Review

Gender-Specific Degeneration of Dementia-Related Subcortical Structures Throughout the Lifespan

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Abstract. Age-related changes in brain structure are a question of interest to a broad field of research. Structural decline has 13 14 been consistently, but not unambiguously, linked to functional consequences, including cognitive impairment and dementia. One of the areas considered of crucial importance throughout this process is the medial temporal lobe, and primarily the 15 hippocampal region. Gender also has a considerable effect on volume deterioration of subcortical grey matter (GM) structures, 16 such as the hippocampus. The influence of age \times gender interaction on disproportionate GM volume changes might be 17 mediated by hormonal effects on the brain. Hippocampal volume loss appears to become accelerated in the postmenopausal 18 period. This decline might have significant influences on neuroplasticity in the CA1 region of the hippocampus highly 19 vulnerable to pathological influences. Additionally, menopause has been associated with critical pathobiochemical changes 20 involved in neurodegeneration. The micro- and macrostructural alterations and consequent functional deterioration of critical 21 hippocampal regions might result in clinical cognitive impairment-especially if there already is a decline in the cognitive 22 reserve capacity. Several lines of potential vulnerability factors appear to interact in the menopausal period eventually leading 23 to cognitive decline, mild cognitive impairment, or Alzheimer's disease. This focused review aims to delineate the influence 24 of unmodifiable risk factors of neurodegenerative processes, i.e., age and gender, on critical subcortical GM structures in 25 the light of brain derived estrogen effects. The menopausal period appears to be of key importance for the risk of cognitive 26 decline representing a time of special vulnerability for molecular, structural, and functional influences and offering only a 27 narrow window for potential protective effects. 28

29 Keywords: Aging, cognitive decline, gender, hippocampus CA1 region, subcortical grey matter

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INTRODUCTION

Age-related changes in brain structure are a question of interest to a number of different fields of research including neuroendocrinology, neurobiology, and neuroimaging, to just name a few. The

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growing body of research evidence has linked struc-35 tural alterations to certain functional and clinical 36 manifestations, including dementia-related disor-37 ders. Dementia has become a major health and public 38 concern worldwide with an increasing prevalence in 39 the aged population. The most common cause of 40 dementia in the general population above 60 years 41 of age is Alzheimer's disease (AD) [1]. AD is char-42 acterized by progressive behavioral, affective, social, 43 and cognitive impairment [2]. The neuropathologi-44 cal changes presumed to stand behind the functional 45 impairment are primarily the amyloid depositions and 46 the neurofibrillary tangles [3, 4]. These histopatho-47 logical alterations have been described in several 48 brain regions involving widespread frontal, pari-49 etal, and temporal cortical and subcortical structures. 50 Among these, medial temporal subcortical structures 51 are typically considered the most commonly empha-52 sized areas affected [5]. The most important risk 53 factor of developing AD that cannot be influenced 54 is age itself [6]. The most recent systematic review 55 and meta-analysis on prevalence and incidence of 56 dementia, and dementia due to AD found that increas-57 ing age was significantly associated with increasing 58 prevalence and incidence rates of dementia [7] and 59 AD [8]. Thus it appears crucial to understand the 60 age-related changes occurring in brain structures of 61 potential key importance. Large sample epidemio-62 logical studies show that women have a significantly 63 higher risk of developing AD for various reasons 64 (e.g., longer lifespan) [9-11]. Interestingly, incidence 65 rates appear to show and age-dependent relationship 66 between sex and likelihood of developing AD. Inci-67 dence of AD has been reported to increase with age 68 for both sexes until about 85-90 years but to continue 69 to increase among women only [12]. Therefore, gen-70 der is also considered a crucial unmodifiable factor 71 in AD pathology with clear differences in structural 72 and functional decline of specific brain areas. 73

This review will be focusing on age and gender dependent changes in grey matter (GM) microand macrostructures-and especially subcortical GM
formations—and related cognitive alterations as a
functional representation in AD pathology.

79 GREY MATTER ALTERATIONS 80 IDENTIFIED IN AD

A number of studies have addressed the neuroanatomical changes in the background of clinical symptoms presenting in AD. A recent large sample

meta-analysis has used anatomic likelihood estima-84 tion aiming to identify more robust and consistent 85 alterations [5]. GM atrophy has been found to pri-86 marily affect bilateral medial temporal lobe (MTL) 87 structures, involving the amygdala, hippocampus, 88 parahippocampal gyrus, uncus, and entorhinal cor-80 tex, as well as the thalamus, caudate, and cingulate an cortices [13]. Strikingly, one significant cluster in 91 the left MTL has been identified as a potential 92 anatomical marker for AD development and pro-93 gression. A robust GM loss has frequently been 94 documented in regions of the MTL bilaterally [14, 95 15]. Furthermore, the microstructure of the white 96 matter fibers in the close vicinity of the mediotem-97 poral structures are also affected by the disease [16]. 98 Hypometabolism as measured by PET studies and 99 hypoactivation as revealed by functional MRI have 100 also been reported [17]. Disrupted functional connec-101 tivity in these regions further supports the critical role 102 of MTL structures in the pathophysiology of AD [18, 103 19]. A main question of debate remains as to what 104 extent these changes reflect the course of the disease. 105 Research evidence indicates that relevant alterations 106 are present primarily in areas of the MTL several 107 years before the clinical signs of AD [20]. More-108 over, morphological abnormalities and atrophy have 109 been detected in the left MTL specifically as the most 110 consistent structure to predict conversion from mild 111 cognitive impairment (MCI) to AD [21]. Thus, based 112 on the pattern of structural atrophy, the left MTL has 113 been suggested as a marker of disease progression in 114 AD [5] (for a summary of referenced findings please 115 see Table 1). 116

AGE-RELATED CHANGES OF RELEVANT GM STRUCTURES

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A great body of research evidence confirms that aging is associated with decrease in total whole-brain volume [22–24], overall GM and white matter (WM) volume [25–29], as well as cortical thickness [30]. It seems evident to state that, parallel to total brain volume, the volume of subcortical brain structures in general decreases with age. However, evidence indicates that the changes are very different in specific brain areas [31, 32]. Even studies reporting no overall significant effect of aging on WM volume did reveal a decline with age in some areas [26, 33].

In order to understand the relevance of the structural loss, we have to decipher their complex neurobiological background and their effect

Age- a	and gender-related changes of medial temporal lobe, with the major focus on the hippocampus
Golomb et al., [160] Murphy et al., [50]	Size of hippocampal formation predicts longitudinal alterations of performance on memory tests. Larger age-related total GM volume loss and atrophy in frontal and temporal areas in males than in females,
	Greater atrophy in females than in males in hippocampus and parietal cortices. Hemispheric metabolic asymmetry in temporal and parietal cortices, Broca's area, thalamus, and also in hippocampus.
Raz et al., [43]	Largest age-related decline: volume of the prefrontal cortices. Slighter age-related alterations: volume of the fusiform gyri, inferior temporal, superior parietal areas. Weak effects of age on hippocampus and postcentral gyrus.
	Larger total brain volume and the hippocampus in males than in females.
Jack et al., [161]	Annual decline in hippocampal volume, increase in temporal horn volume was identified in the elderly. 2.5 times greater rates in patients with AD than in age- and gender-matched controls.
Xu et al., [60]	Larger atrophy with aging in right frontal lobe posteriorly in males compared to females. Age-related atrophy in right temporal lobe medially, in parietal cortices, cerebellum + left basal ganglia in males, but not in females. Smaller left thalamus, parietal, occipital cortices + cerebellum volume compared to the right barriers.
	to the right hemisphere. No age- and gender-related difference in this asymmetry.
Good, et al., [26]	Linear global GM volume loss with age, steeper decline in men.
0000, 01 ali, [20]	Accelerated loss bilaterally in the insula, superior parietal gyrus, central sulcus + cingulum. Little or no age effect in amygdala, hippocampus + entorhinal cortex.
Ge et al., [25]	Constant GM volume loss, linearly with age throughout adulthood, whereas delayed WM volume loss until midlife. No effect of sex.
Scahill et al., [24]	Acceleration in atrophy with age in all analyses, prominently after the age of 70, particularly in the ventricles and in the hippocampus.
Wang et al., [162]	Distinct patterns of hippocampal shape alteration with age, different patterns of hippocampal volume loss may distinguish mild dementia from healthy aging.
Sullivan et al., [52]	Linear thalamic volume loss with age in a similar pace in males and females, whereas more steep cortical GM volume decline during aging in men than in women.
Fleisher et al., [80]	Greater deleterious effect of APOE*E4 genotype status on gross hippocampal pathology and memory functions in women as compared to men.
Lemaitre et al., [55]	Between the ages of 63 and 75 years, largest GM atrophy in primary cortices + in angular gyri, superior parietal gyri, orbitofrontal cortex + in hippocampus. No sex × age interaction.
Ahsan et al., [42] Smith et al., [29]	Larger left caudate, nucleus accumbens + putamen, and larger globus pallidus in men. Relative regional differences in GM volume frontal, parietal + temporal cortices, no volume loss in medial temporal lobe and in posterior cingulate. No gender effects.
Sowell et al., [30]	Thicker right inferior parietal + posterior temporal cortices in females. Gender differences in these areas are detectable from late childhood and are maintained throughout life.
Curiati et al., [35]	Selective focus of accelerated GM reduction only in men, including temporal neocortices, prefrontal cortices, and medial temporal areas.
Neufang et al., [65]	 Larger GM volumes of left amygdala in males, larger right striatal GM volumes and hippocampal GM volumes bilaterally in females. Independently of gender, volumes of amygdala and hippocampus are associated with levels of circulating testosterone.
Ostby et al., [36]	From childhood until adulthood: non-linear decrease in GM in cerebral cortex, linear decrease in caudate, putamen, pallidum, nucleus accumbens, and cerebellum. Small, non-linear increase in amygdala and hippocampal GM volume.
Ystad et al., [163]	Hippocampal volumes are important predictors for memory function in elderly women. Hemispheric asymmetry in hippocampal volumes during aging. In females, volume of left hippocampus has predictive value. Gender and left hippocampal volume may predict verbal memory performance in healthy elderly.
Erickson et al., [82]	Limited time window for hormone replacement therapy to positively influence hippocampal volume.
Fjell and Walhovd, 2010 [38]	Heterogeneous pattern in the atrophy of specific brain areas during aging: largest shrinking in frontal and temporal cortices + in putamen, thalamus, and nucleus accumbens.
Mukai et al., [77]	Important role of hippocampus-derived estradiol in the modulation of synaptic plasticity.
Goto et al., [83]	Reduced GM volume in bilateral hippocampus in females in their fifties (most of them experiencing menopause) compared to females in their forties (most of them not experiencing menopause).
A1 . 1 F	\rightarrow Menopause may correlate with reduction of hippocampal volume.
Skup et al., [45]	Different patterns of decline with age in males and females in AD group and MCI group compared to healthy controls in precuneus and caudate nucleus bilaterally, right entorhinal gyrus, thalamus bilaterally, left incula, and also in right amygdala.
Takahashi et al., [51]	insula, and also in right amygdala. More retained GM concentrations in females during aging in inferior frontal gyri bilaterally, cingulate gyrus anteriorly, hypothalamus and in medial thalamus.

Table 1

(Continuea)	
Devanand et al., [164]	Differences in volumes of hippocampus, entorhinal cortex, and parahippocampal gyrus between MCI and healthy controls.
	In patients converting from healthy to MCI: larger atrophy in the head of hippocampus, specifically in CA1 and subiculum, in entorhinal cortex, especially in bilateral pole of EC.
Borghesani et al., [165]	Improvement of midlife memory positively correlates with larger hippocampal volume in the elderly, compared to those who had decline or no change in their episodic memory in their midlife.
Ooishi et al., [78]	Crucial role of hippocampus-derived estradiol, T, and DHT in modulating synaptic plasticity.
Rijpkema et al., [53]	No gender difference in caudate nucleus and nucleus accumbens.
	Larger globus pallidus and putamen volume.
Spencer-Segal et al., [79]	In females, important role of estrogen receptor signaling in hormone's influence regarding hippocampal synaptic plasticity.
Fjell et al., [34]	Faster estimated decline in the elderly in hippocampus.
Taki et al., [166]	Positive correlations between yearly regional GM volume alterations and age: temporal pole bilaterally, caudate nucleus, insula, hippocampus.
	Negative correlations between age and changes in cingulate gyri bilaterally + cerebellum.
	Age \times gender interaction between annual ratio of regional GM volume change in hippocampus bilaterally.
Crivello et al., [167]	Higher GM decline in females compared to males (persistent throughout age ranges)
	Hippocampus: similarly accelerated decline with age in males and females.
Li et al., [58]	Age-related atrophy in basal ganglia and thalamus.
	Hippocampus atrophy in males only, and no decline in the amygdala.
Perlaki et al., [57]	No sexual dimorphism in the size of hippocampus.
Kiraly et al., [56]	Larger hippocampus volume in females.
	Age-related decrease of caudate nucleus, putamen and thalamic volumes in males.
	Thalamic volume loss in females.
	Faster decrease in total GM volume in males as compared to females.

Table 1

on functionality. Fjell and his co-workers have 133 done tremendous work in an effort to character-134 ize cross-sectional and longitudinal changes in brain 135 aging and to compare healthy normal aging to 136 pathological alterations (i.e., the Alzheimer Disease 137 Neuroimaging Initiative) [34]. Fjell et al. have used 138 a nonparametric smoothing spline approach to assess 139 age trajectories of anatomical structures in a large 140 sample of healthy adults. Cross-sectional as well as 141 longitudinal, follow-up data has been analyzed iden-142 tifying certain critical age periods. These critical ages 143 would account for a more significant rate of change 144 within the estimated range of volume loss. Latter 145 has been described for total brain volume with a 146 stronger correlation above the age of 60, as well as 147 for the cerebral cortex, and, interestingly the pal-148 lidum, with the age of around 25 years correlating 149 most with structural decline. A linear reduction with 150 age has been identified for a number of subcortical 151 structures, i.e., the amygdala, nucleus accumbens, 152 putamen, and the thalamus, also supported by sev-153 eral previous findings [31, 35]. The hippocampus has 154 been previously characterized by a nonlinear pat-155 tern of estimated change through adulthood. This 156 might be explained by a prolonged phase of devel-157 opment [36], a longer stable period and, critically, 158 an accelerated volume loss starting around the age 159

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of 50 and an even more robust negative relationship above 60 [37–39]. Indeed, in the longitudinal analysis, the hippocampus showed the fastest rate of volume reduction (–0.83% per year) among subcortical structures [34]. Changes in brain volume constitute a truly dynamic process with a great number of potential influencing factors, which should be ideally monitored by using longitudinal approaches with a high density of assessments. Nevertheless, more complex and sophisticated methods of analysis as well as large volume data could yield more insight into targeted questions [40].

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Another highly dynamic process throughout the human lifespan is considered the interaction with and accommodation of constant endogenous and exogenous influences. The view of lifespan trajectories of change in brain structure and function might serve as a base of understanding vulnerability to certain agerelated disorders such as MCI and AD. It might be crucial to emphasize the potential significance of life course effects which, in a complex interaction, will eventually separate dementia and cognitive decline from normal aging-related mechanisms. However, it also appears that the relationship between different exogenous and endogenous events and their impact on brain structure and function varies in importance in the light of the time of their occurrence [41].

187 GENDER-RELATED CHANGES OF 188 RELEVANT GM STRUCTURES

Sexual dimorphism of the human brain anatomy
 has gained increasing interest, with subcortical GM
 structures also being investigated more widely [42].

A number of studies have addressed the combined 102 effects of age and gender on human brain structures. 193 A more profound decline in GM volume has been 194 described in males [33, 43, 44]. However, in patients 195 with MCI and AD, GM volume has been found to 196 decline faster in females as compared to males sup-197 porting the evidence of faster progression from MCI 198 to AD [45]. This might be related to the main dif-199 ference in brain anatomy between sexes, i.e., brain 200 size. A larger brain might well have a greater reserve 201 capacity to withstand pathology at the same level of 202 functionalilty and cognitive abilities [46]. This has 203 also been underlined by autopsy studies reporting 204 women to have significantly higher odds of a clin-205 ical diagnosis of AD at the same level of neuronal 206 pathology [47]. 207

The effect of gender on the volume of these 208 structures might be crucial, considering that basal 209 ganglia possess a high density of sex steroid receptors 210 [48]. However, neuroimaging results on the gender 211 dependent volume of subcortical GM are somewhat 212 contradictory. Some studies reported larger volumes 213 of the caudate nuclei [49], hippocampus [50], and 214 thalamus in females [51], while others had oppos-215 ing results [52, 53]. The amygdala [54], pallidum, 216 and the putamen [53] have been consistently found 217 to be larger in males. Thus, research evidence appears 218 inconsistent especially considering the subcortical 219 GM structure [55]. This might also be due to the 220 method of analysis, considering the difficulty to 221 delineate subcortical GM using conventional voxel 222 based morphometric methods. Our research group 223 has applied a deformable surface model based seg-224 mentation approach to address volumetric alterations 225 especially in regions with low tissue contrast [56]. 226 While age, gender, and head size (intracranial vol-227 ume) are the most commonly included 'nuisance' 228 variables when performing neuroimaging analysis, 229 studies vary as to which of these variables are 230 included and which method is used for correction 231 [57]. These factors might widely account for the 232 great variability in the results. Accounting for skull 233 size significantly influences results when it comes to 234 GM volume and it might be of even greater impor-235 tance when considering differences between males 236

and females. Our results revealed larger cortical and subcortical GM volume for females as a result of correction for total intracranial volume in a study involving 103 participants in the age range of 21–58 years. The volume of the hippocampus was found significantly larger in the female group as compared to males. We also detected a significant effect of hemisphere in the male group only, with larger volumes of the right caudate and the left thalamus as compared to their contralateral structures.

Interestingly, we also found an age-dependent decrease in the volume of cortical as well as subcortical GM. Latter remained significant after correction for skull size in the caudate, putamen, and thalamus bilaterally for males and the thalamus bilaterally for females. Within the age range of 21 to 58 years, we found a linear decrease in GM volume with aging. Strikingly, this process proved to occur at a faster pace in males. Converging research evidence emphasizes the importance of considering age and sex interaction effects on the volumetric decline of subcortical structures. Li and his colleagues found this to be of key relevance for the hippocampus specifically, showing a linear negative correlation with age for males only [58]. Strikingly, for females, the pace of hippocampal volume decline has been found to occur at an even slower pace than whole brain volume loss. In contrast with this, a strong effect of aging on basal ganglia and thalamus volume changes has been observed primarily for females. The authors link these results to functional consequences involving predominantly psychomotor performance especially at later ages [59–61]. However, a number of studies did not find a significant effect of gender on cognitive performance or decline with age [62, 63]. While directly linking functional aspects to structural changes in brain anatomy might not be equivocal, elucidating effects of age \times sex interaction on specific subcortical GM regions might well serve the investigation of related psychopathological alterations, such as MCI or AD.

The background of the disproportionate GM volume changes has not yet been elucidated, but the changes in hormone levels and the consequent sensitivity of the brain to hormonal effects are most certainly involved [64]. Sex hormones have been found to critically influence regional maturation of subcortical GM structures, e.g., higher circulating testosterone levels correlated positively with amygdala volume and negatively with hippocampal volume [65]. Estrogen among androgens has gained significant interest for its crucial role

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during brain development. Females with endogenous 287 estrogen deficiency have been found to have dis-288 proportionately reduced hippocampal volumes and 289 increased amygdala volume as compared to age-290 matched controls [66]. This might be related to the 291 complex distribution of estrogen receptors through-202 out the brain. Distinct estrogen receptor subtypes 203 have been identified in nearly all cell types of the 294 central nervous system, and importantly, in brain 295 regions typically associated with cognitive func-296 tion such as memory and affective processing, e.g., 297 the amygdala and the hippocampus [67]. Strikingly, 298 the estrogen-related volume deficiency evidenced by 299 structural neuroimaging has also been associated 300 with functional consequences revealed by cognitive 301 assessment [68]. 302

Epidemiological results support the notion that 303 age-related loss of steroid hormones is associ-304 ated with an increasing risk to develop AD [69]. 305 Above this, AD prevalence is higher in post-306 menopausal women as compared to age-matched 307 men-not explained by the generally higher life 308 expectancy for females [70, 71]. The crucial role of 309 estrogen is supported by several lines of evidence, 310 with early menopause having been associated with 311 an increased prevalence of dementia [72]. Estro-312 gen has been found to modulate neurogenesis and 313 activation of new neurons in response to targeted cog-314 nitive demands in the hippocampus [73, 74]. This 315 might be mostly dependent on brain derived estra-316 diol concentration [75], suggesting the importance 317 of neuronal, and especially hippocampal, estrogen 318 production [76]. Estrogen has a potent effect on 319 inducing neurogenesis, neuronal morphology, and 320 plasticity in specific areas of the hippocampus, 321 such as the CA1 region and the dentate gyrus [74, 322 77-79]. An association between estrogen deficiency 323 and hippocampal volume loss in females with clin-324 ically diagnosed MCI [80] might well serve as a 325 potential common course leading to AD. However, 326 there might be another crucial aspect, which should 327 be emphasized when considering neuronal estro-328 gen related hippocampus structure and function. A 329 significant sex hormone cycle related effect on spe-330 cific cognitive performance has only been found 331 during initial testing and disappeared with repeated 332 examinations of the same parameter, controlling for 333 other confounding factors [81]. This occurred dur-334 ing an 8-week long testing period, which raises 335 interesting questions about a life course perspec-336 tive of hippocampus-related cognitive performance 337 and the risks of consequent dementia. Furthermore, 338

hormone treatment effects on the hippocampus in post menopause detected a limited window of opportunity to influence hippocampal volume. However, the larger hippocampal volumes associated with hormone treatment initiated at the time of menopause did not translate to improved cognitive performance [82].

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Hippocampal volume loss appears to become accelerated in the postmenopausal period [83], which, associated with brain estrogen production decline, might be due to a significant reduction in neuronal plasticity primarily in the CA1 region. While postmenopausal hormone replacement therapy might spare the total hippocampal volume in a limited window of action, this might not be effective on the key areas of neuroproliferation. Consecutively, cognitive performance is not affected beneficially, eventually leading to the development of MCI or AD, due to the impaired cognitive reserve abilities influenced by several other factors (Fig. 1).

FUNCTIONAL CONSEQUENCES OF GM CHANGES RELEVANT FOR DEMENTIA OCCURANCE

Above the structural differences, there is increas-362 ing evidence for the functional sexual dimorphism of 363 subcortical structures. Hippocampus-related memory 364 functions are differently affected by stress in males 365 and females [84]. Peripartum hormonal changes are 366 known to modulate the hippocampal function [85]. In 367 addition to gender effects, recent evidence supports 368 the influence of brain hemisphere showing lateral-369 ization of structure-function relationships, as well as 370 more specific relationships between individual struc-371 tures (e.g., left hippocampus) and functions relevant 372 to particular aptitudes (e.g., vocabulary) [86]. Numer-373 ous differences between the cognitive patterns of the 374 two sexes have been reported [87]. Estrogen and 375 testosterone appear to play a significant and contin-376 uous role in cognition throughout the lifespan [58]. 377 In puberty, adolescents who mature later have better 378 visuospatial skills than those who mature earlier [88]. 379 Furthermore, a longer reproductive period is associ-380 ated with higher levels of verbal fluency later during 381 adulthood [89]. In adulthood, certain differences 382 between male and female cognitive features are well 383 known, e.g., higher performance on visuospatial tasks 384 in males and female advantage in verbal skills [90]. 385 This characteristic pattern of different cognitive abil-386 ities appears to persist later in life [91]. Interestingly, 387

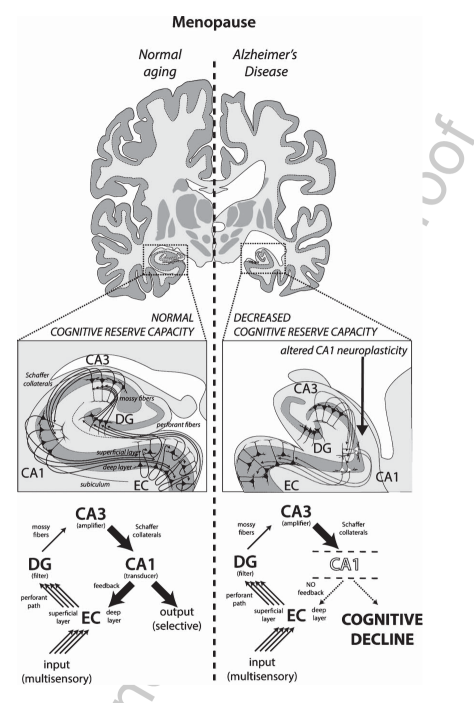


Fig. 1. According to major communication pathways of the hippocampal circuit multisensory input information enters primarily the entorhinal cortex (EC) then projecting towards the dentate gyrus (DG) and the CA3. Pyramidal cells of the CA3 send their axons to the CA1, which then projects to deep layers of the EC and sends the selected information along the output paths of the hippocampus. Additionally, feedback is being provided to the EC. The postmenopausal period and related estrogen loss might be associated with changes in the neuroplastic capacity of especially vulnerable regions of the hippocampus, such as the CA1 region. This region is rich in brain derived estrogen receptors and represents a key area for estrogen related neuronal manifestations. Molecular and pathobiochemical alterations might be present in the background of this deterioration, i.e., mitochondria-related inflammatory, oxidative effects. As a consequence, the selection of relevant information might become impaired or completely altered. In addition, the feedback source of the EC representing the major multisensory input area also becomes disturbed or even absent. In the presence of an impaired cognitive reserve capacity related to several previous internal and external factors, this might be an especially vulnerable time window for hippocampal structural and functional decline. This could result in an accelerated volume loss of the hippocampus and presumably, a consequent significant cognitive decline.

cognitive skills of women tend to decline slower
than those of men [92]. Estrogen has been
suggested as a protective factor against dementia
through facilitating neurogenesis in the hippocampus and thus enhancing hippocampus-related spatial
learning and aspects of memory [74].

Distinguished patterns of cognitive skills were con-304 firmed not only in healthy aging, but also in patients 395 with AD. Assessing AD patient's verbal skills, a 396 meta-analysis revealed a difference in naming tasks 397 and semantic fluency with lower performance in 398 women [93]. As to visuospatial skills, no significant 399 difference was found between women and men with 400 AD [94]. Based on another meta-analysis assessing 401 global dementia severity in men and women, it was 402 found that women reached a significantly lower score 403 compared to men with AD [95]. 404

Apart from the individual's sex and its hormonal 405 influences on cognition through the lifespan, other 406 contributing factors might enhance or prevent cogni-407 tive decline and developing AD. According to a recent 408 cohort study, lower performance in school during 409 childhood may increase the risk for cognitive decline 410 in later life [96]. Greater midlife stress is associated 411 with a higher risk to develop dementia, especially 412 AD among women [97]. Strongly negative life events 413 such as losing a close relative can also increase vul-414 nerability to enhance cognitive decline along with 415 depression; however, milder but chronic stress factors 416 may even stimulate cognitive functioning [98]. 417

Brain areas typically affected in MCI and AD 418 have a specific hierarchical order in which they 419 become altered during the course of the disease based 420 on Braak and Braak's neuropathological model [3]. 421 According to this model, the first lesions can be 422 detected in the MTL, including the hippocampus, 423 parahippocampus, and crucial areas of the limbic 424 circle, e.g., the amygdala, then in several areas of 425 the temporal lobe, followed by other regions of the 426 neocortex. The affected structures have their distinct 427 roles in cognition; however, they contribute alto-428 gether to the characteristic clinical manifestation of 429 AD. As an example of key importance, higher visual 430 perception, including identification and recognition 431 of faces and landmarks, as well as recognition of 432 facial emotions, is dependent on the medial temporal 433 lobe structures [99]. The impairment of these abili-434 ties might have an impact on behavioral disturbances 435 in early AD and might even serve early identification 436 of AD [100]. 437

Being a key structure of the MTL and its memory
 network, the integrity of the hippocampus is required

not only in episodic and semantic memory, but also 440 in spatial information processing and manipulation 441 [101]. The reduced ability to retain new information is 442 one of the earliest core features of dementia and con-443 stitutes a heavy burden on the daily life of patients and 444 caregivers [102]. A significant correlation of reduced 445 hippocampal volume combined with higher levels 446 of cortisol and performance on auditory and ver-447 bal memory subtests of the Wechsler's Intelligence 448 Scale and Block Design tests measuring visuospa-449 tial skills has also been reported [103]. A recent 450 study describes decreased thickness of the hippocam-451 pal GM formation in AD as compared to healthy 452 individuals or patients with MCI [104]. Considering 453 that scores on the Mini-Mental State Examination 454 (MMSE) and the Alzheimer's Disease Assessment 455 Scale-Cognition (ADAS-Cog) correlate with base-456 line entorhinal cortex thickness, its atrophy might 457 be a predictor of subsequent cognitive impairment. 458 The atrophy of hippocampal areas has been asso-459 ciated with more severe deficits in several aspects 460 memory (especially episodic memory) and execu-461 tive function [105]. Associated with lower activity 462 in these areas, AD patients have demonstrated poorer 463 encoding and retrieval than healthy individuals [106]. 464 Simultaneously, increased activation in ventral lateral 465 prefrontal areas may be interpreted as a compensatory 466 mechanism in AD. 467

When considering the broader picture of cognitive disturbances already detectable in early stages of dementia, several other areas need to be mentioned. The thalamus, as a key area of the limbic circuit and the episodic memory network, has also been reported to be affected in early stage AD [107]. Alterations of the amygdala appear to have a profound effect on emotional aspects of memory in AD [108, 109]. Emotional stimuli, especially those with negative valence, have altered influence on memory functions in AD patients [110] and amygdala atrophy has been correlated positively with emotional memory impairment severity [111]. Some recent studies even pointed out other complex functions of the MTL, including path integration, e.g., spatial representation, self-motion sensing, and temporal processing [112]. Lesions of the anterior areas of the hippocampus, parahippocampus, amygdala, and the anterior and lateral section of temporal gyrus are associated with poor performance on tests of delayed memory, long-term memory and spatial memory. Additionally, patients with alterations of these structures have difficulties in target-directed walking because of deficits of allocentric spatial information processing.

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The picture is certainly much more complex and it becomes increasingly difficult to decipher a causal relationship. Nevertheless, the role of the hippocampal region appears to be crucial in the occurrence and progression of the cognitive impairment in MCI and AD.

It is debated whether the extent of MTL structural 498 atrophy is a better predictor of clinical dementia as 499 compared to the memory deficit. Some studies found 500 that the ratio of amygdala volume loss and bilat-501 eral entorhinal cortex shrinkage predicted time until 502 MCI symptom occurrence [113]. Others, for example 503 Visser et al., reported scores on cognitive test batter-504 ies to serve as better predictors than MTL atrophy in 505 a longitudinal study design [114]. 506

Considering that the volume of subcortical GM 507 critically impacts the size of neurons, glia cells, and 508 number of synapses it entails, we might hypothesis 509 that it affects the function and performance of these 510 structures. While deducing cognitive or any other 511 type of functional activity of subcortical GM solely 512 from their structural characteristics would be inad-513 missibly simplified, observing changes in volume of 514 subcortical GM influenced by gender and aging might 515 yield better insight into several pathological condi-516 tions, e.g., MCI and AD [115]. 517

TRANSITION FROM HEALTHY AGING TO MILD COGNITIVE IMPAIRMENT AND AD

MCI is considered a precursor stage of AD with an 521 annual conversion rate of approximately 15% [116]. 522 However, the clinical manifestation of MCI is still 523 not considered a predestination of a future conver-524 sion to AD. One of the crucial biomarkers proposed 525 in the aim of a more valid diagnostic construct is 526 MTL atrophy [117]. A large number of studies have 527 focused on hippocampal volume loss focusing on 528 MCI conversion to AD reporting a non-uniform pat-529 tern of hippocampal shrinkage. Converging research 530 evidence emphasizes the key role of the CA1 region 531 and subiculum showing the most significant involve-532 ment throughout disease progression early on in the 533 course of illness [118-124]. While hippocampus vol-534 ume has been reported to hold the highest predictive 535 accuracy for conversion to AD, the best multivariate 536 model for AD prediction, interestingly, consisted of 537 cognitive variables only [125]. 538

A potential explanation for this seeming discrepancy might be related to methods of imaging analysis

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with more advanced techniques needed to ascertain reliable and accurate data processing. The radial atrophy technique used to investigate subtle changes in distinct regions of the hippocampus might be a useful method in addressing prominent volume loss prior to clinical pathology. Here, the CA1 region might be of crucial importance, considering its robust volumetric loss above the age of 60 also compared to other regions of the hippocampus. However, if this is true for the normal aging process, what could then be the key turning point that eventually leads to the outcome of dementia?

A view that gains increasing support offers an explanation relying on neuroplasticity. Brain regions characterized by high neuroplasticity have been found to be especially vulnerable to neurodegeneration as well [126-128]. The CA1 region of the hippocampus maintains its neuroplastic flexibility well into adulthood presumably serving cognitive capacity in interaction with external and internal demands. Converging evidence supports the finding that high level abilities of neuroplasticity are retained late in life [129–131], especially in areas with long axonal connections, such as the hippocampal region [127]. The neurons in these regions might be able to maintain their morphological and functional flexibility to serve cognitive processes, however, these abilities might on the other hand increase their vulnerability to neurotoxic effects eventually resulting in structural and functional decline [132, 133]. The hippocampal region is undoubtedly a key area for high-order cognitive processes, such as memory and learning, associated with high demands for neuroplasticity and neuronal flexibility [134, 135]. In addition to this, other neuronal morphological processes, such as dendritic spine plasticity, might also play a crucial role in cognitive flexibility throughout the lifespan [136]. This mechanism might be involved in cognitive processes related to the CA1 region of the hippocampus [137, 138]. However, this might also be a vulnerability component for pathological effects, i.e., disturbed neurogenesis and neuronal flexibility in the hippocampus has been suggested as a crucial early component in cognitive decline and even AD [139]. The relatively rapid structural decline observed in postmenopausal women in these vulnerable regions might further accelerate the deterioration resulting in a vicious circle [140]. This is supported by findings of an age \times gender \times subcortical structural dependent interaction with an impact on cognitive reserve abilities [141].

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RELEVANT MICROSTRUCTURAL AND PATHOBIOCHEMICAL CHANGES IN THE BACKGROUND OF STRUCTURAL AND FUNCTIONAL DETERIORATION

In the light of the presumably impaired neuro-596 plasticity consequently leading to macrostructural 597 changes in the hippocampal formation, one has to 598 certainly address the microstructural neuropathology 599 behind it. Focusing on specific hormonal effects, it 600 has been shown that neuronal substrates associated 601 with cognitive decline are significantly impacted by 602 estrogens [142]. Research evidence indicates that 603 most of estrogens' neuronal effects are related to 604 brain derived estrogen, synthetized within the cen-605 tral nervous system [143, 144]. While levels of brain 606 estrogen might largely differ from that of circu-607 lating estrogen, female brain estrogen levels have 608 been found to relate well with blood estrogen lev-609 els measurable on the periphery [145]. Strikingly, a 610 significant decline in brain-derived estrogen charac-611 terizes the postmenopausal period. It has also been 612 suggested that this decline occurs mainly around 613 menopause and, paired with a significant reduction 614 in brain derived estrogen synthesis, it might lead to 615 consequent cognitive deterioration [146, 147]. One 616 key neuronal substrate that integrates several estrogen 617 regulated molecular pathways is the mitochondria 618 [148-150]. Estrogen receptors have been found in 619 the mitochondria and the key role of mitochondria 620 in estrogen associated neuroprotection has been sup-621 ported by several different lines of evidence involving 622 anti-inflammatory actions, anti-oxidant effects, and 623 glutamate-related mechanisms among others (for an 624 excellent review, see [151]). New evidence also 625 indicates that a mitochondrial estrogen receptor defi-626 ciency found in the female AD brain results in 627 impaired anti-inflammatory and anti-oxidative capac-628 ity of the mitochondria indicating vulnerability for 629 neurodegeneration [152]. Our research has focused 630 on the mitochondrial disturbances critical in aging, 631 neurodegeneration, and AD specifically also involv-632 ing the kynurenine system [153-155], glutamatergic 633 mechanisms [156], and bioenergetic effects [157]. 634 The complex interaction of these processes might 635 well serve as a pathobiochemical and molecular 636 background for the structural and functional alter-637 ation described in neurodegeneration. This is also 638 supported by the relationship between worse patho-639 logical changes (i.e., amyloid depositions and total 640 tau levels) and a more rapid hippocampal atrophy and 641 cognitive decline in females, marking a potentially 642

increased vulnerability for the clinical manifestations of MCI and AD [158]. In the female brain, the menopausal period brings deterioration in the above mentioned bioenergetical balance with a potential lack of compensatory mechanisms representing a vulnerability to cognitive decline [159].

CONCLUDING REMARKS

AD is a growing healthcare issue worldwide demanding more and more precise characterization and identification of potential turning points from healthy aging to MCI and AD. An increasing body of research evidence has confirmed specific subcortical GM alterations in the brain during this process, evolving based on a hierarchical model. The firstly affected and most crucial areas are the components of MTL, especially the hippocampus. Endogenous and exogenous factors interacting with each other contribute to continuous alterations of these areas from our birth throughout adulthood. There are non-modifiable variables, such as age and gender, which have specific effects during aging, involving hormonal influence. In women, hippocampal volume loss appears to be accelerated in the postmenopausal period. This volume loss might be associated significantly and in a beginning stage with the neuroplasticity of the CA1 region in hippocampus, considering its high sensitivity to pathological alterations. The atrophy and consequent structural decline and functional impairment of this region evolving to other hippocampal and MTL areas might lead to the clinical manifestation of cognitive decline. This risk might be the greatest in the case of an already narrowed cognitive reserve capacity or subclinical cognitive impairment. Serving as a potential biomarker, specific structural hippocampal changes might be associated with consequent functional patterns of cognition, potentially supporting the identification of MCI and AD prior to the clinical symptoms of the disease. The interaction of age and gender combined with individual variables such brain-derived estrogen receptors, bioenergetical balance, and compensatory mechanisms should be taken altogether into consideration when assessing a potential occurrence of MCI and AD.

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