Brain Structural and Functional Changes in Cognitive Impairment Due to Alzheimer's Disease

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Cognitive neuropsychology seeks a potential alignment between structural and functional brain features to explain physiological or pathological processes, such as Alzheimer’s disease (AD). Several structural and functional brain changes occurring during the disease, including cognitive impairment, are found at the end of the patient’s life, but we need to know more about what happens before its onset. In order to do that, we need earlier biomarkers at preclinical stages, defined by those biomarkers, to prevent the cognitive impairment. In this minireview, we have tried to describe the structural and functional changes found at different stages during AD, focusing on those features taking place before clinical diagnosis.

Keywords: structural changes, functional changes, early markers, reversion, cognitive impairment (CI)

INTRODUCTION

Alzheimer’s disease (AD) is the most prominent among neurodegenerative disorders, being responsible for structural and functional brain changes that may result in the functional changes that can be found during the development of the disease (Masters et al., 2015; Jack, et al., 2018).

We have an extensive knowledge of the features of the disease, but we are unable to cure or prevent it. Indeed, when a clear clinical diagnostic arrives, it is too late to prevent the disease, whose diagnosis is only definitely confirmed when the hallmarks of the disease are characterized at brain level in the autopsy after patients’ death, as firstly described by Alzheimer (1907) (Alzheimer et al., 1995).

Thus, we know several brain structural and functional changes occurring during the disease that are found only after the patient’s life, but we need to know more about what happens not only during the development and the end of the disease, but before its onset. In order to do that, we need earlier markers that could be obtained from liquid samples [blood or cerebrospinal fluid (CSF)] (Barthélemy et al., 2020) or image analysis like PET (Vlasenko et al., 2021), which in the case of tau pathology (Villemagne et al., 2015) can recapitulate Braak stages found in patients’ autopsies. These earlier markers, complemented with functional analysis, are needed to prevent the disorder.
In this minireview, we have tried to start describing the structural and functional brain changes found in the disease, and those features taking place before clinic diagnosis.

We also comment on the possible alignment between structural and functional brain changes that may occur at different times in the life of a human being, that could end in dementia. Many structures can be involved in AD development. Indeed, brain regions involved in the Default Mode Network (DMN) may play a role (Alves et al., 2019) at different time and levels.

We propose, as it has been indicated by others, that long longitudinal studies to follow those possible changes several years before dementia onset will facilitate the step-by-step analysis of AD development. Since some of those analyses have already been carried out, we will comment on some of the obtained results looking at structural and functional time-dependent brain changes.

**STRUCTURAL CHANGES RELATED TO AD DEMENTIA**

As indicated by Alzheimer et al. (1995), the presence of two main aberrant structures, senile plaques and neurofibrillary tangles, precede the neuronal death and brain degeneration found later on the disease. Plaques are aggregates of beta amyloid peptide that progressively appear during the development of the disease following a specific pattern (Thal et al., 2002). This pattern starts in the neocortex and continues through the allocortex, hippocampus, basal ganglia, midbrain, and cerebellum. This pathway starts many years before clinical diagnostic (Bateman et al., 2012). Later on, it can be found by the presence of neurofibrillary tangles (tau protein aggregates), that also propagate, but follow a different pathway (Braak and Braak, 1991), starting at the transentorhinal region and expanding through the entorhinal cortex, hippocampus and neocortical areas. In this way, there are two different pathologies (amyloid and tau pathologies, with different structural changes) that appear at different times during the development of the disease. Indeed, it has been suggested that the disorder can become a unique disease when both amyloid and tau pathologies overlap (Hojjati et al., 2021). Before that time, different features may be related to the presence of only plaques in some brain regions.

**FUNCTIONAL CHANGES RELATED TO AD DEMENTIA**

Memory loss, cognitive impairment, loss of executive functions, and loss of consciousness, among others (Masters et al., 2015; Jack, et al., 2018), occur in AD dementia. They can appear at different times during disease development, step by step through a continuum that ends in dementia and an extensive brain degeneration. Being aware of one’s surroundings and considering social behavior as part of the interaction with that environment, it is important to note that it changes with aging (Rosati et al., 2020), which is relevant taking into account that aging itself constitutes the main risk for dementia. Indeed, there is a loss of consciousness, related to loss of individual awareness or awareness related to the world around those future patients.

**MILD COGNITIVE IMPAIRMENT**

We now know that before dementia there is a mild cognitive impairment (MCI), that could be more related to tau pathology. At the end of the past century (Petersen et al., 1999), it was suggested that before dementia and probably related to the onset of the first Braak stages, MCI could result in changes in cognition, while maintaining the capacity for executive functions and the independence to carry out daily activities. MCI definition can be classified into two different types, amnesic and non-amnesic. The first one is more related to memory changes, the second one may maintain an intact executive functionality (Carmasin et al., 2021). Moreover, MCI could be related to Braak stages 1 and 2 and CA1 region could be involved in the appearance of MCI.

The existence of a so-called AD continuum could indicate the presence of MCI before AD (Petersen et al., 1999). Although there are several types of MCI, including amnestic, non-amnestic, and mixed, that could behave differently to progress into dementia, we will mainly comment on MCI as a whole.

Around 10–15% of subjects with MCI could progress to dementia per year (Petersen, 2000) and it has been estimated that overall more than 40% of subjects with MCI could develop dementia (Panpalli Ates and Yılmaz Can, 2020). Thus, it is paramount to know why the other 60% do not progress similarly.

The percentage of transition from MCI to dementia depends on factors like age, education, family history of dementia, vascular risk factors or ApoE4 status (Kryscio et al., 2006). Also, lifestyle-related factors like alcohol consumption have a role on this proportion (Xu et al., 2009). Some of these factors could be modified to prevent the development of the disease (Sanz-Blasco et al., 2021).

**SUBJECTIVE COGNITIVE DECLINE**

It has been suggested that subjective cognitive decline (SCD), expressed by a frequent confusion and transitory memory loss could be a cognitive decline without being an objective (testable) mild cognitive impairment. Thus, it has been suggested that SCD could be a previous step to MCI (Jessen et al., 2014).

**TRANSITION FROM SUBJECTIVE COGNITIVE DECLINE TO MILD COGNITIVE IMPAIRMENT**

In a 7-year longitudinal study, it was described that around 20% of SCD subjects can progress to MCI.
A working hypothesis for the AD continuum, starting with a putative change in consciousness, related to amyloid pathology, followed by features that could facilitate the development of the continuum like chronic stress, SCD, and MCI. The probability of transition (upper arrows) is increasing step by step. These steps could be reverted but not that from AD (dementia), with an inversion probability (lower arrows).

(Bessi et al., 2018). Similar results were found in other studies (Avila-Villanueva and Fernandez-Blazquez, 2017).

Thus, SCD is a clear risk factor for MCI, like MCI is a risk factor for dementia.

Again, it will be of interest to identify the causes for the transition of that 20% SCD subjects to MCI, looking for a possible prevention.

**EXECUTIVE FUNCTIONS**

In addition to episodic memory loss, related to changes in the CA1 hippocampal region, a main feature on the development of AD is the loss of executive functions, like planning, working memory, self-control, flexible thinking, or organization. Executive functions have been mainly located in prefrontal regions (Stuss, 2011), although other regions like nucleus accumbens (NAcc) could also play a role in such functions (Floresco, 2015; Prasad, 2018). More recently, this role has been proposed again (Jenkins et al., 2021). Since hippocampal CA1 can connect to NAcc (Zhou et al., 2020), a damage in CA1 could later-on have an effect on NAcc and on executive functions or specific types of memories (Prasad, 2018).

In this way, it will be of interest to know if some features of cognitive decline related to CA1 damage could take place, or not, earlier than those specific executive functions. Further analysis should be done to test if it is the case.

**IS THERE A CHANGE BEFORE SUBJECTIVE COGNITIVE DECLINE? COULD BE THAT CHANGE CHRONIC STRESS**

We have previously discussed the role of chronic stress as a trigger for the AD continuum, being a possible step before SCD (Avila-Villanueva et al., 2020).

Structurally, chronic stress may affect structures like amygdala (Liu et al., 2020). Amygdala could activate other brain areas,
such as hypothalamus and brainstem, altering prefrontal cortex (PFC) function (Arnsten, 2009). Also, chronic stress may induce changes in the sympathetic nervous system altering the hypothalamic-pituitary-adrenal axis and producing an increase of cortisol, a compound that can cross the brain-blood barrier and is able to bind to hippocampus, amygdala, or prefrontal cortex receptors (de Kloet et al., 1999; Li et al., 2019). Thus, damage in those structures may be a previous step to SCD. Indeed, people with chronic stress in midlife could have a higher risk for SCD and MCI (Avila-Villanueva et al., 2020). This also agrees well with the fact that subjects with SCD tend to have a higher level of cortisol, a marker for chronic stress (Fiocco et al., 2006).

Additionally, depression or anxiety could be functional factors, taking place before MCI or dementia and they may correlate with changes in structural areas like amygdala (Liu et al., 2020). Chronic stress in turn can be consequence of the lifestyle, being poverty, the main cause of chronic stress (Fernandez-Blazquez et al., 2021). On the other hand, cortisol secretion is linked to circadian rhythm and a relation between sleeping time, cortisol secretion, and dementia has been recently indicated (Antypa et al., 2021).

CONSCIOUSNESS AND “HIDDEN” STRUCTURES THAT COULD BE INVOLVED BEFORE THE APPEARANCE OF COGNITIVE IMPAIRMENT

In Familial Alzheimer Disease (FAD), consciousness changes have been considered as an early marker of the disease (Aschenbrenner et al., 2020), and claustrum has been proposed to be a brain area controlling consciousness (Crick and Koch, 2003). Claustrum is a “hidden” structure, located below the insula cortex, that can only be visualized when other parts of the cortex are pulled aside. Claustrum dysfunction may precede amyloid accumulation and aggregation in FAD (Goutagny et al., 2013). Additionally, claustrum can establish connections with entorhinal cortex (Kurada et al., 2019) and hippocampal areas (Amaral and Cowan, 1980), which have been related to tau pathology, and cognitive impairment. Thus, we suggest that further studies analyzing the possible role of claustrum in very early stages of AD should be performed, not only on FAD, but also in sporadic Alzheimer’s disease (SAD). If there is a role of claustrum in SAD, a very early functional change in the AD continuum could be related to controlling consciousness (Figure 1A).

REVERSION FROM MILD COGNITIVE IMPAIRMENT TO NORMAL COGNITION

Recently, it has been shown that the likelihood of progression from MCI to dementia is very similar to the reversion from MCI to normal cognition (Sanz-Blasco et al., 2021). Some factors involved in that reversion have been described (Sanz-Blasco et al., 2021), but there are other features that should be analyzed, based on the previous history of the patient. For instance, we have previously commented on this minireview the possible role of chronic stress as a very early risk factor for dementia. Suitably, reversion of chronic stress correlates with reversion to a normal healthy cognition. However, in some cases that stress results in the irreversible atrophy of dentate gyrus neurons (Bai et al., 2012), which remains as a risk signature that could facilitate the future progression to dementia. In addition, morphological (unreversible?) reorganization, in hippocampus, nucleus accumbens, and amygdala has been reported after corticosterone administration (Morales-Medina et al., 2009).

In this way, we would like to comment that new psychological tests to determine changes in social and emotional behavior may be needed to account for possible changes related to the presence of amyloid plaques at specific brain locations at very early times of the continuum. Recently, changes in emotion and generosity in older adults have been reported (Carstensen and Chi, 2021), but little was done in MCI/AD patients, especially at early stages of the disorder.

In humans, social and emotional processing is mainly localized at cerebral neocortex in areas like the orbital frontal cortex. In the first Thal stage, this area already shows an evident amount of amyloid aggregation. Although it has been reported that orbital frontal cortex is involved in emotional enhancement of memory (Kumfor et al., 2013), there are not many studies
looking for possible behavioral changes at those early Thal stages. Curiously, orbital frontal cortex pathology related to AD has been probably more analyzed by examining at tau pathology than amyloid pathology (Tekin et al., 2001). It has also been indicated that damage on the orbitofrontal cortex and the anterior cingulate cortex correlate with behavioral changes, for example dealing with empathy (Avila-Villanueva et al., 2021). Thus, we suggest the possibility of preparing behavioral psychological tests to explore changes at very early timepoints of the continuum. These tests may analyze behavioral changes, expressed by symptoms like agitation, disinhibition, elation, anxiety, or depression (Cajanus et al., 2019), and other features related to subjective wellbeing (van Zonneveld, 1961), similar to EuroQoL-5D (Dolan, 1997). These tests could be a good complement for the previous ones measuring memory changes, cognition, or executive functions, more related to the development of tau pathology, even though during the development of the disease there is an overlapping of both pathologies. Indeed, there such overlapping during the spreading of tau pathology is linked to the fact that tau spreading is favored by the presence of amyloid aggregates (Busche and Hyman, 2020). This tau spreading from the hippocampal area to cerebral cortex is related to an initial memory (and cognitive) impairment, which endpoint could be dementia. Thus, in Figure 1B, we have shown a relation between structural changes (with the presence of amyloid, tau, or both pathologies) with specific functional (behavioral, emotional, memory, or cognitive) changes, occurring at different times of the AD continuum.

CONCLUSION

We propose that further analysis should focus on indicating the features that favor the progression into dementia or those that could be involved in the reversion to a normal cognitive situation (Sanz-Blasco et al., 2021). The knowledge of the later features may facilitate the use of therapeutic tools at very early stages of the AD continuum. The possible reversion will probably need some extent of in-depth knowledge of the patient through the recently proposed precision (personalized) medicine (Hampel et al., 2018, 2020) and also by taking into account the possible correlation between image studies to analyze structural changes with functional failures analyzed by neurological and neuropsychological studies.

AUTHOR CONTRIBUTIONS

All authors were responsible for the conceptualization, reviewing the literature, and critically editing the manuscript, approved the submitted version of the manuscript, and were accountable for the accuracy and integrity of the work.

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