Cardiovascular Changes in Menopause

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Abstract: Menopause is associated with changes consistent with cardiovascular aging. The effects of cardiac disease are multifaceted, affecting endothelial function, coronary artery physiology and metabolic dysfunction leading to structural changes in the coronary anatomy. A systematic review of literature from 1986 to 2019 was conducted using PubMed and Google Scholar. The search was directed to retrieve papers that addressed the changes in cardiovascular physiology in menopause and the current therapies available to treat cardiovascular manifestations of menopause. The metabolic and clinical factors secondary to menopause, such as dyslipidemia, insulin resistance, fat redistribution and systemic hypertension, contribute to the accelerated risk for cardiovascular aging and disease. Atherosclerosis appears to be the end result of the interaction between cardiovascular risk factors and their accentuation during the perimenopausal period. Additionally, complex interactions between oxidative stress and levels of L-arginine and ADMA may also influence endothelial dysfunction in menopause. The increased cardiovascular risk in menopause stems from the exaggerated effects of changing physiology on the cardiovascular system affecting peripheral, cardiac and cerebrovascular beds. The differential effects of menopause on cardiovascular disease at the subclinical, biochemical and molecular levels form the highlights of this review.

Keywords: Dyslipidemia, endothelial dysfunction, oxidative stress, metabolic syndrome, menopause, cardiovascular aging.

1. INTRODUCTION

The life expectancy of women has increased over the years and is currently at 81 years for American women [1]. Menopause (MP) is a normal physiologic event reflecting the loss of ovarian follicle function, decreasing ovarian hormone production, and permanent cessation of menstrual cycles. The average age of natural MP in industrialized countries is around 52 years. Hence, women spend about a third of their lives in MP. Quality of life (QoL) is very important in women as they age.

Coronary Vascular Disease (CVD) is the most common cause of death in post-menopausal women worldwide, even more than cases of breast or other gynecologic cancer deaths. Traditional CVD risk factors include age, smoking, sedentary lifestyle, poor diet, body mass index, hypertension, diabetes mellitus, dyslipidemia, and family history of premature CVD.

There are some sex differences in the incidence of cardiovascular disease (CVD), with women having a lower incidence of CVD before menopause (MP) compared to age-matched men. Pre-menopausal women have a lower prevalence of CAD, likely due to the protective effects of estrogens in females [2, 3]. There is a marked increase in CAD in women after MP, typically seen about 10 years post MP [3, 4].

It is unlikely that MP per se leads to this change, but it is more likely that increasing adverse risk factors like dyslipidemia, insulin resistance, fat redistribution, high blood pressure after MP causing metabolic and vascular changes contributes to the accelerated risk for CAD and cardiovascular disease (CVD). These may be related to adverse peripheral artery endothelial function. The increased CVD risk is thought to begin in the menopausal transition period.

This review attempts to discuss the role of cardiovascular factors that contribute to the increased CVD risk, which begins with MP transition and into post-MP years. CVD risks increase at reproductive aging due to premature MP before age 40 years, early surgical MP by bilateral oophorectomy, or at natural MP.

2. VASCULAR AGING AND CORONARY ENDOTHELIAL FUNCTION

The progressive stiffening of the arteries with a decline in the ability of the vessels to dilate is called vascular aging. It progresses differently in men and women. At the onset of MP, there is accelerated vascular aging, which is different from the gradual loss of vascular function seen with chronologic aging. Vascular aging and endothelial dysfunction contribute to the development of CVD with MP. Vascular aging contributes to the development of hypertension and atherosclerosis [5]. Endothelial dysfunction is a contributor to vascular aging and a key initiator in the development of atherosclerosis [6].
Estrogen is crucial to maintain normal endothelial function. Estrogen increases the synthesis of nitric oxide (NO) by vascular endothelium, which then diffuses into vascular smooth muscle cells, causing its relaxation. This is called Endothelium-dependent vasodilation (EDV). Impaired EDV is a hallmark feature of endothelial dysfunction. Estrogen preserves endothelial function, and declining ovarian hormones with reproductive aging and MP rapidly affects EDV. MP has no effect on endothelial-independent smooth muscle function [7]. Estrogen decreases the synthesis of Endothelin-1 (a potent vasoconstrictor) by the endothelial cells. Hence in MP transition, there is a decrease in EDV and increased synthesis of endothelin 1, both promoting vasoconstriction. Studies have shown that estrogen has antioxidant and anti-inflammatory properties. Estrogen deficiency upregulates oxidative stress or systemic inflammation leading to decreased endothelial function [8]. Thus, estrogen has multiple effects like increasing NO synthesis, antioxidant and anti-inflammatory properties; its deficiency in MP causes endothelial dysfunction [9].

Endothelial dysfunction precedes the development of CAD (obstructive or non-obstructive). In women, endothelial dysfunction and its associated metabolic implications are of greater relevance (than obstructive CAD) in the development of CVD in women. The endothelium of blood vessels synthesizes and secretes substances involved in vasodilation and vasoconstriction. Hence, coronary endothelial function (CEF) serves as a marker for the vascular health of the coronary arteries. It is to be noted that the coronary and systemic vascular beds differ in their biology. Measures of endothelial function in these two vascular beds do not show any strong relations [10, 11]. It is known that normal CEF is affected by traditional and non-traditional CVD risk factors and may respond favorably to modification of these risk factors [12, 13]. Impaired function of coronary endothelium is an early mechanism that plays a role in the development of atherosclerosis and can predict CVD events [14-18]. So far, assessments of CEF was done by invasive methods, thus limiting its use to low risk populations.

Arterial blood flow generates a frictional force per unit area on the endothelium of the vessel wall and is known as shear stress. Disturbed shear patterns act as weak stimuli to release vasoactive substances, cause endothelial inflammation and adhesion of leukocytes to the endothelium. This acts as a proatherogenic environment and can lead to increased production of superoxide radicals and contributes to endothelial dysfunction.

Recently, non-invasive methods to assess CEF with coronary MRI have been developed, and this enables the test to be more widely used [19]. In this technique, the changes in the lumen of coronary arteries (the cross-sectional coronary artery area CSA) and changes in blood flow (coronary blood flow CBF) are measured in response to isometric handgrip exercise (IHE), which is an endothelial-dependent stressor. These luminal and flow changes are dependent on coronary vasodilation caused by endothelial nitric oxide (NO), and thus is a marker for CEF. The changes in CSA with IHE reflect macrovascular endothelial reactivity, while changes in CBF are reflective of both macro and microvascular reactivity. Mathews et al., in the above study, concluded that there are sex differences in CEF, and pre-MP women have better CEF compared to post MP women. The peri MP may be associated with the onset of rapid adverse changes in CEF and MP accelerates the CV risks [20].

3. CHANGES IN LIPID PROFILE

Changes in the lipid profile of women are noted to start around the periMP years, with increases in total cholesterol (TC), LDL cholesterol, triglycerides (TD). The Study of Women’s Health Across the Nation (SWAN) study was a prospective study of MP transition in Caucasian and minority women (African American, Hispanic, Japanese, Chinese) not on hormone therapy. It provided evidence that MP transition is linked to adverse lipid profiles. It showed that TC, LDL and apolipoprotein-B all increase in the 1-year interval around the final menstrual period- independent of the age at which that occurs. All of these are linked to endothelial dysfunction and lead to atherosclerosis. An increase in LDL in the peri MP period is linked to carotid plaques post MP [21, 22]. These changes are distinct from the linear changes in chronologic aging.

The trend of HDL cholesterol or its presumed cardioprotective role is inconsistent over the MP transition. In young women, high HDL cholesterol plays an independent cardioprotective role. This may be because the HDL particles can promote cholesterol efflux, which is a means for HDL to remove cholesterol from peripheral cells [23, 24]. In peri and post MP women, high HDL cholesterol may be linked to higher CVD risk. Carotid intima-media thickness (cIMT) is a marker for vascular health and remodeling. In post MP women with higher HDL levels, the cIMT is greater. This may be due to changes in the quality of the HDL particles over the MP transition.

4. FAT DEPOTS OF HEART

Epicardial adipose tissue (EAT) directly covers the heart between the myocardium and visceral pericardium. Paracardial adipose tissue (PAT) located anterior to the EAT, outside the parietal pericardium, EAT and PAT are now recognized as novel coronary heart disease risk factors [25]. These fat depots may have a more adverse effect than visceral fat due to the proximity to the heart [26]. In the SWAN cardiovascular fat ancillary studies, late periMP/post-menopausal women had 9.9% more EAT and 20.7% more PAT than premenopausal women [28]. Thus, PAT may be a specific MP specific CHD risk marker [26].

5. SUBCLINICAL CVD

Markers like cIMT, coronary artery calcification (CAC), which is a marker of atherosclerotic plaques, aortic calcification, measures of vascular stiffness like aortic pulse wave velocity or flow-mediated dilation (suggestive of endothelial function) are suggestive of subclinical CVD. These can predict CV events. Late periMP, when dyslipidemia and
metabolic syndrome worsen in women, is characterized by increased cIMT [27]. There also seems to be a link between the risk of endothelial dysfunction, and MP transition [28].

6. PREVALENCE OF METABOLIC SYNDROME

Metabolic syndrome is defined as a coexistence of several metabolic risk factors like hypertension, dyslipidemia, impaired glucose tolerance and central adiposity.

Estrogen plays an important part in fat storage and fat distribution. Before menopause, the fat is deposited in thighs, buttocks and hips. Women tend to gain weight (total body fat) during midlife and beyond as a function of chronologic aging. However, when women go through the MP transition, there is a change in the body composition (fat: lean body mass) as well as the distribution of fat. Many women in the MP transition and in post-MP complain of gaining weight in the midsection (android appearance) and trouble with losing weight despite maintaining a healthy lifestyle [29]. MP transition may thus contribute to an increase in visceral (abdominal) fat, insulin resistance, diabetes and inflammatory diseases, leading to the development or worsening of metabolic syndrome in women [29-33].

7. OXIDATIVE STRESS AND MP

Oxidative stress and aging go hand in hand. Overproduction of free radicals such as reactive oxygen species (ROS), and decreased antioxidant levels can lead to atherosclerosis [34]. This decline, combined with a gradual loss of estrogen in the female reproductive system, is highly associated with the various sequelae of MP such as heart disease and vaso-motor disturbances in addition to non-cardiac effects such as osteoporosis. Estrogens exert an antioxidant effect at high concentrations by inhibiting the 8-hydroxylation of guanine DNA bases. Paradoxically at low concentrations, estrogen becomes pro-oxidative. Estrogens have been implicated in DNA adduct production as well as oxidation of bases via ROS formation [34]. Oxidative stress increases inflammatory cytokines and pro-oxidants such as glutathione, 4-hydroxynonal, and malonaldehyde [35], which in turn contribute to further pathology secondary to augmented inflammation.

Oxidative stress increases levels of oxidized LDL [35, 36]. Increased expression of AT1, the angiotensin receptor -1, leads to endothelial dysfunction and secondary increased vasoconstriction observed in atherosclerosis [37]. Decreased levels of nitric oxide play an important role in increased smooth muscle proliferation, inflammation, and atherogenic effects on the vasculature [38]. Nitric oxide exerts cardioprotective effects via inhibition of smooth muscle propagation [38]. Oxidative stress propagates vaso-motor disturbances in MP such as hot flashes and night sweats particularly. Such vaso-motor disturbances result in persistent increases in metabolism, leading to imbalances in pro-oxidants and antioxidants [38].

Disruption of the nitric oxide pathway in MP leads to endothelial dysfunction. The exact mechanisms still remain to be elucidated. Asymmetric dimethylarginine (ADMA) is an endogenous methylated arginine which competitively inhibits nitric oxide (NO) synthesis by competing with L-arginine, the substrate of NO. The competitive inhibition by ADMA leads to a decrease in NO production, translating into endothelial dysfunction contributing to atherosclerosis. ADMA has been implicated as an independent risk factor for cardiovascular disease. Hormone therapy (HT) lowers ADMA concentrations in healthy post-menopausal women. The effect of estrogens on ADMA levels, although small, is considered significant, as the physiological variation of ADMA is limited. Larger randomized trials are necessary to establish that estrogens significantly lower ADMA levels [39]. Current literature reports show that L-arginine appears to be decreased in MP transition. The ratio of L-arginine to L-arginine metabolism biomarker called citrulline, N\(^\text{\textsuperscript{\text{O}}}\)-methyl-L-arginine [L-NMMA] was also decreased. The decreased ratio showed a significant positive correlation with flow-mediated vasodilation of the brachial artery. These findings could suggest a role for L-arginine deficiency in endothelial dysfunction noted in MP transition [40].

8. ESTROGEN DECLINE AND CEREBROVASCULAR DISEASE

Estrogens decrease vascular tone and therefore increase blood flow in the cerebrovascular system, while androgens increase tone. Increased angiogenesis is another function of estrogens and androgens. Inflammation and oxidative stress are reduced by estrogen and therefore exert a neuroprotective effect by preserving the blood-brain barrier and reduce oxidative stress. In the presence of cardiovascular comorbidities, MP changes in hormone levels contribute to cerebrovascular dysfunction and may influence adverse cognitive effects. Further research is needed in this area to elucidate pathophysiology, which in turn could lead to therapeutic developments [41, 42]. Estradiol or E2 declines rapidly over the menopausal transition, which influences cognition, mood and sleep [43]. Further studies are needed to elucidate the effect of E2 definitively in the human brain.

9. MANAGEMENT OF CARDIOVASCULAR SYMPTOMS IN MP

VMS (vasomotor symptom) is the major symptom in MP. Narrowing of the thermoneutral zone so that slight changes in core body temperature brings on compensatory flushing and sweating, leading to hot flashes and night sweats. VMS has been linked to CV risks [44, 45]. VMS also contributes to poor sleep quality, irritability, difficulty in concentration and overall reduced QoL [46]. Lifestyle changes, non-hormonal medications and systemic hormone therapy may be recommended for the management of VMS. The following discussion on the management of VMS and genitourinary syndrome of MP is based on the 2017 position statement from the North American MP Society [47].

Hormone therapy (HT) is not recommended for the primary prevention of any condition- like to preserve cardiovascular health, prevent osteoporosis, prevent memory loss, etc. However, the primary indication for HT is for the management of moderate to severe VMS. Hormone therapy is the
gold standard for the relief of VMS. This may include the use of estrogen alone (Estrogen therapy ET) in women who have had a hysterectomy or estrogen and progesterone therapy (EPT) in women who still have their uterus. Women with a uterus need endometrial protection (against endometrial neoplasia) and are provided by either progestogens or the SERM bazedoxifene. Management of VMS requires the use of systemic hormones that may be given with different routes of administration, including oral (PO) and transdermal (T/D). In general, it is recommended to use the lowest dose of hormones needed for symptom relief for the shortest period of time needed. The type, dose, regimen and duration of use of HT should be individualized.

The decision to offer HT to a menopausal woman for management of VMS requires careful consideration of individual risks and benefits. Based on the Timing Hypothesis, the benefits of HT outweigh the risks in most healthy PM women under the age 60 years or less than 10 years from the final menstrual period. The Women’s Health Initiative (WHI) showed an increased risk of breast cancer with 3-5 years of EPT while 7 years of ET alone did not show the increased risk for breast cancer. It is to be noted that systemic HT is contraindicated in breast cancer survivors.

Lower doses of HT are associated with a lower risk for venous thromboembolism (VTE), less unscheduled vaginal bleeding and less breast tenderness [48, 49]. Lower doses of HT may take 6-8 weeks to provide symptom relief. The formulations of estrogen may include: Oral conjugated equine estrogen (CEE) 0.5mg, oral 17 beta-estradiol 0.5 mg, estradiol patch 0.025mg. If progestogens are indicated for the patient, it may be in the form of oral medroxyprogesterone acetate (MPA), oral progesterone.

The use of progestogens (natural progesterone or synthetic progestogens) alone is a treatment option for VMS; however, they are not as effective as estrogen therapy and have limited long-term safety data. The concern with long-term use is the risk for breast pathology.

Formulations of progestogens include oral MPA 10 mg/day, oral megestrol acetate 20mg or micronized progesterone 300 mg nightly [50-53].

The combination of a SERM (Selective Estrogen Receptor Modulator) called Bazedoxifene with (CEE) is Tissue Selective Estrogen Complex (TSEC) that is FDA approved for management of VMS in women with a uterus. It has the additional benefit of the prevention of osteoporosis. Here, bazedoxifene offers endometrial protection. Hence additional progesterone is not indicated.

The government approved Bioidentical Hormone Therapy (BHT), i.e., hormones similar to endogenous hormones include formulations of estradiol, estrone and micronized progesterone are monitored for their purity, safety and efficacy. The “Compounded BHT” that are marketed as BHT is not approved by the FDA. There are unique concerns here, esp. of safety. These are usually prepared by a pharmacist in a compounding pharmacy based on the provider’s prescription. The compounded BHT may combine many hormones (estrone, estradiol, estriol, DHEA, testosterone and progesterone). Hence the purity, efficacy or safety of ingredients cannot be relied on. The concentration of hormones in these formulations is also uncertain as is its bioavailability. Hence there is a potential for overdosing or underdosing. Compounded BHT usually does not outline its risks. They may contain unapproved combinations of medications and may be used for untested routes of administration, including hormone pellets, troches or subdermal implants [54-56]. Compounded BHT has minimal government regulation and monitoring. They present safety concerns due to potential over or under dosing, presence of potential impurities, unknown sterility of ingredients, lack of safety/efficacy data and label outlining the risks of use.

Compounded BHT should only be considered if patients cannot tolerate the FDA-approved hormones due to issues like an allergy to components or if there is a lack of a dose or formulation of the FDA-approved hormones.

10. MANAGEMENT OF GENITOURINARY SYNDROME OF MP

Genitourinary syndrome of MP (GSM) is a collection of symptoms caused by estrogen deficiency that includes changes in the clitoris, labia, vestibule, vagina, urethra and bladder. Patients may report genital, vaginal dryness, burning, itching, irritation-urinary symptoms of urinary urgency, frequency, dysuria and frequent UTIs. Sexual symptoms include secondary dyspareunia from vaginal dryness. Unlike VMS that improves over the course of time, GSM does not improve with time but instead gets progressively worse over time. Vaginal lubricants and moisturizers may be tried initially for symptom relief. ET is the most effective treatment for GSM [57, 58].

Low-dose vaginal ET is generally safe and effective for the treatment of GSM [59, 60]. Topical ET may be delivered as tablets, suppositories, ring, or creams. The formulation may be estradiol or CEE. Topical ET usually has minimal systemic absorption. It is to be noted that with topical ET, it is not required to use progestogens for endometrial protection, even in women who have a uterus. There are no safety data for topical ET beyond 1 year of use. In women with a history of breast cancer, topical ET for GSM should be prescribed after consulting the patient’s oncologist. In women on aromatase inhibitors, even topical ET is of concern [61, 62].

HT does not have FDA approval for treating any urinary health issues. However, studies show that vaginal ET can improve urge urinary incontinence, overactive bladder and recurrent UTIs. This may possibly due to increased vascularity around the urethra and bladder neck and promoting relaxation of the detrusor muscle. However, systemic HT does not improve urinary incontinence and may actually worsen stress urinary incontinence [63, 64].

Ospemifene is a SERM that is FDA approved for the relief of moderate to severe dyspareunia associated with vulvovaginal atrophy [65]. Being a SERM, it has a class effect of
Fig. (1). Shows the possible impact of cardiovascular risk factors and menopause on the progression of atherosclerosis. (A higher resolution / colour version of this figure is available in the electronic copy of the article).

Fig. (2). Shows the accentuation of cardiovascular risk factors in the setting of menopause.

CONCLUSION

Accelerated atherosclerosis (Fig. 1) appears to be the end result of a complex interaction between cardiovascular risk factors and their accentuation during the perimenopausal period. Additionally, complex interactions between oxidative stress and levels of L-arginine and ADMA may also influence endothelial dysfunction in menopause. The increased cardiovascular risk in MP stems from the exaggerat-
ed effects of changing physiology on the cardiovascular system, which affects vasculature in the peripheral, cardiac and cerebrovascular systems [43, 68]. Changes in lipid profile, vascular stiffness, metabolic parameters, advanced glycation end products (AGE) and oxidative stress (Fig. 2) all contribute to worsening cardiovascular risk in women in the perimenopausal period [68-71]. Treatment strategies should include tight control of cardiovascular risk factors to prevent accelerated cardiovascular disease in menopausal women.

FUTURE DIRECTIONS

Dynamic changes in estradiol and follicle-stimulating hormone levels that lead to vasomotor symptoms should be monitored and treated effectively. Research into developing metabolic markers should be augmented so that possible prophylactic therapeutics can be developed to combat the metabolic syndrome that is accentuated during MP. The female gender-specific aspects and drug therapy in women need more research investigations. Tight regulation of inflammation influences a balanced immune response. The nuclear factor erythroid 2-like 2 (Nrf2) and its role in the inflammation from the standpoint of the development of therapeutics need further investigation. The biochemical basis of nrf-2 activation needs to be studied in the human system. Nrf-2 pathway modulation and its role in cardiovascular aging need more research investigations. Tight regulation of inflammatory cytokines in cardiovascular disease enterprises: Part E: aging arteries: a “set up” for vascular disease. Circulation 2003; 107(1): 139-46.

REFERENCES

[1] Arias E, Xu J. National Vital Statistics Reports, CDC. United States life tables 2017. 2019; 68(7): 1-65.
[2] Mosca L, Hammond G, Mochari-Greenberger H, Towfighi A, Albert MA. American Heart Association Cardiovascular Disease and Stroke in Women and Special Populations Committee of the Council on Clinical Cardiology, Council on Epidemiology and Prevention, Council on Cardiovascular Nursing, Council on High Blood Pressure. Fifteen-year trends in awareness of heart disease in women: results of a 2012 American Heart Association national survey. Circulation 2013; 127(11): 1254-1263, e1-e29. http://dx.doi.org/10.1161/CIR.0b013e318287cf2 PMID: 23429926
[3] Lerner DJ, Kannel WB. Patterns of coronary heart disease morbidity and mortality in the sexes: A 26-year follow-up of the Framingham population. Am Heart J 1986; 111(2): 383-90. http://dx.doi.org/10.1016/0002-8703(86)90155-9 PMID: 3946178
[4] Stampfer MJ, Colditz GA. Estrogen replacement therapy and coronary heart disease: a quantitative assessment of the epidemiologic evidence. Prev Med 1991; 20(1): 47-63. http://dx.doi.org/10.1016/0091-7435(91)90086-P PMID: 1826173
[5] Lakatta EG, Levy D. Arterial and cardiac aging: major shareholders in cardiovascular disease enterprises: Part E: aging arteries: a “set up” for vascular disease. Circulation 2003; 107(1): 139-46. http://dx.doi.org/10.1161/01.CIR.0000048892.83521.58 PMID: 12515756
[6] Rossi R, Nuzzo A, Origliani G, Modena MG. Prognostic role of flow-mediated dilation and cardiac risk factors in post-menopausal women. J Am Coll Cardiol 2008; 51(10): 997-1002. http://dx.doi.org/10.1016/j.jacc.2007.11.044 PMID: 18325438
[7] Taddei S, Virdis A, Ghiadoni L, et al. Menopause is associated with endothelial dysfunction in women. Hypertension 1996; 28(4): 576-82. http://dx.doi.org/10.1152/ajpheart.00396.2018 PMID: 30216121
[8] Sumino H, Ichikawa S, Kasama S, et al. Different effects of oral conjugated estrogen and transdermal estradiol on arterial stiffness and vascular inflammatory markers in postmenopausal women. Atherosclerosis 2006; 189(2): 436-42. http://dx.doi.org/10.1016/j.atherosclerosis.2005.12.030 PMID: 16469323
[9] Stannewicz AE, Wenner MM, Stachenfeld NS. Sex differences in endothelial function important to vascular health and overall cardiovascular disease risk across the lifespan. Am J Physiol Heart Circ Physiol 2018; 315(6): H1569-88. http://dx.doi.org/10.1152/ajpheart.00396.2018 PMID: 30216121
[10] Iantorno M, Hays AG, Schir M, et al. Simultaneous non-invasive assessment of systemic and coronary endothelial function. Circ Cardiovasc Imaging 2016; 9(3): e003954. http://dx.doi.org/10.1161/CIRCIMAGING.115.003954 PMID: 26919997
[11] Anderson TJ, Uchata A, Gerhard MD, et al. Close relation of endothelial function in the human coronary and peripheral circulation. J Am Coll Cardiol 1995; 26(5): 1235-41. http://dx.doi.org/10.1016/0735-1097(95)00327-4 PMID: 7594037
[12] Reriani MK, Lerman LO, Lerman A. Endothelial function as a functional expression of cardiovascular risk factors. Biomarkers Med 2010; 4(3): 351-60. http://dx.doi.org/10.2217/bmn.10.61 PMID: 20550469
[13] Hadi HA, Carr CS, Al Suwaidi J. Endothelial dysfunction: cardiovascular risk factors, therapy, and outcome. Vasc Health Risk Manag 2005; 1(3): 183-98. PMCID: 17319104
[14] Reriani M, Sara JD, Flammer AJ, et al. Coronary endothelial function testing provides superior discrimination compared with standard clinical risk scoring in prediction of cardiovascular events. Coron Artery Dis 2016; 27(3): 213-20. http://dx.doi.org/10.1161/CIR.0000000000000347 PMID: 26882018
[15] Libby P. Inflammation in atherosclerosis. Nature 2002; 420(69170): 868-74.
[16] Schächinger V, Britten MB, Zeiher AM. Prognostic impact of coronary vasodilator dysfunction on adverse long-term outcome of coronary heart disease. Circulation 2000; 101(16): 1899-906. http://dx.doi.org/10.1161/01.CIR.101.16.1899 PMID: 1079454
[17] Suwaidi JA, Hamasaki S, Higano ST, Nishimura RA, Holmes DR Jr, Lerman A. Long-term follow-up of patients with mild coronary artery disease and endothelial dysfunction. Circulation 2000;
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e230421187681

Current Cardiology Reviews, 2021, Vol. 17, No. 4

10(19): 94-58.

http://dx.doi.org/10.11161/01.CIR.101.9.948 PMID: 10704159

[18] von Mering GO, Arant CB, Wessel TR, et al. National Heart, Lung, and Blood Institute. Abnormal coronary vasomotion as a prognostic indicator of cardiovascular events in women: results from the National Heart, Lung, and Blood Institute-Sponsored Women’s Ischemia Syndrome Evaluation (WISE). Circulation 2004; 109(6): 722-5.

http://dx.doi.org/10.11161.CIR.000115525.92645.16 PMID: 14970106

[19] Mathews L, Iantorno M, Schar M, et al. Coronary endothelial function is better in healthy premenopausal women than in healthy older postmenopausal women and men. PLoS One 2017; 12(10): e0186448. http://dx.doi.org/10.1371/journal.pone.0186448 PMID: 29073168

[20] Mathews KA, Koller LH, Sutton-Tyrrell K, et al. Changes in cardiovascular risk factors during the perimenopausal post menopausal and coronary artery atherosclerosis in healthy women. Stroke 2001; 32(5): 1104-11.

http://dx.doi.org/10.1161/01.STR.32.5.1104 PMID: 11340217

[21] Mathews KA, Crawford SL, Chae CU, et al. Are changes in cardiovascular disease risk factors in midlife women due to chronological aging or to the menopausal transition? J Am Coll Cardiol 2009; 54(25): 2366-73.

http://dx.doi.org/10.1016/j.jacc.2009.10.009 PMID: 20082925

[22] Mathews KA, El Khoudary SR, Brooks MM, et al. Lipid changes around the final menstrual period predict carotid subclinical disease in postmenopausal women. Stroke 2017; 48(1): 70-6.

http://dx.doi.org/10.1161/STROKEAHA.116.014743 PMID: 27909203

[23] El Khoudary SR. HDL and the menopause. Curr Opin Lipidol 2017; 28(4): 328-36.

http://dx.doi.org/10.1097/MOL.0000000000000432 PMID: 28459707

[24] El Khoudary SR, Hutchins PM, Matthews KA, et al. Cholesterol efflux capacity and subclasses of HDL particles in healthy women transitioning through menopause. J Clin Endocrinol Metab 2016; 101(9): 3419-28.

http://dx.doi.org/10.1111/jce.2016-2144 PMID: 27399353

[25] Iacobellis G, Gao YJ, Sharma AM. Do cardiac and perivascular adipose tissue play a role in atherosclerosis? Curr Diab Rep 2008; 8(1): 20-4.

http://dx.doi.org/10.1007/s11892-008-0005-2 PMID: 18366994

[26] El Khoudary SR, Shields KJ, Janssen I, et al. Cardiovascular fat, MP and sex hormones in the SWAN cardiovascular fat ancillary study. J Clin Endocrinol Metab 2015; 100(9): 3304-12.

http://dx.doi.org/10.1210/jc.2015-2110 PMID: 26176800

[27] El Khoudary SR, Wildman RP, Matthews K, Thurston RC, Bromberger JT, Sutton-Tyrrell K. Progression rates of carotid intima-media thickness and adventitial diameter during the menopausal transition. Menopause 2013; 20(1): 8-14.

http://dx.doi.org/10.1097/gme.0b013e3182611787 PMID: 22990755

[28] Samargandy S, Matthews K, Janssen I, et al. Central arterial stiffness increases within one-year interval of the final menstrual period in midlife women: Study of Women’s Health Across the Nation (SWAN) heart. Circulation 2018; 137: AP362.

http://dx.doi.org/10.1016/j.lipid.2018.11.014 PMID: 30471324

[29] Russell JK, Jones CK, Newhouse PA. The role of estrogen in brain and cognitive aging. Neurotherapeutics 2019; 16(3): 649-55.

http://dx.doi.org/10.1016/j.neuint.2018.11.014 PMID: 30471324

[30] El Khoudary SR, Wildman RP, Matthews K, Thurston RC, Bromberger JT, Sutton-Tyrrell K. Progression rates of carotid intima-media thickness and adventitial diameter during the menopausal transition. Menopause 2013; 20(1): 8-14.

http://dx.doi.org/10.1097.gme.0b013e3182611787 PMID: 22990755

[31] Lovejoy JC, Champagne CM, de Jonge L, Xie H, Smith SR. Inverse relationship. J Am Coll Cardiol 2009; 54(25): 2374-5.

http://dx.doi.org/10.1016/j.jacc.2009.10.008 PMID: 20082926

[32] Wang Z, Chandrasena ER, Yuan Y, et al. Redox cycling of catechol estrogens generating apurinic/apyrimidinic sites and 8-oxo-deoxyguanosine via reactive oxygen species differentiates equine and human estrogens. Chem Res Toxicol 2010; 23(8): 1365-73.

http://dx.doi.org/10.1021/tr1011282 PMID: 20599668

[33] Signorelli SS, Neri S, Sciacchitano S, et al. Behaviour of some indicators of oxidative stress in postmenopausal and fertile women. Maturitas 2006; 53(1): 77-82.

http://dx.doi.org/10.1016/j.maturitas.2005.03.001 PMID: 16325025

[34] Thernorph A, Calles-Escandon J, Sites CK, Pochlman ET. Menopause, central body fatness, and insulin resistance: effects of hormone-replacement therapy. Coron Artery Dis 1998; 9(8): 503-11.

http://dx.doi.org/10.1097/00001505-199809080-00006 PMID: 9847982

[35] El Khoudary SR, Wildman RP, Matthews K, Thurston RC, Sutton-Tyrrell K, Everson-Rose SA, Hess R, Powell PM, Moreau KL. A relative L-arginine deficiency contributes to endothelial dysfunction across the stages of the menopausal transition. Physiol Rep 2017; 5(17): e13409.

http://dx.doi.org/10.14814/phy2.13409 PMID: 28904082

[36] El Khoudary SR, Wildman RP, Matthews K, Thurston RC, Bromberger JT, Sutton-Tyrrell K. Progression rates of carotid intima-media thickness and adventitial diameter during the menopausal transition. Menopause 2013; 20(1): 8-14.

http://dx.doi.org/10.1097/gme.0b013e3182611787 PMID: 22990755

[37] Nelsen HD. Menopause. Lancet 2008; 371(9614): 760-70.

http://dx.doi.org/10.1016/S0140-6736(08)60346-3 PMID: 18313505

[38] The NAMS 2017 Hormone Therapy Position Statement Advisory Panel. The 2017 hormone therapy position statement of The North American Menopause Society. Menopause 2017; 24(7): 728-53.

http://dx.doi.org/10.1097/GME.0000000000000921 PMID: 28650869

[39] Canonico M, Plu-Bureau G, Lowe GD, Scarabin PY. Hormone replacement therapy and risk of venous thromboembolism in postmenopausal women: systematic review and meta-analysis. BMJ 2008; 336(7655): 1227-31.
