Aspirin for Primary Prevention in Persons with Diabetes Mellitus), ARRIVE (Use of Aspirin to Reduce Risk of Initial Vascular Events in Patients at Moderate risk of Cardiovascular Disease), and ASPREE (Effect of Aspirin of All-Cause Mortality in the Healthy Elderly), published in 2018, found a lack of net benefit, suggesting that prophylactic aspirin should not be used in the routine primary prevention of ASCVD (98-100). The ASCEND trial evaluated low dose aspirin versus placebo in over 15,000 adults who had diabetes but no ASCVD and found that the absolute benefit for reduction in serious vascular events conferred by aspirin were largely counterbalanced by the increased risk of bleeding (98). The ARRIVE trial, evaluating over 12,000 adults at intermediate estimated ASCVD risk, found no benefit of aspirin for reducing vascular
events but increased risk of gastrointestinal bleeding (99). Finally, the ASPREE trial of over 19,000 adults age >65 years (including 56% women) found no reduction in cardiovascular events with aspirin, but there was an increased risk of bleeding and risk of death (100,101). Finally, an updated 2019 meta-analysis found that the number needed to treat to cause major bleeding was lower than the number needed to treat to prevent an ASCVD event (210 vs. 265), suggesting more harm than benefit (102).

These findings guided the updated aspirin recommendations in the 2019 ACC/AHA Guideline on the Primary Prevention of CVD (6). The 2019 guidelines state that most healthy people do not need to take aspirin, and there were no sex-specific recommendations. These recommendations differ from prior AHA guidelines, which recommended that aspirin could be considered for patients with 10-year ASCVD risk ≥10%. There may still be select patients age 40 to 70 years who have a high ASCVD risk who may benefit from aspirin if they are at low risk for bleeding. One might consider low-dose aspirin (75 to 100 mg/day) among current smokers, those with a strong family history of premature ASCVD, those with very elevated cholesterol suboptimally treated with statins, those with subclinical atherosclerosis such as a coronary artery calcium (CAC) scores ≥100, and other select patients at high ASCVD risk. However, these decisions are needed in the context of a clinician-patient risk discussion. Clinicians should qualitatively evaluate for bleeding risk and withhold aspirin in primary prevention patients with prior gastrointestinal bleeding, known bleeding disorder, severe liver disease, thrombocytopenia, concurrent anticoagulation or NSAID use, or uncontrolled hypertension.

The more recent trials differ than prior trials, since in the modern era, smoking rates are lower and there is more contemporary preventive therapy, including greater prevalence of statin use and BP control. The percent of patients taking statins in ASPREE, ARRIVE, and ASCEND was 34%, 43%, and 75%, respectively (98-100). Population-specific modeling might help identify those anticipated to derive a net benefit of aspirin for primary prevention, but most primary prevention patients are unlikely to benefit (Figure 4) (103).

**STROKE PREVENTION FOR AF.** Many studies have shown that women are at greater risk for AF-related stroke than men. The reason for this higher risk is unclear. Even after adjusting for differences in stroke risk factors and stroke prevention treatment with oral anticoagulants, women have about a 20% to 30% higher risk of stroke than men with AF (104,105). As a
result of this higher risk, female sex was incorporated into the commonly used algorithm, CHA$_2$DS$_2$-VASc score to predict the risk of stroke in patients with nonvalvular AF (106,107).

In 2018, a consensus statement regarding sex differences in arrhythmias was published by the European Heart Rhythm Association and endorsed by the Heart Rhythm Society and Asia Pacific Heart Rhythm Society (108). The statement emphasized the residual stroke risk in women compared with men using vitamin K antagonists and recommended the use of the novel anticoagulants as the first choice (109,110). Compared with men with AF, women with AF had worse stroke severity and more permanent disability after a stroke (111). The statement also highlighted the lower risk of bleeding seen in women compared with men with the use of the novel anticoagulants (108). The statement noted that since a meta-analysis of all 4 novel anticoagulants showed no significant difference with regard to their safety and efficacy in women compared with dose-adjusted warfarin, the novel anticoagulant can be used interchangeably in women depending on personalized needs (112).

The 2019 AHA/ACC/Heart Rhythm Society update on AF guidelines changed the Class I recommendation for anticoagulation, increasing the CHA$_2$DS$_2$-VASc score from ≥2 to ≥3 for women and no change in recommendation for men (CHA$_2$DS$_2$-VASc scores of ≥2) (5). Table 3 is a comparison of the recommendations of the American and European guidelines as well as the updated European recommendations (5,113). Table 4 is key points in atrial fibrillation and women.

There are no sex-specific recommendations for left atrial appendage closure devices or surgical occlusion of the left atrial appendage orifice. However, in a pooled patient-level analysis of the PROTECT-AF (WATCHMAN Left Atrial Appendage System for Embolic PROTECTION in Patients With Atrial Fibrillation) and PREVAIL (Evaluation of the WATCHMAN LAA Closure Device in Patients With Atrial Fibrillation Versus Long Term Warfarin Therapy), in women, LAA closure significantly reduced bleeding compared with patients treated with warfarin (HR: 0.17; 95% CI: 0.074 to 0.369; p < 0.001) (114).

**MENOPAUSAL HORMONE THERAPY.** At this time, there is no role for menopausal hormone therapy (MHT) for CVD prevention. This recommendation is consistent with the American College of Obstetrics and Gynecology statement published in 2013 and reaffirmed in 2018 (115). Since the publication of the HERS (Heart and Estrogen/progestin Replacement Study) (116) secondary prevention trial of MHT, and WHI (Women’s Health Initiative) Study (117), a primary prevention trial of MHT for CVD, long-term use of MHT for CVD prevention is not recommended, as both trials failed to demonstrate cardiovascular benefit and suggested potential harm.

However, there has been much discussion regarding the “timing hypothesis” of MHT. In a

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**Table 3** Comparison and Summary of the Recommendations for Stroke Prevention for Patients With Nonvalvular AF

<table>
<thead>
<tr>
<th>CHA$_2$DS$_2$-VASc Score</th>
<th>ACC/AHA/HRS (5)</th>
<th>ESC (113)</th>
<th>EHRA/HRS/AP HRS (108)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No anticoagulant</td>
<td>No antithrombotic</td>
<td>No antithrombotic</td>
</tr>
<tr>
<td>1</td>
<td>OAC or ASA or no antithrombotic (IIb)</td>
<td>OAC for men (IIa)</td>
<td>OAC for men (IIa)</td>
</tr>
<tr>
<td>2</td>
<td>OAC for men (I)</td>
<td>OAC for men (I)</td>
<td>OAC for men (I)</td>
</tr>
<tr>
<td>≥3</td>
<td>OAC for men and women (I)</td>
<td>OAC for men and women (I)</td>
<td>OAC for men and women (I)</td>
</tr>
</tbody>
</table>

AF = atrial fibrillation; ASA = acetylsalicylic acid; EHRA = European Heart Rhythm Association, HRS = Heart Rhythm Society; OAC = oral anticoagulant; other abbreviations as in Table 2.

**Table 4** Key Points in AF and Women

<table>
<thead>
<tr>
<th>Stroke Prevention in AF in Women</th>
<th>(Ref. #)</th>
<th>First Author (Year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex is a risk modifier and adding female sex to the CHA$_2$DS$_2$-VASc score matters for age &gt;65 yrs or &gt;2 non-sex-related stroke risk factors.</td>
<td>(107)</td>
<td>Nielsen et al. (2018)</td>
</tr>
<tr>
<td>Women with AF have a greater stroke severity and worse long-term outcome in terms of permanent disability, compared with men with AF.</td>
<td>(108,111)</td>
<td>Linde et al. (2018); Martin et al. (2017)</td>
</tr>
<tr>
<td>Women with AF have a greater residual stroke risk even with well-controlled VKAs, which was not seen in randomized controlled trials of the novel anticoagulants.</td>
<td>(109,110)</td>
<td>Sullivan et al. (2012); Pancholy et al. (2014)</td>
</tr>
<tr>
<td>Women taking novel anticoagulants have lower major bleeding rates compared with men.</td>
<td>(108)</td>
<td>Linde et al. (2018)</td>
</tr>
<tr>
<td>There were no significant differences among the novel anticoagulants in terms of safety and efficacy for women.</td>
<td>(112)</td>
<td>Moseley et al. (2017)</td>
</tr>
</tbody>
</table>

AF = atrial fibrillation; VKA = vitamin K antagonist.
combined analysis of the 2 WHI trials, estrogen + progesterone and unopposed estrogen alone, women who started MHT closer to menopausal onset appeared to have lower risk developing subclinical atherosclerosis (118,119) and lower risk of developing CVD (118). However, these findings have not been seen consistently in other trials (120,121).

The most recent meta-analysis in 2017 combining similar long-term MHT studies showed that increased risk of MHT outweighs any benefit in regard to prevention of CVD (122). An increased risk of venous thromboembolism with hormone therapy has been shown with all forms of hormone therapy except for transdermal estrogen (122). Thus, it is imperative that even younger patients who are being considered for treatment for post-menopausal vasomotor symptoms with MHT be assessed for personal and familial risk of venous thromboembolism.

DEPRESSION AND PSYCHOLOGICAL ISSUES IN WOMEN

A large body of epidemiological, experimental, and clinical observations have long linked acute and chronic emotional stress and psychological disturbances, such as depression, to physiological perturbations of the cardiovascular system and the risk of CVD (123,124). Psychosocial stress tends to be a more important risk factor for cardiometabolic diseases in women than in men, not only because women in general have higher exposures to psychosocial stress and adversity than men, but also because they may be more vulnerable to the effects of such exposures (125). In particular, depression, early-life adversities, socioeconomic deprivation, and post-traumatic stress disorder (PTSD) are more prevalent in women than in men and tend to show more robust associations with cardiometabolic risk in women than in men, especially in younger populations or with early exposure (125).

Depression affects approximately 7% of the population each year, and is about 2-fold more common in women than in men (126). Depression is a recognized risk factor for incident MI and cardiac death (127). Among women, a clinical diagnosis of depression is associated with a doubling of risk of CVD even over a period of decades (128,129). Although few studies have examined sex-related differences, available data suggest that depression may be an especially strong risk factor for early-onset CVD in women (130,131).

Compared with men, women have a higher exposure to severe childhood adversities, such as physical and sexual abuse and child neglect, which are increasingly recognized as risk factors for CVD (132). Similar to depression, exposure to adversity in early life appears to be a stronger predictor of CVD in women than in men (133). These early exposures are also predisposing factors for depression and PTSD, as well as strong correlates of adverse lifestyle behaviors.

Although general symptoms of anxiety, measured with a variety of scales, have been associated with incident CVD in a number of studies, individual study results are heterogeneous and the effect sizes are in general modest (134). In contrast, symptoms of PTSD, a condition previously classified among anxiety disorders, have been consistently related to increased risk of CVD (135). In the United States, PTSD affects 9.7% of women (past year prevalence) versus 3.6% of men (136). In a prospective study of women, those with ≥5 PTSD symptoms had an over 3-fold higher risk of ischemic heart disease compared with those without PTSD symptoms, independent of CVD risk factors and depression (137). In the Nurses’ Health Study II, women who reported ≥4 PTSD symptoms had a 60% higher risk of CVD; those with a history of trauma but no PTSD symptoms also showed an elevated CVD risk (45% higher) (138).

There are multiple possible mechanisms linking depression, PTSD, psychological stress, and trauma to CVD. All of these conditions and exposures are associated with poor health behaviors, such as smoking, poor dietary habits, and physical inactivity. Alterations in neurobiological stress response pathways can also play a role, leading to increased inflammation, chronic autonomic dysregulation, endothelial dysfunction, and hypercoagulability (122). Therefore, recognition and management of psychosocial stressors should be useful in promoting a healthy lifestyle and preventing cardiometabolic risk. Currently there are no national guidelines or recommendations on the assessment of these factors in preventive cardiology care. Although there is currently limited understanding of whether interventions addressing psychosocial and emotional disturbances prevent progression to cardiometabolic diseases, recognition and management of these factors should help the quality of life of patients with these conditions, many of whom are women.

CONCLUSIONS

Women have different manifestations of CVD, and studies have shown sex differences in their response to risk factors and treatments. In addition, unique aspects that pertain to women, such as pregnancy-associated conditions that increase future risk, PCOS, and treatment-related issues specific to
women, need to be considered when treating women. Knowledge of updated guideline recommendations are critical in shared decision-making plans to treat women and men to improve CVD outcomes.

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KEY WORDS adverse pregnancy outcomes, aspirin, atrial fibrillation, cardiovascular disease, gestational diabetes