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# Female Sex and Alzheimer's Risk: The Menopause Connection

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### Abstract

Along with advanced age and apolipoprotein E (APOE)-4 genotype, female sex is a major risk factor for developing late-onset Alzheimer's disease (AD). Considering that AD pathology begins decades prior to clinical symptoms, the higher risk in women cannot simply be accounted for by their greater longevity as compared to men. Recent investigation into sex-specific pathophysiological mechanisms behind AD risk has implicated the menopause transition (MT), a midlife neuroendocrine transition state unique to females. Commonly characterized as ending in reproductive senescence, many symptoms of MT are neurological, including disruption of estrogen-regulated systems such as thermoregulation, sleep, and circadian rhythms, as well as depression and impairment in multiple cognitive domains. Preclinical studies have shown that, during MT, the estrogen network uncouples from the brain bioenergetic system. The resulting hypometabolic state could serve as the substrate for neurological dysfunction. Indeed, translational brain imaging studies demonstrate that 40-60 year-old perimenopausal and postmenopausal women exhibit an AD-endophenotype characterized by decreased metabolic activity and increased brain amyloid-beta deposition as compared to premenopausal women and to age-matched men. This review discusses the MT as a window of opportunity for therapeutic interventions to compensate for brain bioenergetic crisis and combat the subsequent increased risk for AD in women.

#### Keywords

Perimenopause; women; Alzheimer's disease; early detection; brain imaging

## Alzheimer's Disease Risk and Female-Specific Hormonal Changes

Along with advanced age and apolipoprotein E (APOE)-4 genotype, female sex is one of the major risk factor for developing late-onset Alzheimer's disease (AD) (1). Epidemiologic

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studies have for years consistently documented that women comprise two-thirds of people living with AD, regardless of age and ethnicity (1–3).

These findings cannot be fully accounted for by the fact that women tend to have greater longevity than men (4). Further, there is consensus that AD is not primarily a disease of the old but rather begins decades prior to clinical symptoms with changes in brain biochemistry (4), which has led investigation into gender differences that may explain the increased risk for females of developing AD independent of longer lifespan in the national effort to accelerate and improve early detection of AD (3).

One major sex difference heralded by pre-clinical and human studies as having early prognostic import for AD is the menopause transition (MT) (5). Though commonly thought of as a neuroendocrine transition state in females ending with reproductive senescence, MT also manifests a number of neurological symptoms, including disruption of estrogen-regulated systems such as thermoregulation, sleep and circadian rhythms, depression, and impairment in multiple cognitive domains (5). Preclinical studies demonstrate that uncoupling of estrogen-mediated brain energy metabolism both marks the onset of MT and induces a hypometabolic state associated with accompanying neurological symptoms (5).

Pathophysiological mechanisms of AD that are activated years to decades prior to clinically detectable symptoms form the basis of a prodromal (preclinical) stage of the disease (5, 6). While symptoms of late-onset AD typically arise in the early to mid-70s, AD pathology appears during the prodromal phase, which is proximate to MT (5, 6). Menopausal changes therefore coincide with the timespan between average age of menopause, in the early-to-mid-50s (7), and average age of AD diagnosis, in the mid-seventies.

Recent translational brain imaging studies have found that 40-60 year-old perimenopausal and postmenopausal women exhibit an AD-endophenotype characterized by decreased metabolic activity (including reduced glucose metabolism and mitochondrial function) and increased brain amyloid-beta deposition (A $\beta$ , a hallmark of AD pathology) as compared to premenopausal women and age-matched men (8, 9). These findings substantiate the eclipsing effects of endocrine aging on chronological aging in the female's brain several years, if not decades, before possible clinical symptoms emerge and the progressively increased risk of AD in women due to MT (6).

Increased risk for AD in females as initiated by brain modifications during the MT is the focus of this review. It also considers the importance of further investigation of the MT as a risk-stratifying method for early detection of AD progression and subsequent intervention, which will be increasingly needed to curb the growing AD epidemic (3, 5, 6).

#### **Clinical Characterization of the MT**

Menopause, by definition, is the final menstrual period a woman experiences and is diagnosed after 12 months of amenorrhea. This universal aging process of a woman's reproductive system occurs at an average age of 51 years with a range of 40-58 years for 88% of women (5, 7). From a clinical perspective, premenopausal women have regular menstrual cycles that have less than 7 days variability in cycle length. Perimenopausal

women have irregular menstrual cycles, with variation in cycle length greater than 7 days. Postmenopausal women have had no menstrual cycles for 12 months or longer (10).

From an endocrinological perspective, perimenopause is characterized by increased variability in the length of the menstrual cycle and levels of circulating hormones. During perimenopause, decreased ovarian reserve results in fluctuations in levels of hypothalamic, pituitary and ovarian hormones, and, notably, a decrease in ovarian secretion of estrogen and progesterone (10). Hormonal changes and clinical symptoms occur over a period leading up to and immediately following menopause. This period is frequently termed climacteric, or perimenopause, but is increasingly referred to as the MT. Determination of these well-defined endocrine features has enabled a symptom and indicator classification system, known as The Stages of Reproductive Aging Workshop (STRAW) criteria (10), that describes the stages of reproductive aging across the transition from perimenopause to postmenopause (Figure 1).

Neurological symptoms of MT include temperature dysregulation (hot flashes), disturbed sleep, insomnia, pain, depression, and cognitive dysfunction (5). Extensive epidemiological analyses designate these symptoms as risk factors for AD (5). Neurological symptoms tend to cluster together in occurrence and severity and, in many instances, persist into postmenopause (11). The breadth of neurological symptoms that can occur during the perimenopausal transition is indicative of disruption in multiple systems, whereas the simultaneous emergence of such disruptions is indicative of a common controlling factor: estrogen.

#### The Role of Estrogen in the Brain

One of the brain's master regulatory systems is the estrogen receptor network. Under its influence, the brain effectively responds at proper timescales to regulate brain energy metabolism, such as in the ovarian-neural estrogen axis (5). Changes in either the availability of estrogen or its receptor network can affect intracellular signaling, neural circuit function, and energy availability (5).

Neural structures in the brain that control the multiple functions affected during MT are populated with estrogen receptors (12, 13). Such receptors are localized to plasma membranes, mitochondria, and in the nucleus of cells, and are particularly abundant in the hypothalamus, which is the primary thermoregulatory center and regulator of sleep and circadian rhythms (14). Brain regions that are crucial to learning and memory, including the prefrontal cortex, hippocampus, amygdala, and posterior cingulate cortex, also contain substantial estrogen receptors (14). Finally, the serotoninergic neurons of the raphe nucleus are positive for estrogen receptors, as are the adrenergic neurons of the locus coeruleus (14).

MT is also associated with the emergence of estrogen receptor splice variants, changes in protein expression, alterations in receptor degradation, and possible epigenetic reconfigurations, all which lead to plummeting estrogen levels (5). The brain compensates for decreased estrogen and reduced receptor activity during the MT. But for some women this adaptive compensation is diminished, lacking, or only expressed in a sub-set of

estrogen-regulated neural networks, giving rise to the complex neurological phenotype of menopause (7, 15).

In particular, estrogenic regulation of brain glucose metabolism is dismantled during perimenopause, inducing a hypometabolic state (12, 16, 17). Preclinical studies indicate that during perimenopause, when brain estrogen plummets, the systems required for estrogen activation of cerebral glucose metabolism rates (CMRglc) and suppression of the ketogenic pathways disassemble (18). Following the decline in CMRglc is induction of an adaptive starvation reaction to increase fatty acid metabolism for the generation and utilization of ketone bodies by mitochondria as an alternative fuel (5, 18–21). Hypometabolism, reduced mitochondrial function and subsequent oxidative damage are known to promote accumulation of Aß pathology and neuronal dysfunction (22), therefore increasing risk of developing AD later in life.

These findings collectively provide additional evidence for the contribution of declining estrogen levels to dysregulated glucose metabolism in brain regions that confer cognitive functioning.

#### Imaging the Menopausal Brain

Recent brain imaging studies have investigated AD risk during endocrine transition states in clinically and cognitively normal women and men to characterize sex-dependent emergence of an AD endophenotype, characterized by A $\beta$  deposition (11C-PiB PET), glucose hypometabolism (18F-FDG PET) and brain atrophy (MRI) (9).

Both postmenopausal and perimenopausal groups exhibited AD biomarker abnormalities as compared to age-matched men (9). These abnormalities included reduced CMRglc, increased A $\beta$  accumulation, and gray and white matter loss in key brain regions for AD, such as parieto-temporal, posterior cingulate, and frontal cortices. Biomarker deficits were most pronounced in postmenopausal women, intermediate in perimenopausal women, and lowest in premenopausal women (9).

Additionally,  $A\beta$  deposition was exacerbated in postmenopausal women positive for APOE-4 (9), a major genetic risk factor for AD that disproportionally affects AD risk in women (1, 23). Even in the absence of dementia, APOE-4 is known to significantly increase brain atrophy and decrease brain connectivity much more strongly in women than in men (24, 25). While it seems likely that loss of estrogen during perimenopause may be driving the observed AD biomarker shifts, other theories exist. For example, the drop in estrogen levels may act as a surrogate biomarker of other underlying molecular changes, including age-related declines in bioenergetic function. As mitochondria are the generators of ATP required for cognitive and other brain functions, and mitochondrial deficits have long been implicated in AD (26), a follow-up PET study investigated whether AD biomarker changes in postmenopausal women were related to altered mitochondrial activity (8).

On 18F-FDG PET, as compared to premenopausal women, perimenopausal and postmenopausal women exhibited hypometabolism in the same brain regions as clinical AD patients, as well as correlated reductions in platelet mitochondrial cytochrome oxidase

(COX, electron transport chain complex IV) activity, a rate limiting step in ATP production (8). A gradient effect was observed so that bioenergetic abnormalities were again most pronounced in postmenopausal, intermediate in perimenopausal, and lowest in premenopausal women (Figure 2) (8). Results were independent of age, education, and APOE genotype (8).

Further investigation is required to accurately determine the role of bioenergetic dysfunction in the pathogenesis of AD. The amyloid cascade hypothesis situates Aß dysmetabolism upstream of oxidative stress, a mechanism typically associated with the rare, early-onset familial AD cases arising from autosomal dominant genetic mutations (27). In early-onset familial AD cases, brain imaging demonstrates that amyloid deposition indeed occurs prior to hypometabolism (28). As an alternative to the amyloid cascade hypothesis, reduced metabolic activity in perimenopausal and postmenopausal women could potentially arise as a consequence of the hypothesized bioenergetic and mitochondrial crisis in the brain during MT (5). Further imaging studies are needed to confirm where bioenergetic dysfunction sits in the common, late-onset form of AD.

Finally, early age of menopause was found to influence cognitive decline and Alzheimer's amyloid pathology in a large cohort of older women which all had surgically-induced menopause (29). Prospective studies are warranted to provide accurate determination of how age of menopause influences the observed biomarker changes in naturally-occurring as well as surgically-induced menopausal women.

#### Interventions

As noted, although the outcome of the MT is inevitable for all women, the experience of the MT and need for intervention can be highly variable. For a thorough discussion of available therapeutic options, we recommend (30). Below is a brief summary of the literature so far, with a focus on AD.

Hormone Replacement Therapy. Hormone Replacement Therapy (HRT) replaces estrogen and progesterone to treat common symptoms of menopause, such as vasomotor symptoms. HRT can help to provide relief of vasomotor symptoms (e.g., hot flashes), reduce the risk of unwanted pregnancy, avoid the irregularity of menstrual cycles, and preserve bone density. However, current FDA guidelines only approve HRT for severe vasomotor symptoms (31). The FDA also advises women to use HRT for the shortest time and at the lowest dose possible to control menopausal symptoms. The FDA lists blood clots, heart attacks, strokes, breast cancer, and gall bladder disease as potential risks of HRT (31).

Whether the risks of HRT outweigh its benefits is an area of active debate. The most comprehensive evidence about risks and benefits of HRT comes from two randomized clinical trials that were sponsored by the National Institutes of Health as part of the Women's Health Initiative (WHI), a 15-year study tracking over 161,800 healthy, postmenopausal women (32). The clinical trials included the WHI Estrogen-plus-Progestin Study, in which women with a uterus were randomly assigned to receive either a hormone medication containing both estrogen and progestin (Prempro<sup>TM</sup>) or a placebo; and the WHI

The WHI trials have also shown some benefits related to use of HRT, including one-third fewer hip and vertebral fractures, and one-third lower risk of colorectal cancer relative to placebo (32, 34). However, a follow-up study found that neither benefit persisted after the study participants stopped taking combined hormone therapy medication (34). Longer-term follow-up of the participants continues to provide new information about the health effects of HRT. Specifically, HRT when instituted prior to the onset of menopause provides protective effects on cardiovascular health (35).

The impact of HRT on the risk of AD remains controversial. The WHI clinical trials reported that combined hormonal therapy doubled the risk of developing dementia among postmenopausal women age 65 and older (33). However, other clinical trials have shown that HRT is effective at preserving CMRglc in AD-regions and preventing cognitive decline, if initiated prior to menopause (36, 37). There is also evidence that, among women who undergo bilateral oophorectomy prior to menopause, those who do not use HRT may double their risk for AD compared to those who initiate HRT soon after the procedure and continue usage at least until the age of natural menopause (38). Altogether, research indicates that HRT may be protective against AD if initiated within a maximum of 5 years after menopause (39–41), whereas HRT can be deleterious if initiated later (42, 43) or in type 2 diabetic women (36, 44).

Two ongoing hypotheses offer explanations for the positive effects of early HRT on AD risk. The "window of opportunity" hypothesis proposes that long-term estrogen depletion (due to MT) may cause decreased levels of estrogen-receptor-alpha in the hippocampus, rendering it irresponsive to HRT (45). The other hypothesis, termed "healthy cell bias of estrogen benefit," suggests that age-related damage to mitochondria may blunder the potentially positive effects of estrogen, or even negatively suffer from them (46). It is likely that both mechanisms contribute to differential benefits of HRT when initiated at different times during the MT.

Non-hormonal therapy. In June 2013, the FDA approved paroxetine mesylate (Brisdelle<sup>™</sup>) as the first non-hormonal therapy for vasomotor symptoms (hot flashes) associated with menopause (47). In women who suffer from hot flashes but either cannot undergo estrogen or hormone therapy because of a history of breast cancer and/or history of deep vein thrombosis, or choose not to do so, some FDA-approved anti-depressants (SSRIs and SNRIs —in particular, venlafaxine) have also been shown to alleviate vasomotor symptoms (48). More work is needed to evaluate these antidepressants for AD prevention in menopausal women.

Alternative products and lifestyle interventions. Phytoestrogens, estrogen-like compounds derived from sources such as soy products, oil seeds (primarily flaxseed), whole-grain

cereals, legumes like chickpeas, or the plant black cohosh, are reputed to ease menopausal symptoms (49). A double-blind, randomized controlled trial reported mild but favorable effects of soy protein supplementation on body composition (e.g., weight, body mass index [BMI], and percent body fat) in treated postmenopausal women as compared with those who received milk protein, over a 6 month period (50). Each group also received isoflavones (50). However, other randomized, double-blind trials have not shown that any of these remedies is superior to a placebo in relieving hot flashes (51). Trials of other herbal remedies, such as evening primrose oil, ginseng, and wild yam, have also not shown efficacy (52).

Clinical trials are needed to test whether dietary patterns rich in these foods, rather than isolated food categories, are protective against AD in women. Two large cohort studies have hypothesized an association but reported conflicting findings (53, 54). The European Prospective Investigation into Cancer and Nutrition (EPIC) study found that women with a higher intake of total fat, protein, and meat experienced a delayed onset of natural menopause (53). The Shanghai Women's Health Study found that, in women aged 40-70 who experienced natural menopause during the study, intake of protein, fruit and tea influenced later age at menopause, whereas fat intake was inversely correlated (54). It is possible that conflicting results may depend on the type of fat in the diet. Recently, the UK Women's Cohort Study showed that a higher intake of oily fish and fresh legumes was associated with delayed onset of natural menopause by 3.3 and 0.9 years per portion/day, respectively, whereas consumption of refined grains was associated with an earlier age of menopause (-1.5 years per portion/day) (55). A higher intake of vitamin B6 and zinc was also associated with later age at menopause (55). By further establishing a predictive relationship between diet and age at natural menopause, and given findings that AD pathology starts to accumulate during the MT in women, dietary interventions would thus have a valuable impact on etiological research and prevention of AD.

In addition to diet, some evidence suggests that exercise and cardiorespiratory fitness are associated with a reduced risk of cognitive decline and AD in women (56, 57), though not all studies agree (58). These conflicting results may be due to the stage of life in which exercise is measured. In a very large cohort study, women who were physically active at age 30 and 50 had a reduced risk for AD as compared to more sedentary women (59). This advantage was significantly stronger in those women who were also physically active as teenagers (59).

#### **Further Directions**

Beyond statistics that expose this greater global burden of AD for females, empirical data about sex differences in AD pathogenesis is warranted to guide clinical and scientific approaches to AD for the benefit of both women and men. Since AD disproportionately affects women, clinical trials may ensure greater progress by focusing on the effects of sex differences on therapy response rates.

To date, though no disease-modifying drug has been proven effective, there is consensus that interventions aimed at preventing or delaying AD progression would be more successful if

instituted earlier in the disease course (3, 6). The MT might well represent a window of opportunity to prevent age-related neurological diseases in women.

Translational brain imaging findings indicate a progressively increased risk of AD in women undergoing the MT, and suggest that endocrine aging outweighs the effects of chronological aging in the female brain by several years, if not decades, before possible clinical AD symptoms emerges (8, 9). Imaging the menopausal brain is of relevance given the current understanding of AD as a progressive disorder characterized by an extended preclinical phase during which the disease is underway but has not led to any recognizable clinical or cognitive symptoms (6). Imaging data suggesting that bioenergetic deficits may be driving amyloid deposition in at least some women (8) may serve to revive zeal in pursuit of new treatments and preventative sex-targeted interventions for AD.

It is important to highlight the fact that not all menopausal women develop AD. Just as importantly, only approximately 80% of women experience neurological symptoms during the MT, while 20% do not (5). It is possible that estrogen-mediated brain networks of those women who do not experience symptoms are not dismantled to the same extent observed in those who experience those symptoms, therefore reducing risk of AD. Alternatively, it is possible that postmenopausal women who do not develop AD may be compensating against menopausal brain changes thanks to known AD-protective factors including healthy diet, exercise, intellectual stimulation, and lack or management of vascular risk factors such as hypertension, diabetes, obesity and cardiovascular disease (60). More research is needed to identify female-specific AD-protective factors.

Biomarker results support further investigation into the potential efficacy of estrogen-based and other therapies in preventing decline in brain bioenergetic capacity in women at the perimenopausal and postmenopausal stage. While it is currently unknown as to the most optimal time to begin potential risk reducing interventions, primary prevention and secondary prevention have received growing attention toward determining which stage of the MT may have the greatest potential for success. Many factors contribute to the confounding HRT reports, including unresolved issues regarding the timing of intervention, hormone formulation, dose, route of administration, metabolic comorbidities, and genetic factors, among other considerations. Optimizing and personalizing hormone therapy remains an unmet need in women's health and a central issue for precision medicine to treat neurological symptoms that develop during, and persist beyond, the perimenopause.

We propose that, given the complexity of the MT, clinical trials examining the effectiveness of estrogen-based and other pharmacological interventions and lifestyle interventions would benefit from stratifying female participants into categories according to their symptoms so as to target specific MT phenotypes rather than enrolling a heterogeneous female population.

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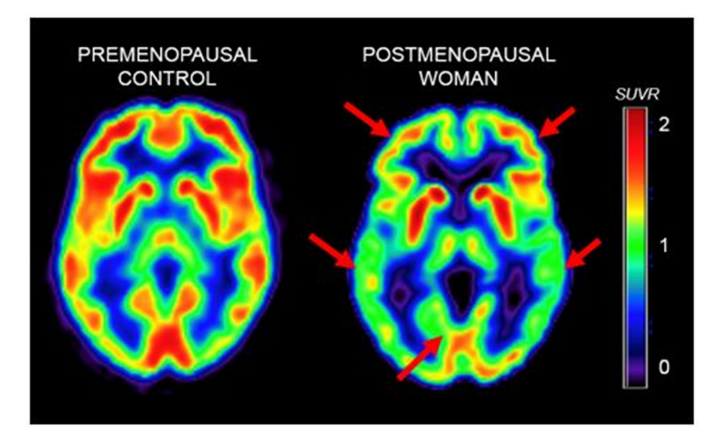
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Menarche					Final Menstrual Period (0)					
Stage:	-5	-4	-3b	-3a	-2	-1	+1 (a)	+1 (b)	+1 (c)	+2
Terminology:	REPRODUCTIVE				MENOPAUSAL TRANSITION		POSTMENOPAUSE			
	early	peak	late		early	late	early			late
		3	22		PERIMENOPAUSE					
Duration:	variable				variable	1-3 years	2 years 3-6 years (1+1)		Rest of lifespan	
Menstrual Cycle <sup>4</sup>	Variable- to-regular	Regular	Regular	Subtle Changes in Flow and Length	Variable length: persistent ≥7-day difference in length of consecutive cycles	Interval of amenorrhea of≥60 days				I
Laboratory find	ings				<b>I</b>	-				
FSH <sup>2</sup>			Low	Variable	Elevated variability*	>25 IU/L**	Elevate variabi		Stable	
AMH <sup>2</sup>			Low	Low	Low	Low	Low		Very Low	
Inhibin B²				Low	Low	Low	Low		Very Low	
Antral Follicle Count <sup>3</sup>			Low	Low	Low	Low	Very L	.ow	Very Low	
Symptoms <sup>4</sup>						Vasomotor symptoms <i>likely</i>	Vasomo sympton most lik	ms		Increasing symptoms of urogenital atrophy

#### Figure 1.

The Stages of Reproductive Aging Workshop + 10 staging system for reproductive aging in women

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#### Figure 2.

FDG-PET scans of two representative female participants at the premenopausal (left side of figure) and menopausal stages (right side of figure)