Clinical epidemiology of Alzheimer’s disease: assessing sex and gender differences

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Abstract: With the aging of the population, the burden of Alzheimer’s disease (AD) is rapidly expanding. More than 5 million people in the US alone are affected with AD and this number is expected to triple by 2050. While men may have a higher risk of mild cognitive impairment (MCI), an intermediate stage between normal aging and dementia, women are disproportionately affected with AD. One explanation is that men may die of competing causes of death earlier in life, so that only the most resilient men may survive to older ages. However, many other factors should also be considered to explain the sex differences. In this review, we discuss the differences observed in men versus women in the incidence and prevalence of MCI and AD, in the structure and function of the brain, and in the sex-specific and gender-specific risk and protective factors for AD. In medical research, sex refers to biological differences such as chromosomal differences (eg, XX versus XY chromosomes), gonadal differences, or hormonal differences. In contrast, gender refers to psychosocial and cultural differences between men and women (eg, access to education and occupation). Both factors play an important role in the development and progression of diseases, including AD. Understanding both sex- and gender-specific risk and protective factors for AD is critical for developing individualized interventions for the prevention and treatment of AD.

Keywords: Alzheimer’s disease, dementia, sex, gender, risk factors, dimorphic medicine

Introduction

Alzheimer’s disease (AD) is the most prevalent type of dementia, comprising about 60%–70% of all dementia cases.1 Beta-amyloid plaques, neurofibrillary tangles, and neurodegeneration are the hallmark pathologic characteristics of AD. Clinically, AD is a progressive disorder characterized by loss of memory and overall cognitive functioning and by behavioral symptoms such as apathy, depression, and anxiety; vocabulary and crystallized abilities are preserved. The burden of AD is high, with more than 5 million people currently affected in the US alone. Presently, one in nine people aged 65 and older has AD and more than one in three people aged 85 and older are affected.2 With the increasing age of the population, it is estimated that 14–16 million Americans will be diagnosed with the disease by 2050 unless new treatments to prevent or delay the onset of AD are identified.3,4 Women are disproportionately affected by AD; they are more likely to become caregivers to AD patients, and are also more likely to develop AD.5 In contrast, some studies suggest that men are at greater risk of developing mild cognitive impairment (MCI), a state between the normal cognitive changes associated with aging and early dementia.5,6 Sex-related differences in the rate of progression after a diagnosis of AD and in the response to treatments have also been reported.
In the present review, we will first summarize sex differences in cognitive aging, in the prevalence and incidence of MCI and AD, and in the rate of progression after an AD diagnosis. We will then discuss sex-specific differences in the neuroimaging measures used to study AD. Lastly, we will review potential reasons for the described differences between men and women considering factors both related to biology (sex) and to society and culture (gender).7–9

**Sex versus gender**

An Institute of Medicine report published in 2001 concluded that “being male or female is an important fundamental variable that should be considered when designing and analyzing basic and clinical research.” In medical research, sex refers to biological differences such as chromosomal (eg, XX versus XY chromosomes), gonadal, or hormonal differences. In contrast, gender refers to psychosocial and cultural differences between men and women (eg, access to education and occupation). Both factors play an important role in the development and progression of diseases, including AD. In this review, potential sex and gender differences that may influence the difference in prevalence and incidence rates of MCI and AD among men and women will be discussed.

**Sex differences in cognitive aging, in the prevalence and incidence of MCI and AD, and in the rate of progression after a diagnosis of AD**

**Cognitive aging**

The shrinking of the brain and expansion of the ventricles is part of the natural maturational process of the brain during normal aging. Longitudinal studies that have used “normal” cognitive status as an inclusion criterion have found that even “normal” aging may contribute to subtle declines in cognitive functioning.11,12 There are significant sex differences in the normal aging process.13 The most consistent cross-sectional difference at all ages is that women perform better on verbal memory tasks and men perform better on visuospatial tasks.14–16 However, longitudinal studies have shown inconsistent sex differences, either reporting steeper annual rates of cognitive decline in men,17 women,14 or no sex differences.18 One of the major factors causing these inconsistent findings may be the cognitive reserve profiles (discussed later) of specific cohorts (eg, the Lothian Birth cohorts of 1921 and 1936).19

The trajectory of cognitive decline due to ongoing pathological insults to the brain (“pathological aging”) has been shown to deviate from the normal aging process. There is significant accelerated decline in cognitive functioning and brain volume loss years before the onset of MCI or dementia.20–23 However, there have been no systematic studies of sex differences in the cognitive decline prior to onset of clinical symptoms (ie, preclinical stages of the disease).

**MCI**

MCI is considered an intermediate state between the cognitive changes associated with aging and mild dementia, particularly of the Alzheimer type.24–26 Indeed, the risk of dementia is higher in persons with MCI compared to cognitively normal individuals.27–30 The prevalence of MCI in persons older than 65 years of age ranges from 10% to 20%, depending on the population studied and on the diagnostic criteria utilized.5,27,28,31–34 Some studies suggest a higher prevalence of MCI in men,6,35,36 while others suggest either a higher prevalence in women37,38 or no sex difference.39–40 The incidence rate of MCI has been estimated to be about 1%–4% per year in cognitively normal individuals aged 65 and older.27,40–42 Some studies examining the incidence of MCI also report that there may be sex differences, but reports vary based on the study design, diagnostic criteria, and age distribution of the sample. In general, women have a higher incidence of MCI at older ages. Men consistently have a higher incidence of the non-amnestic type of MCI.43 Amnestic MCI is defined as cognitive impairment that includes the memory domain whereas non-amnestic MCI refers to impairment in other domains (eg, executive functioning, visuospatial, language), but no impairment in memory. While amnestic MCI is considered prodromal for AD, non-amnestic MCI is considered prodromal for non-AD dementias, such as vascular dementia.25

**AD**

The prevalence of AD is significantly higher in women compared to men. Recent estimates suggest that almost two-thirds of the individuals diagnosed with AD are women. A reason for the higher prevalence among women may be that they live longer, on average, than men.44,45 By contrast, incidence studies examining sex differences in AD are equivocal. The majority of studies conducted in the US have not observed sex differences in the rates of developing AD.46–52 In contrast to these studies, the Cache County Study (Cache County, UT, USA), did report a higher incidence of AD in men than women until age 78, after which women had a higher incidence than men.53 Similarly, the Mayo Clinic Study of Aging recently reported that the rate of progression from
MCI to AD was similar in men and women aged 70–79, but higher in women than men after age 80.\(^\text{54}\) Consistent with these last two studies, most studies from European\(^\text{55-59}\) and Asian\(^\text{60,61}\) populations have also observed a higher incidence in women after the age of 80–85 years.

The reasons for these disparities across studies and geographic regions are not clear. Discrepancies could be due to the use of different diagnostic criteria for AD versus other forms of dementia, such as vascular dementia or Lewy body dementia. The differences may also be due to small sample sizes at the upper range of the age distribution, resulting in unstable estimates. Finally, some differences across Europe, Asia, and North America may be due to social, cultural, and historical events. For example, the impact of World War II and the following Cold War era have been very different across continents. Some of these historical events may have affected men and women differently. Notably, a meta-analysis of 13 studies of populations in the US, Europe, and Asia did show that women were at a significantly greater risk of developing AD, but not other dementias.\(^\text{62}\) Interestingly, women also have a faster rate of cognitive and functional decline after a diagnosis of AD.\(^\text{63,64}\)

### Sex differences in neuroimaging measures of brain reserve: structure and function

The concept of brain reserve posits that subjects with higher reserve have a greater capacity to cope with pathological insults than those with low reserve, and that these individual differences in reserve mechanisms help explain why cognitive decline may be initiated at different times in relation to the onset of pathology for each individual. Specifically, the concept of brain reserve stemmed from the observation by Katzman et al that subjects with larger brains have greater capacity to withstand more pathology at the same level of cognitive performance.\(^\text{65}\) Cognitive reserve is discussed later in the review.

### Structure

The most striking difference between the brain anatomy of men and women is the larger head size and cerebral brain volume in men (~10%).\(^\text{66}\) Therefore, one would expect men to be able to withstand more pathology compared to women. This hypothesis was supported by an autopsy study that found that women had significantly higher odds of a clinical diagnosis of AD at the same level of pathology.\(^\text{67}\) While overall larger head sizes may suggest larger brain reserve in men, studies have consistently shown faster age-associated brain volume decline in men compared to women in cognitively normal individuals.\(^\text{68-71}\) However, in patients with MCI and AD, brain volumes have been found to decline faster in women than men, supporting the evidence of faster progression of women from MCI to AD.\(^\text{72}\) Thus, even after considering differences in head size, sexual dimorphism in the brain anatomy exists.\(^\text{56,73-75}\) For example, women typically have a higher percentage of grey matter in several brain regions, whereas men have a higher percentage of white matter.\(^\text{76}\) While many of these differences are likely due to sex chromosomes and sex hormones,\(^\text{77-79}\) the exact mechanism through which sex hormones influence brain structures is still poorly understood. Notably, one major flaw of the studies examining changes in brain structure with age has been the modeling of brain volume loss over the life span without taking into account hormonal changes in men and women over time.

### Function

Functional imaging measures such as \(^{18}\)F-fluorodeoxyglucose positron emission tomography (FDG-PET) for measuring metabolism and resting state functional magnetic resonance imaging for measuring brain connectivity have shown significant differences between men and women.\(^\text{76,80}\) Typically, cerebral blood flow and connectivity have been found to be higher in women in the parietal association cortices and higher for men in the visual and motor cortices,\(^\text{81,82}\) providing evidence for brain function and behavior differences between the sexes. Several imaging studies have shown that sex differences in the brain circuitry contribute to significant performance differences on specific cognitive tasks; for example, men perform better on visually oriented tasks. In the context of cerebral metabolic deficits associated with cognitive impairment in dementia, two studies have shown that men have more pronounced cerebral metabolic deficits compared to women at the same level of cognitive impairment, suggesting that the greater brain reserve in men may be helping them withstand more pathology than women at the same level of dementia severity.\(^\text{83,84}\) Given the current hypothesis that regions of high connectivity in the brain harbor amyloid deposition,\(^\text{85}\) there is clearly a need to investigate sex differences in the pathological cascade of AD.

### Biological explanations for the sex differences

#### Genetics

While many studies have examined and reported the relationship (or lack of) between numerous genes and single-nucleotide polymorphisms (SNPs) and risk of AD,
few studies have specifically examined whether the relationships vary by sex. One reason for this is the high number of individuals needed for genetic analyses, and lack of power to examine sex differences, particularly for genome-wide association studies. As a result, studies adjust for sex rather than stratify by sex, or examine interactions with sex. The identification of the different genetic processes that may affect the risk of MCI and AD in men and women is imperative for individualized preventive and treatment plans.

The e4 allele of the apolipoprotein ε (APOE) gene is the strongest known genetic risk factor for late-onset AD. Compared to non-carriers, heterozygous carriers of one e4 allele are 3–4 times more likely to develop AD, whereas the risk for homozygous carriers is even higher. The APOE e4 allele is specifically associated with an earlier age of onset of AD. The majority of studies, including a large meta-analysis of 8,607 controls and 5,930 AD cases, have reported that the effects of the e4 genotype are more pronounced in women than in men. Three studies reported that women with one e4 allele had about a four-fold risk of AD, whereas men with one e4 allele showed little increased risk. The APOE e4 allele also has a greater deleterious effect on hippocampal pathology, functional connectivity changes in the default mode network, cortical thickness, and memory performance in women compared with men at different stages of AD. Additionally, a large autopsy study found that amyloid plaque and neurofibrillary tangle pathology was greatest among women who were e4 carriers.

Other genes and SNPs have also been shown to increase risk and progression of AD in one sex, but not the other. A large study consisting of 16 research centers worldwide (including 4,711 patients and 4,537 controls) reported that the Met66 allele of Brain Derived Neurotrophic Factor (BDNF) gene, which reduces the transport of BDNF, is associated with an increased risk of AD in women (odds ratio =1.14, 95% confidence interval 1.05–1.24, P=0.002), but not in men. This finding is biologically plausible since estrogen plays an important role in the expression of BDNF. Post-menopausal women with the MET66 allele would therefore have both reduced transport and expression of BDNF, thus causing an increased risk of AD.

SNPs found to pose a risk of AD among men, but not women, include a SNP (rs688) of the low-density lipoprotein receptor and functional apolipoprotein E receptor, the rs17571 SNP of the lysosomal protease cathepsin D, involved in signal transduction pathways, including insulin signaling, and PCK1 (catalyzes the first step in hepatic gluconeogenesis). Interestingly, a few SNPs have also been found to have an opposite predictive value for women compared with men. A diabetes-related gene, the G allele of NSP65 of the peroxisome proliferators-activated receptors gamma was associated with a significantly increased odds of AD in men, but a reduced odds in women. In contrast, the 219K allele of the ATP Binding Cassette Transporter 1 (ABCA1) gene had a 1.75-fold increased risk of developing AD in women, but was found to be protective in men.

The biological explanations for these sex differences are not fully understood, in part because the physiological effects of many of the genetic polymorphisms have not been completely determined. Most studies finding sex differences link the association to sex hormone levels. For example, some of the physiological benefits of estrogen have been linked to ABCA1-mediated pathways. However, there could also be gene–gene interactions (epistasis) of genes on an autosomal chromosome with genes on chromosome X or Y. With the continued observation of sex differences in the risk of AD for identified SNPs, better understanding of the resulting physiological changes that contribute to the sex difference is needed.

Hormones

Gonadal hormones act as critical neurotrophic factors in the perinatal period and throughout the lifespan. Both hormones and genetic differences (ie, X and Y chromosomes) contribute to the physiological mechanisms underlying sexual dimorphism of the brain, including neurogenesis, axon guidance, synaptogenesis, and neurovascular development. Following menopause, women experience relatively rapid loss of the ovarian sex hormones 17 beta-estradiol and progesterone. A bilateral oophorectomy prior to menopause causes an abrupt deficiency of estrogen, progesterone, testosterone, and a disruption of the hypothalamic–pituitary–ovarian axis. Men also experience significant declines in testosterone levels with age, but these declines are more gradual. Bioavailable testosterone declines 2%–3% per year after the age of 30. Because testosterone can be metabolized to estrogen, men do not have the severe estrogen loss, even in late-life, that is experienced by women after menopause or abruptly after a bilateral oophorectomy prior to menopause.

Animal and cellular models have consistently shown the neuroprotective effects of estrogen which include: improving synapse formation on hippocampal dendritic spines, maintaining hippocampal function during aging; improving cerebral blood flow and glucose metabolism, increasing choline acetyltransferase activity in the basal forebrain and hippocampus, (choline acetyltransferase is involved in the
synthesis of acetylcholine, a neurotransmitter reduced in AD and implicated in memory function), reducing the aggregation of amyloid-beta and associated neurotoxicity, and preventing mitochondrial damage. Despite the apparent benefits in animal and cellular models, the impact of estrogen loss (due to natural menopause or surgically induced) and of hormone replacement therapy (HRT), on the risk of AD in women remains controversial. To date, observational studies generally report reduced risks of AD in women who initiate HRT within a short period (typically <3 years) after natural menopause and after oophorectomy performed prior to menopause. For example, the Mayo Clinic Cohort Study of Oophorectomy and Aging showed an almost doubled risk of dementia in women who underwent bilateral oophorectomy before menopause. However, women who initiated HRT after the bilateral oophorectomy, and continued utilizing HRT at least until the age of natural menopause (approximately 51 years), did not experience an increased risk of AD. In contrast to these studies showing a beneficial effect of estrogen use, the Women’s Health Initiative Memory Study (WHIMS), a large randomized clinical trial of HRT, reported a two-fold increased risk of dementia in women randomized to HRT after age 65 years. One explanation for the differences between observational studies and clinical trials is that observational findings could be the result of confounding. Women who use HRT typically have a higher socioeconomic status, higher education, and/or better health and therefore may be at lower risk of AD. However, another possibility is the timing of the estrogen therapy.

Observational studies show that the use of HRT, when initiated around the time of menopause but not years after, reduces the risk of AD. In the Cache County Study, women who initiated HRT within 5 years of menopause had a 30% lower risk of AD compared to women who reported no use of HRT. However, women who began hormone therapy more than 5 years after menopause did not have a lowered risk. In fact, those who started hormone use when they were 65 years or older had almost a two-fold increased risk. Similarly, in both the Multi-Institutional Research on Alzheimer Genetic Epidemiology (MIRAGE) study and in the Northern California Kaiser Permanente study, initiation of HRT in mid-life was associated with reduced risk of AD, whereas initiation of HRT several years after menopause was associated with an increased risk.

In light of the observational results suggesting that the initiation of estrogen in the immediate years after menopause is protective, whereas later administration increases AD risk, the WHIMS trial results are not surprising. WHIMS subjects were aged 65–79 years old at baseline. Thus, HRT was initiated 10–20 years after the onset of natural menopause.

There are two ongoing hypotheses for the lack of benefit, or even detrimental effects, when estrogen is initiated years after menopause or bilateral oophorectomy. The first, “window of opportunity,” hypothesis is based on the mechanistic findings that long-term estrogen depletion (LTED) can cause decreased levels of estrogen receptor (ER)-alpha, in the CA1 region of the hippocampus, a highly responsive region to estrogen therapy, resulting in cognitive enhancement and neuroprotection. Therefore, the initiation of estrogen after LTED, when ER-alpha receptors are already down-regulated, does not result in the neuroprotective benefits of estrogen. The second, “healthy cell bias of estrogen benefit,” hypothesis suggests that estrogen only yields neuroprotective benefits when applied to healthy neurons. Neurons with damaged mitochondria, a feature of aging, will not benefit, and estrogen may even be detrimental under these conditions. It is likely that both hypotheses contribute to differential benefits of estrogen when initiated peri-menopausal compared to after LTED.

**Social explanation for the gender differences**

In addition to several biological explanations for the observed sex differences in the prevalence and incidence of MCI and dementia, the effects of sociocultural aspects, ie, gender differences, should also be studied. Gender refers to the cultural and psychosocial factors that impact our identity and modify our risk of disease via health perception, risk behavior, social and work-related stressors, personal and societal perceptions of men’s and women’s role, patient–doctor relationships, and adherence to therapy. Specific factors related to gender identity that may contribute to the risk of AD include education, occupation, diet and exercise, and smoking and drinking behaviors. Gender is also strongly linked with the concept of cognitive reserve such that a higher education/occupation and greater engagement in cognitive activities provides higher reserve against disease and results in varying cognitive aging trajectories among individuals. In this section, we discuss gender-related risk and protective factors for AD. All of these factors can be tied to the concept of cognitive reserve as proposed by Stern. This theory posits that subjects with higher cognitive reserve (eg, higher education, better diet, or less stress) may have a greater capacity to cope with pathological insults to the brain, or that it may take longer for them to reach the threshold of dementia detection. Thus, individuals with...
high cognitive reserve would be less likely to display the cognitive symptoms associated with dementia compared to individuals with the same pathology and low cognitive reserve.131,132

**Intellectual lifestyle: education, occupation, and cognitive activity**

Low education and low occupational history (eg, unskilled versus skilled worker) have repeatedly been associated with either a higher prevalence133–137 or incidence of AD.138–141 Cognitive activities have been shown to reduce the risk of dementia in the elderly.142,143 Intellectual lifestyle (education, occupation, and current cognitive activity) explains more than 10% of the variance in an individual’s cognitive performance.131 Innate cognitive ability is also important, and can lead to higher education and better occupation. Indeed, low childhood mental ability and IQ is associated with lower cognitive ability in late-life,144 with an increased risk of dementia,145 and with increased mortality.146

Recent Pittsburgh compound B positron emission tomography (PiB-PET) and FDG-PET imaging studies have also shown that subjects with higher education or occupational engagement have more pathological changes when compared to subjects with lower education at the same level of cognitive performance (ie, they have greater brain reserve).147–150 The mechanism by which low education and occupation are thought to increase risk of AD is by lowering cognitive reserve. A longitudinal study that followed 9,000 people semi-annually for 15 years found that the main effect of education was to increase the baseline cognitive performance of individuals.151 Thus, subjects with higher education take longer to reach the dementia threshold. Sex differences in cognitive reserve and risk factors will further bias studies by causing different thresholds for detection of disease.

In the past century men have had more opportunities for higher education and higher occupational attainment than women. This is particularly true for individuals aged 70 and older who are now at greatest risk of developing AD, suggesting a higher education/occupation related reserve in men. In contrast, women generally engage in more cognitive activities such as reading books, arts and crafts, group, and social activities. While these cognitive activities impact reserve, the effect is much less than the impact of education and occupation.131

Indeed, differential age, period, and cohort effects in educational and occupational attainment may play particularly important roles for late-life cognitive trajectories and risk of AD. The Seattle Longitudinal Study showed that individuals born in later (1914–1948) versus early (1886–1913) cohorts have better cognitive performance at the age of 70 years, and also slower rates of cognitive decline.152,153 Notably, the differences between the younger and older cohorts in cognitive gains were much greater for women than men. This research highlights the importance of gender-specific societal changes in intellectual lifestyle over time by cohort and specific historical periods (eg, during versus after World War II), and its subsequent impact on cognitive aging trajectories and risk of AD.

At the most recent census, the educational attainment in the US was higher in women than men,154 and there also has been a dramatic shift in occupational engagement due to changing gender roles. For example, men and women have experienced different access to education and occupation in North America compared to Europe and Asia in the early part of this century. These gender-related differences may explain the observed geographic differences in the prevalence and incidence of AD that are described above.146 Indeed, it is possible that with greater educational and occupational attainment in women, the sex differences will diminish. The changing trends of intellectual lifestyles in men and women may contribute to changing epidemiologic patterns for AD and dementia across countries and over time.

**Exercise**

Gender roles can affect exercise participation as parenthood and marital status have been shown to be significantly related to whether women exercise.155 Several studies suggest that exercise and cardiorespiratory fitness are associated with a reduced risk of MCI and AD156–158 and with a slower rate of decline after a diagnosis of AD.161 While women are thought to be more “health-seeking” than men, it has been estimated that American women tend to get less exercise than men over the lifespan.162 Studies on sex differences in exercise patterns and risk of AD are ambiguous. Some studies suggest that exercise lowers the risk of cognitive decline and AD more in women than men.163,164 In contrast, another study suggested that women who exercise tend to receive less of a protective effect than men.165 These conflicting results may be due to the stage of life in which exercise is measured because most studies have measured self-reported physical activity in late-life. One study of over 9,000 women collected self-reported information on physical activity when the women were in their teens, age 30, age 50, and in later-life.158 Physical activity at all time points was associated with a reduced risk of cognitive impairment in late-life. However, physical activity in the teenage years was associated with the greatest
reduction in risk. Women who were not active as teenagers, but who were physically active at age 30 and 50, also had a reduced risk, but not as much as those who were active as teenagers. Further, among women who were physically active as teenagers, late-life physical activity did not appear to further reduce the risk of cognitive impairment, suggesting that early activity, when the brain is developing, may be most important.

The benefits of teen activity on late-life cognition are likely multifactorial. Teens who are active have better cognitive performance. Youth physical activity may contribute to a cognitive reserve, similar to the effects of education, which would therefore have long-term effects on cognition. Physical inactivity in the teenage years is also associated with obesity and diabetes; both are risk factors for AD. Physical activity and exercise are much more strongly encouraged for girls and teenage women now compared to the early- and mid-20th century; time will determine the impact that this trend may have on the sex difference in the prevalence of AD. Notably, while there is increasing focus on exercise, overall lifestyle is becoming more sedentary. Low activity throughout the day may be more beneficial than 30 minutes of moderate physical activity combined with 10 hours of sedentary behavior. Little research to date has focused on sex differences in sedentary behavior and how these differences may relate to risk of AD.

Smoking

Acetylcholine is a neurotransmitter that is decreased in Alzheimer's patients. Indeed, the current US Food and Drug Administration-approved medications for AD primarily focus on inhibiting the degradation of acetylcholine. Nicotinic acetylcholine receptors are especially reduced in AD. Therefore, it was hypothesized that nicotine could be used to prevent or delay the progression of AD, and that smoking may be associated with a reduced risk of AD. Indeed, nicotine has been shown to increase cognitive performance in both animals and human smokers. However, the results of clinical trials examining the use of a nicotine patch in AD patients have been mixed with some studies showing reduced cognitive decline, and others showing no beneficial effect. Larger studies are ongoing. Although it is possible that nicotine could be beneficial for AD, cigarette smoking contains several other toxins, has carcinogenic effects, is a known risk factor for cardiovascular and pulmonary disease, and therefore may increase the risk of AD. Additionally, many smokers also drink, and the interaction between cigarette smoking and heavy alcohol use may be especially detrimental for cognition. A recent study suggests a strong interaction between smoking and alcohol use in predicting rate of cognitive decline, such that cigarette smokers who were heavy alcohol users had significantly faster rates of decline than smokers who were moderate alcohol users.

Cigarette smoking exacerbates Alzheimer's pathology in transgenic mice and rats, including amyloidogenesis, tau phosphorylation, neuroinflammation, and neurodegeneration. Among humans, initial case–control studies reported that cigarette smoking was associated with a reduced risk of AD. However, these studies may have been biased because smoking is strongly associated with cardiovascular disease and premature death. Thus, smokers who survive to old age, when they are at greater risk of AD, may be more resilient to the negative effects of smoking and aging-related diseases. Subsequent cohort studies, especially those examining smoking in mid-life, have found that smoking is a risk factor for AD. Cigarette smoking has also been associated with greater regional brain atrophy in cognitively normal individuals.

Some studies suggest that men who smoke are at greater risk of developing AD compared to women who smoke, whereas other studies did not show a sex difference. Traditionally, men have had a higher prevalence of smoking because it was more socially acceptable for men to smoke. It wasn’t until the 1920s and 1930s that more women began to smoke. However, smoking among women was still less than in men. In 1965, 51.9% of men versus 33.9% of women smoked. In recent years, the gender gap has been narrowing such that in 2009, 23.5% of men and 17.9% of women were current smokers. The differential changes in smoking by gender may impact the subsequent incidence rate of AD among women and men.

Conclusion

With the aging of the baby boomer generation, the prevalence of AD is reaching an epidemic size. By 2050, 14–16 million Americans will be diagnosed with the disease. Many more individuals will provide either formal or informal care for AD patients. There is currently no cure for this devastating disease. Current approved medications are symptomatic, and do not modify the underlying disease pathology. Although randomized clinical trials of medications to reduce amyloid and other targets are ongoing, a push towards understanding the factors associated with the risk and progression of AD is critical to identify possible preventive measures and potential new treatment targets. Future clinical trials of new therapies for AD should consider a deliberate stratification...
by sex, and should have adequate sample size to test for a therapeutic effect in men and women separately. A drug may have efficacy in only one sex, or the effect may be stronger in one sex.

A sex-specific or gender-specific focus in AD research is still not mainstream. However, as described in this review, the prevalence and incidence of AD, and brain structure and function, vary by sex and gender. There are also clear sex- and gender-specific risk factors for AD. Ignoring these differences will impede research and treatments. Further, this information is critical for predicting the future disease burden. For example, at the beginning of this century, men had higher education and occupational attainment. However, currently women, on average, have higher educational attainment than men. It is important to study these historical, social, and cultural trends to determine their impact on the future prevalence and incidence of AD. Understanding these sex differences and gender differences will help to define individualized treatment and preventive interventions for AD.

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