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Review

Novel Development and Prospects in Pathogenesis, Diagnosis, and Therapy of Alzheimer's Disease

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Abstract. Alzheimer's disease (AD) is the most prevalent neurodegenerative disease with cognitive decline and behavioral dysfunction. AD will become a global public health concern due to its increasing prevalence brought on by the severity of global aging. It is critical to understand the pathogenic mechanisms of AD and investigate or pursue a viable therapy strategy in clinic. Amyloid- β (A β) accumulation and abnormally hyperphosphorylated tau protein are the main regulating variables in the pathological phase of AD. And neuroinflammation brought on by activated microglia was found to be one risk factor contributing to changes in A β and tau pathology. It is important to investigate the unique biomarkers of early diagnosis and advanced stage, which may help to elucidate the specific pathological process of AD and provide potential novel therapeutic targets or preventative measures.

Keywords: Alzheimer's disease, amyloid-B, biomarker, neuroinflammation, tau

INTRODUCTION

As the primary cause of dementia, Alzheimer's disease (AD) is the most prevalent neurodegenerative disease for which there is no effective treatment available. Short memory loss is the early clinical symptom of AD. Other features of AD include the cognitive decline and behavioral dysfunction at a moderate level [1]. The hallmark clinical aspect of AD is the progressive impairment of behavioral and cognitive abilities, including memory, mood, emotions, language, attention, and judgement. Nearly 60%–80% of AD patients will eventually lead to dementia [2].

Aging is a risk factor for AD. According to the latest findings in epidemiological research, AD will become a global public health concern due to its increasing prevalence brought on by the severity of global aging. Every five years, the number of new instances of AD in adults over 60 doubles. For those over 60, the prevalence rate is 1%, and for those over 85, it is 40% [3]. In the future, more elderly people will be diagnosed with AD due to the increasing illness risk brought on by the aging population.

In 2020, there were more than 50 million people globally who have AD, up from 35 million people who received a diagnosis in 2010. According to the recent Alzheimer's Disease International report from 2021 [4], 75% of dementia patients worldwide, mostly in low- and middle-income countries, remain undiagnosed due to the limitations of early diagnosis. It is anticipated by 2030, 80 million people will

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be diagnosed with AD, and by 2050, the number will rise to around 150 million. Furthermore, current studies indicate that one third of men and two thirds of women are at risk of developing AD in their lives. It is yet unknown why women have a higher risk of disease than men, as well as the underlying biochemical mechanisms and motivations. AD is currently estimated to cost \$1 trillion globally in medical expenses, which will have a significant negative impact on society's ability to sustain its economy. Therefore, it is critical to understand the pathogenic mechanisms of AD and investigate or pursue a viable therapy strategy in clinic.

PATHOGENESIS

According to differing clinical processes and pathogenic variables, AD is separated into earlyonset family AD (EOAD) and late-onset sporadic AD (LOAD). Less than 5% of cases of AD are familial (fAD), while sporadic AD accounts for 95% of cases (sAD). Most fAD patients are diagnosed before the age of 65 and have a background of mutations in numerous genes, including $A\beta PP$ (amyloid- β precursor protein), PSEN1 (presenilin 1), and PSEN2 [5]. EOAD typically includes a family history of inherited disease. LOAD is primarily sporadic and is associated with the multi-risk factors including type 2 diabetes, traumatic brain injury, stroke and APOE 4 (apolipoprotein 4) gene. The genetic factor of sAD is APOE 4 [6], and sAD is also influenced by sortilinrelated receptor 1, clusterin, complement component receptor 1, CD2AP, EPHA1, and MS4A4/MS4A6E. Additionally, there are other factors that can affect the LOAD, including as depression, lower education, and even dietary changes or environmental alterations [7, 8]. It has been also demonstrated that ApoE 4 is produced by astrocytes, and that ApoE 4 can exacerbate the inflammatory response by further stimulating the production of inflammatory chemokines and cytokines. Microglia activation with lipopolysaccharide (LPS) triggers an inflammatory response in a mouse mode that is associated with APOE 4 and may worsen neurodegeneration [9].

Amyloid-cascade hypothesis

In the pathogenic phase of AD, amyloid- β (A β) accumulation and abnormally hyperphosphorylated tau protein are the primary regulatory factors. The development of amyloid plaques was caused by extracellular A β accumulation [10], and neurofibrillary

tangles were brought on by intracellular tau phosphorylation [11]. And the development of neurofibrillary tangles and amyloid plaques induced the further synapse loss and cognitive dysfunction. In line with the amyloid-cascade theory of A β production [12], sA β PP β , A β , and the intracellular domain of A β PP (AICD) were released from A β PP by β -secretase and γ -secretase cleavage, respectively (Fig. 1). γ secretase [13] consisting of nicastrin, PSEN 1, PEN 2 (presenilin enhancer 2) and APH 1 (anterior pharynx defective 1) as the essential components plays the crucial role in above process.

According to the recent finding on clinical drug trials for AD, γ -secretase enzyme may be a promising therapeutic target for controlling AB level regulation [14]. Therefore, specific γ -secretase inhibitors, modulators that particularly reduce of $A\beta_{42}$ production, or stabilizers that produce longer A β peptides (\geq 42) [15], in combination with other therapies that target hyperphosphorylated tau or neuroinflammation, may be used to effectively slow down the pathological progression and development of AD from early to later stages. Such AB production, which largely includes $A\beta_{1-40}$ and $A\beta_{1-42}$ in above pathological process of amyloidogenic pathway played the major role in the development of the pathological process of AD [16]. A $\beta_{42/40}$ is currently a critical biomarker of AD clinical diagnosis by positron emission tomography (PET)-computed tomography (CT) or cerebrospinal fluid (CSF) checking [17]. Additionally, mutations in the genes for $A\beta PP$, PSEN1, and PSEN2 could lead to or accelerate the accumulation of AB peptides [18], and these alterations were the specific genetic background on fAD etiology.

More studies have recently offered various newly proofs that intracellular A β (iA β) [19], a biomarker or intracellular modulator, may have a great impact on the pathological progression of AD. The production of iAB, the import of extracellular AB, and the clearance of $iA\beta$ [20], however, would help to clarify the mechanism of iAB accumulation and provide more information on the effects of AB toxicity on synapse dysfunction. Our latest study [21] also evidenced that γ -secretase inhibitor regulated iA β toxicity by slowing down endocytosis, which was associated to N-cadherin-CTF, a synaptic cell adhesion molecule. These recent findings may offer novel insight into the molecular mechanism of AD and provide new prospective targets for drug discovery and development. The regulation of synaptic morphology and plasticity was associated with the ABPP [22], which functions as a synaptic adhesion molecule. Other



Fig. 1. In line with the amyloid-cascade theory of A β production, sA β PP β , A β peptides, and the intracellular domain of A β PP (AICD) were released from A β PP by β -secretase and γ -secretase cleavage, respectively. The essential components of γ -secretase include nicastrin, PSEN 1, PEN 2 and APH 1. In the non-amyloidogenic pathway, sA β PP α , P3 peptides, and AICD were produced by α -secretase and γ -secretase cleavage, respectively.

synaptic adhesion molecules, such as cadherin [23], neurexin, neuroligin [24], LRRTM3 [25], and NCAM [26], also have significant effects on AD's pathophysiology. Additionally, a plausible cause of synapse loss or dysfunction could be an interaction between the synaptic adhesion molecules and A β toxicity on the synapses.

These $A\beta$ effects were the main cause of neurodegeneration in familial AD, and $A\beta$ accumulation and aggregation may be responsible for the pathogenic changes in hyperphosphorylated tau protein. Thus, synergistic effect of $A\beta$ and tau should be taken into consideration.

Synergistic effect of $A\beta$ and tau

A β and tau play the crucial roles in the pathological progression of AD and are, in theory, the most significant therapeutic targets for ongoing new medication research in this area. Many options for anti-AB deposition and anti-tau protein hyperphosphorylation were able to exhibit good pharmacological effects in pre-clinical investigations [27, 28] in accordance with the needs of drug design by reducing A β protein deposition or lowering tau protein hyperphosphorylation, respectively. It would also be in keeping with the original purpose of anti-AD drug creation and might dramatically slow the trajectory of cognitive loss in AD animal models [29]. Although the abovementioned candidate drugs could still limit the level of tau protein hyperphosphorylation and reduce AB deposition in the brains of AD patients. Most clinical trials to far have failed because additional clinical investigations [30, 31] have demonstrated that the decline in cognitive function of these patients has not been significantly reversed or modified.

PET molecular neuroimaging was able to detect both total or hyperphosphorylated tau and AB distribution in the brain, it has been proved that the pathological changes of tau protein were independent with AB accumulation [32], both tau and AB might be occurred in distinct "time windows" of AD development. Additionally, neuroimaging data also demonstrated [33] that such changes to $A\beta$ and tau were seen in several targeted regions during the early stages of AD, with tau showing up earlier among these changes. In the early stages of AD, hyperphosphorylated tau is dispersed alone in the entorhinal cortex of the brain without the presence of A β [34]. Such hyperphosphorylated tau detected solely by PET in the particular neurofibrillary tangles in the absence of AB accumulation provided the additional proof that the pathological process of tau protein may be the alternative pathway independent of AB in the development of AD pathology. Furthermore, it is evident that AB accumulation in the late stages of AD could facilitate or enhance tau distribution in the neocortex, promoting AD-related neurodegeneration [35].

As a result, this concept offers the novel insights into the pathological process of sporadic AD, and studying the synergistic interaction between A β and tau will help us clarify more details of the mechanism of AD. Furthermore, tau-mediated neurodegeneration, which is independent with the A β pathway, has an additional impact on the pathological process of AD [33], which may be part of the reason why anti-A β therapies applied solely by only removing A β obviously from clinical AD patients were unable to improve or reverse the cognitive decline. In addition, Phosphokinases CDK 5 [36], GSK 3[37], and ERK2 [38], may also control tau protein hyperphosphorylation in sporadic AD.

Even though it has been established that the pathological changes of tau protein were independent of the accumulation of A β , the relationship between the two remains controversial. Since the molecular mechanism underlying AD pathogenesis is still unclear, more experimental data and research will be required to investigate it in the years to come. These additional investigations will support a theoretical framework for clarifying the pathological mechanisms and signaling pathways associated with AD, as well as present new pharmacological targets and therapeutic approaches for the clinical treatment of AD patients. Due to the synergistic effects of A β and tau discussed above, a combined therapeutic strategy using anti-tau and anti-AB may have an anticipated positive impact on the future clinical care of AD patients.

Neuroinflammation

In addition to the amyloid cascade hypothesis of AD, neuroinflammation brought on by activated microglia was found to be one risk factor contributing to changes in A β and tau pathology [39, 40]. Pathological tau protein can induce the production of IL-1 via exacerbating neuroinflammatory reactions. In hippocampus, tau phosphorylation is decreased by inhibiting adenosine A2A receptors [41]. In the rTg4510 model, LPS may lead to the form of neurofibrillary tangles and hyperphosphorylation of tau [42]. Numerous research on the biomarkers for neuroinflammation have shown that the activation of microglia in such an inflammatory response within the brain led to the release or production of cytokines or chemokines such as IL-3, IL-12, or IL-10, etc. In animal models, stimulation of astrocytes and microglia promotes amyloid accumulation and speeds up inflammatory responses through secretion of cytokines. Research has indicated that there is a greater presence of TNF- α and IL-6 in the brains of individuals suffering from clinical AD. Neurotoxic astrocytes may arise in a neuroinflammatory environment due to the activation of microglia and the

release of IL-1, tumor necrosis factor (TNF- α). Furthermore, these neurotoxic astrocytes are unable to support the development of new synapses [43, 44].

Recent clinical studies [45] have demonstrated that AD patients who have higher amounts of IL-12 in their brains, even in the presence of a significant quantity of AB accumulation concurrently, will not experience severe cognitive impairment. Intriguingly, tau tangles are rarely seen in the brains of AD patients if IL-12 levels are higher. Additionally, raising the levels of IL-12 and IFN- γ may postpone or slow down cognitive loss, but the precise molecular pathways driving the degenerative course of AD are still unknown. The most recent clinical research report also confirmed that IL-3 released from astrocytes in the brain could control the activation of microglia and reprogram neuroinflammation [46], as well as clear the accumulation of extracellular AB plaques and reduce the formation of intracellular tau tangles. However, it is still unknown whether this action of IL-3 could regulate or reverse cognitive decline. This means that the pathogenic changes brought on by these cytokines may dramatically restrict the progression of AD. Furthermore, since the development of synapses is the basis of learning and memory, microglia have the ability to alter or remove existing synapses as well as control synaptic plasticity by remodeling the extracellular matrix. The balance between synapse formation and synapse elimination will be regulated by the effect of neuron-glia communication system [47]. Our latest research [48] shown that hemisynapse formation could be induced on cortical or cerebellar astrocytes following interaction with cortical explant axon by overexpression of the synaptic cell adhesion molecules LRRTM2. This finding may offer a potential approach for treating AD by artificially inducing the newly synapse formation. The process of the neuron-glia communication system also involves the secretion or release of IL-33 and alterations in microglia [49]. Age-related alterations in dendritic spines are all linked to IL-33 and its receptor IL1RL1. The loss of dendritic spines that occurs with aging can be stopped by artificially raising IL-33 levels, which encourages synapses' remodeling and growth. The aforementioned novel biomarker from blood or CSF may be used in the clinical diagnosis or treatment of AD patients.

According to recent breakthroughs in research on AD therapeutic medications, if anti-A β therapy is taken as early as feasible in conjunction with the multi-targeted treatments, it may primarily be successful or delay the onset of clinical symptoms.

As a result, early diagnosis using a combination of precise techniques like magnetic resonance imaging (MRI), PET, biomarker measurement, or cognitive score evaluation becomes increasingly crucial and urgent in order to prevent the destructive progression of AD from the early stages to the late phases.

DIAGNOSIS

The diagnosis of AD is based on the cognitive, functional, and behavioral scores of the patient, medical evaluation by doctors, combined with PET or MRI brain imaging, and analysis of biomarkers such as $A\beta_{42/40}$, p-tau, glia fibrillary acid protein, and neurofilament light from CSF (cerebrospinal fluid) or blood.

Basic brain imaging

The use of MRI [50], which helps to comprehend and observe the changes in brain structure related to AD pathology, is currently one of the key medical examination techniques used in hospital settings for the diagnosis of AD. A prominent clinical hallmark of dementia brought on by AD is the loss of brain tissue, and an MRI will reveal morphological alterations in AD patients, such as the atrophy of specific brain regions. By assessing the distribution of AB or phosphorylated tau in different brain regions from preclinical stage or moderate cognitive impairment to dementia of AD, PET imaging [51] can offer the metabolic changes in AD brain. It's important to elucidate the pathogenic mechanism of neuroinflammation in animal models and AD patients by using PET molecular imaging. The most neuroinflammatory PET imaging targets is mitochondrial 18 kDa translocator protein (TSPO), and the PET tracer ^[18F]flortaucipir was used to detect tau accumulation and neuroinflammation [52, 53].

Novel CSF or blood biomarkers

 $A\beta_{42/40}$ is one important biomarker for the diagnosis of AD patient through PET-CT or CSF examination [17]. The clinical diagnostic level of AD will be raised by using single molecule array (SIMOA) or mass spectrometry as high sensitivity immunoassays to measure novel biomarkers in patient plasma or CSF. Therefore, creating simple, timely, affordable, and accurate diagnostic methods based on blood samples can aid us in promptly screening AD patients as early as feasible and in taking early preventative action to slow down the clinical pathological process of AD patients. Neurofilament light [54, 55] has been quantified from patient CSF or plasma samples and utilized as a general indicator of neurodegenerative disease to represent neuronal loss or injury in numerous countries to date. Another novel blood-based AD biomarker is glia fibrillary acid protein [56], which was linked to the pathological alterations on AB and released during astrocytic activation. New biomarkers from inexpensive blood tests can change the drawback and limitations on CSF investigation caused by the necessity of lumbar puncture, as well as the existing expensive clinical diagnosis of AD patients using PET imaging. It will aid in the development of novel AD therapeutics. Although the patient does not exhibit any signs of cognitive impairment in the preclinical stage of AD, there are nonetheless neuropathological changes in the tau or A β proteins. In the laboratory, it has been reported that several new phosphorylated tau specificities, including p-tau 181, p-tau 217, and p-tau 231 [57-59], have been used by research scientists to assess tau and AB pathological process as the novel blood biomarkers for the potential clinical diagnostic method of AD.

Applications of artificial intelligence

Additionally, due to the recent rapid advancement of computational intelligence techniques, computeraided diagnosis systems for disease using medical artificial intelligence algorithms such as machine learning, deep learning, and convolutional neural network model [60] will provide the most accurate diagnosis and early prevention for AD patients in conjunction with professional physician clinical judgment through the cognitive tests, imaging data, or biomarkers analysis. Using a deep learning algorithm and ¹⁸F-FDG PET imaging [61], it can predict the AD diagnosis. By applying machine learning in conjunction with drawing characteristics and Montreal Cognitive Assessment scores [62], it has been suggested that early identification and diagnosis of cognitive deficiencies will assist enable preventative dementia intervention in clinic. The Digital Neuro Signature from Altoida was used to assess neurocognitive function on an individual basis [63] and was approved as a breakthrough device by the FDA in August 2021. Therefore, in the next years, a basic clinical early evaluation system may screen cognitive capacities of high-risk groups early on by utilizing deep learning and machine learning algorithms

in addition to the Mini-Mental State Examination, drawing clock, AD8 test, and other tests. This will play a certain auxiliary role in the early prevention of AD.

THERAPIES

AD will unavoidably place significant societal burdens on people's lives all around the world before the development of efficient treatment medications or procedures. It is very advantageous economically and socially to investigate the pathogenesis of the illness and identify potential new treatment targets or preventative measures (Table 1) in order to lessen clinical symptoms and enhance the quality of life for AD patients [64]. The FDA has approved the use of the therapeutic medications Galantamine, Donepezil, Rivastigmine (AChE inhibitors), and Memantine (NMDA receptor antagonists) to treat the clinical symptoms of AD patients, although none of them reverse or improve cognitive impairment [65]. Additionally, the FDA approved one A β immunotherapy using Aducanumab from Biogen in June 2021, while this choice is still causing debate among scientists [66, 67]. In January 2023, Lecanemab was approved by FDA for the treatment of AD through the accelerated approval pathway. It was obtained traditional approval from FDA in July 2023 [68-70] (Table 2).

Even though they performed well for AD animal models in the preclinical research stage, the majority of medication candidates undergoing clinical trials are unable to slow down or reverse the clinical symptoms of cognitive loss in individuals with AD. Therefore, novel regenerative therapies using stem cells, such as mesenchymal stem cell or human induced pluripotent stem cell (hiPSC) application, or artificial intelligence technologies used in new drug discovery and development would also have a potent chance or possibility in the clinical treatment or prevention strategies of AD in the future, in addition to the traditional drug candidates in various stages of preclinical or clinical trials of AD.

With the aid of hiPSC technology, the pathological basis of AD has been studied as well as potential treatment options. After using cell culture techniques, hiPSC acquired from AD patients will be developed into neural progenitor cells and neurons [71]. Combining this cell model with gene-editing techniques like CRSIPR-Cas9 offers the ability to quickly assess the properties of possible medications while shedding light on the cellular and molecular illness causes, which can lower the cost and shorten the development cycle for novel therapies. The human iPSC model was also employed to investigate the pathological features of AD in depth and identify potential therapeutic targets for AD medication development in our lab [72]. In addition, human iPSC technology may be employed for individualized treatment like cell transplantation, based on neuro-regeneration therapeutic for AD, to stimulate neuronal and synaptic regener-

10p 5 list for the CADRO target in an phases of chinical trans (<i>until 1 January 2025</i>)								
CADRO target	Number of agents (all clinical phase)	Percentage of total agents	Agents (phase 1)	Agents (phase 2)	Agents (phase 3)			
Transmitter/Receptor	29	15.5%	6	12	11			
Inflammation	25	13.4%	6	17	2			
Amyloid	25	13.4%	7	11	7			
Synapse plasticity	21	11.2%	1	14	6			
Tau	14	7.5%	4	8	3			

 Table 1

 Top 5 list for the CADRO target in all phases of clinical trials (until 1 January 2023)

CADRO, Common Alzheimer's Disease Research Ontology; There were 187 total agents in all phases of clinical trials until 1 January 2023.

Table 2 New representative drugs for Alzheimer's disease								
Agent	Name	Company	Mechanism of action	Therapeutic purpose	Status			
Aducanumab	Aduhelm	Biogen	Monoclonal antibody	DMT, biologic	Accelerated approval from FDA, June 2021			
Donanemab	LY3002813	Eli Lilly	Monoclonal antibody	DMT, biologic	Applied for full FDA approval, <i>June 2023</i>			
Lecanemab	Leqembi	Eisai/Biogen	Monoclonal antibody	DMT, biologic	Traditional approval from FDA, <i>July 2023</i>			

DMT, disease modifying therapy.

ation in cognition of AD [73], which would provide innovative therapeutic strategy for AD patients.

Additionally, the objective of nonpharmacological treatment is to enhance the daily functioning of AD patients by influencing cognitive function through art therapy or memory training to lessen depressive symptoms and correct sleeping abnormalities [74]. Word games, music therapy, and other activities have been proven in certain studies to be beneficial for training cognitive function. As a result, non-drug therapy can be utilized as a supplement to other methods of treating clinical AD patients. More practical answers for AD prevention techniques will also be provided by outcomes from linked studies on diet, genetics, or environmental variables. Antioxidant mediators, including vitamin E, have been shown to be beneficial at delaying the progression of AD [75, 76].

CONCLUSION

Therefore, by investigating the unique biomarkers of early diagnosis and advanced stage, it will elucidate the specific pathological process of AD and offer fresh suggestions for clinical treatment techniques, along with significant and valuable scientific research value.

AUTHOR CONTRIBUTION

Zenghui Teng (Conceptualization; Data curation; Writing – original draft; Writing – review & editing).

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CONFLICT OF INTEREST

The author has no conflict of interest to report.

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