Alzheimer’s Disease Silent Killer of Memory: A Review on Pathological Mechanisms

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Abstract

Alzheimer’s Disease (AD) is a Central Nervous System (CNS) related neurological disorder characterized by impairment of memory and finally turbulence in learning, reasoning, planning, language and perception. AD is a complex disease caused by an accretion of β-Amyloid (AB) plaques and neurofibrillary tangles formed by Tau amyloid fibrils linked with loss of synaptic plasticity and neurodegeneration. Apart from several other environmental factors, the rise in Oxidative Stress (OS), failure of mitochondria and Golgi plays a crucial role in neuronal dysfunctioning leading to neurodegeneration. However, oxygen is necessary for life and any imbalance between metabolism and excess Reactive Oxygen Species (ROS) generation leads to the origin of disorders such as Alzheimer’s disease. The ROS induced Mitochondrial (Mt) dysfunctions and excitotoxicity through over Ca²⁺ influx are two hallmarks behind neurodegeneration in AD. Still, there is no medicament presently available which can prevent further progression or decrease the symptoms of this disease. Some anti-oxidants were used but they were not effective due to their bioavailability problem into brain. The absence of effective therapeutic drugs and increased severity of AD day by day is a big challenge for against researchers. Therefore this review basically focused to explore the underlying pathological mechanism behind neuronal dysfunctioning and the various pathways involved in the pathogenesis of AD.

Keywords: Alzheimer’s disease; Beta-amyloid; Caspases; Neurofibrillary tangles; Oxidative stress; Tau protein

Abbreviations

AD - Alzheimer’s Disease
AB - Amyloid Beta
NFTs - Neurofibrillary Tangles
CASP - Caspases
TNF-α - Tumor Necrosis Factor
APP - Amyloid Precursor Protein
ROS - Reactive Oxygen Species
IL - Interleukins

Introduction

Alzheimer’s Disease (AD) is a slowly progressive neurological disorder characterized by impairment of memory and ultimately disturbances in learning, reasoning, planning, language, and perception [1]. It is also known as Senile Dementia of the Alzheimer Type (SDAT). Globally more than 45 million peoples are living with dementia, most suffering from AD. The prevalence of dementia will reach 75.6 million in 2030 and 135.5 million in 2050, of which 6.46% are from USA [2]. Many scientists believe that AD results from an increased oxidative stress due to the accumulation of a specific protein (beta-amyloid protein) in the brain that leads to neuronal cell death [3].

AD has been named after the German psychiatrist Alois Alzheimer who first described this disease in 1901 in Auguste D, a 50-year-old lady. Depending on its time of onset AD may be; Early-onset AD (usually affecting people aged 30 to 60 years) and Late-onset AD (this generally disturbs people over age 65 years) [4]. In AD patient may face problems in recognizing family and friends due to excessive neurodegeneration or neurotransmitter imbalance. The common symptoms in AD include alteration in mood, behavioral, hallucinations, and delirium. It usually starts slowly, silently and gets worse over time. With the advancement of disease physiological symptoms also alter like problems with language, alteration of mood, loss of inspiration and behavioral issues [5]. As a person’s condition declines, she or he often withdraws from family and society, gradually body functions are lost leading to neuronal loss and death.

In addition, the accumulation of neurofibrillary tangles, free radical damage, low antioxidant enzymes level (Superoxide Dismutase (SOD) and catalase), mitochondrial failure have been observed in AD patients [6]. The increased oxidative damage to lipids and proteins with reduced activity of glutathione and other antioxidant enzyme are...
more localized to the neuronal junctions and correlate with the severity of the disease, suggesting to play key role in AD-related neurodegenerative disorders [7]. The postmortem reports suggest that oxidative stress may be one of the prominent factors that is responsible for the initiation and development of AD. The exact pathophysiology of AD is still inadequately understood but mitochondria dysfunction and reduced level of pericyle are the progressive pathological factors [8].

Available evidence indicates that fragmentation of Golgi body increase Amyloid Precursor Protein (APP) trafficking and increase Amyloid Beta (Aβ) accumulation [9]. Joshi, et al., has suggested that the accumulation of Aβ peptides which leads to fragmentation of Golgi complex [10]. Nearly 70% risks of factors are of genetic background due to the involvement of various genes [11]. Additional risk factors include: person suffering from depression, accidental brain injury or hypertension. The major hallmark includes deposition of protein plaques (Tau proteins) and neurofibrillary tangles in the brain [12].

Nowadays there is no specific treatment available to prevent AD but the non-pharmacological measures like physical exercise as well as reducing body fat may decrease the risk of AD. Some of the antioxidants and vitamins from a long time are being under pre-clinical evaluation but these supplements cannot cure AD entirely or reverse its progression, though for the moment reduce symptoms [13]. Several drug therapies are available but they provide symptomatic relief to the patient suffering from this disease. The patients suffering from this disease become dependant on others for their daily life activities often increase a burden on the caregiver. Exercise programs are advantageous in the daily life and potentially improve outcomes. Antipsychotic agents are commonly used for the management of behavioral or psychotic problems due to dementia but not usually recommended due to their stumpy brain penetration and less therapeutic potential. Hence further studies are required to explore the hidden targets to achieve better drug therapy to secure life [14].

Pathophysiological Mechanisms in Alzheimer’s Disease

Role of Amyloid Beta (Aβ) in neurodegeneration

Amyloid Beta (Aβ) are peptides of 40-42 amino acids which produce various pathological changes in AD [15]. The accumulation of the proteins like β-amyloid is responsible for structural changes in AD [16]. The peptides are formed by proteolytic cleavage of the large transmembrane Amyloid Precursor Protein (APP) by β and γ secretases. These Aβ protein molecules can combine to form oligomeric bundles which are soluble and exist in several forms [17]. It is hypothesized that there are certain misfolded oligomers (known as “seeds”) responsible for inducing or misfolding of other β-amyloid molecules by a series of a chain reaction. This results in accumulation of β-amyloid and produces a toxic effect in the nerve cells [18]. The other type of protein present in AD is tau proteins which also exist in misfolded oligomers. The number of evidence support that misfolded Aβ can induce misfolding of tau protein. The actual physiological function of Aβ is not fully clear but some animal study reports have shown that the lack of Aβ proteins does not lead to loss of any physiological function [19]. There are several potential actions have been exposed for Aβ, including activation of kinase family of enzymes, defense against oxidative stress, lowering or regulation of cholesterol transport, and anti-microbial activity (mainly related to Aβ’s pro-inflammatory activity). The lymphatic system is responsible for clearance of β-Amyloid proteins from the mammalian brain particularly but the pace of elimination is significantly increased during the sleeping time [20]. Aβ proteins are by the sequential cleavage of the Amyloid Precursor Protein (APP) through the action of β and γ secretases respectively [21]. A number of isoforms (36-43 amino acid residues in length) are generated by the action of γ secretases on the C-terminal end of the Aβ peptide. The Aβ40 and Aβ42 are the most common isoform, produced by cleavage in the endoplasmic reticulum and trans-Golgi network respectively [22]. The Aβ40 is the more common form than Aβ42 but Aβ42 is fibrillogenic and therefore responsible for the disease states [23]. The increased production of Aβ42 has been noted during the mutations in APP responsible for early-onset Alzheimer’s which suggested the involvement of modulation of the activity of β and γ secretases as a possible therapy for Alzheimer’s disease.

In 2010 it was noted that the structural sequence on the oligomeric state of amyloid beta is still sparse. Low-temperature and low-salt conditions allowed isolating pentameric disc-shaped oligomers devoid of beta structure [24]. The amyloid beta after accumulation cause loss of neuronal functioning by inhibiting mitochondrial function, increase generation of ROS, further cause’s lipid peroxidation and the generation of a toxic aldehyde called 4-hydroxynonenal which in turn decreases ATPase’s activity, glucose transport and consequently neuronal loss occur. The β amyloid increases glutamate-dependent depolarization of the synaptic membrane which increases calcium influx dependent mitochondrial impairment [25]. Amyloid hypothesis stated that the fundamental cause of AD is extracellular Amyloid Beta (Aβ) deposits. This postulate has been evidenced by the position of the gene that regulates Amyloid Precursor Protein (APP) on chromosome 21, implying that individuals with Down’s syndrome develop typical Alzheimer neuropathology because they produce too much Aβ life-long [26]. Another fact that supports the above-mentioned postulate is that the people with trisomy 21 almost universally exhibit AD by 40 years of age [27]. Also, a specific form of Apolipoproteins (ApoE4) is the strongest known genetic risk factor for the late onset of AD [28]. Further, in the transgenic mice, it has been seen that those mice which express a mutant type of the human APP gene are more prone toward the development of fibrillary amyloid plaques and have more chances of AD. An experimental vaccine was used in early human trails to find out its effect for clearing the amyloid plaque but it was only effective for the clearance of plaque without significant effect on dementia. Researchers have been led to suspect non-plaque Aβ oligomers as the primary pathogenic form of Aβ. These noxious protein oligomers are also known to as Amyloid-Derived Diffusible Ligands (ADDLs). These oligomers easily bind to a cell membrane receptor of neurons and modify the structure of the synapse thereby distracting neuronal communication. A single receptor for Aβ protein oligomers may behave like prion protein, i.e. the same proteins which are associated with disease and the related human situation like Creutzfeldt-Jakob [29] (Figure 1).

Golgi body and Amyloid beta protein: Available experimental reports suggest that the inhibitors of plasminogen activator inhibitor-1 decrease the plasma and brain Aβ levels in transgenic animals. Amyloid targeting immunotherapy can be done by bapineuzumab, gantenerumab etc. These monoclonal humanized antibodies mainly act on the CNS reduce Tau proteins level either by inhibiting their
phosphorylation or clearance mechanism [30]. Exogenous antioxidants are being used like flavonoids, carotenoids and ubiquitous antioxidants show a powerful neuroprotective effect but have some bioavailability problems. The previous finding has reported that rutin, lutein, and curcumin also attenuates Aβ induced inflammation and reasonably inhibits β-secretase and AChE [31]. Normally the Aβ are matured and released by Golgi body but excessive accumulation or tauopathy cause fragmentation of this Golgi assembly that leads to loss of functioning [10].

Role of Tau protein in Alzheimer’s disease

AD also has a concern with abnormal aggregation of tau protein in the neurons and therefore it is also considered as tauopathy. Normally these tau proteins are present in neurons which helps to stabilize the microtubule during phosphorylation and therefore known as microtubule coupled protein [32]. In AD, a tau protein undergoes chemical alterations, become hyperphosphorylated and starts to form a couple with other and thus appear like Neurofibrillary Tangles (NFTs). The presence of NFTs disrupts normal synaptic plasticity, which would lead to a diminution in neurotransmission, death of neurons bearing tangle and dementia [33].

The tau hypothesis, has proposed that any structural and functional tau protein abnormalities initiate disease cascade or disease like condition. Once the hyperphosphorylated tau accumulates inside nerve cell bodies, they cause disintegration of microtubules, destroy the cell’s cytoskeleton that collapses the neuron transport system [34]. This may lead to malfunction of neurochemical communication among neurons that can cause neuronal loss or death [35]. Thus, according to Tau hypothesis it is believed that the abnormal aggregation of the tau protein is the cause of AD. Neurofibrillary tangles are thought to develop when an unknown trigger leads to the dissociation of tau from microtubules. Some studies revealed that hyperphosphorylated tau accumulates as tangles even when tau levels are decreased by transgenic suppression. This might be due to the action of tangles as a sink for tau intermediates that are predicted to have formed as part of the pathogenic process [33]. However, tangle formation is not associated directly with cognitive decline, which suggests that tau intermediates are the neurotoxic species; similar intermediates composed of other proteins such as Huntington or Aβ protein have been suggested to be neurotoxic elements in other diseases [36].

Tau-base targeted strategies prevent Alzheimer’s disease: Tau protein accumulation due to hyperphosphorylation loses microtubule binding and stabilizing role, contributing to neuronal deterioration [37]. Targeting tau phosphorylation is in basically needed to focus on Glycogen Synthase Kinase-3 (GSK3), one of the primary enzymes concerned in tau phosphorylation. A temperate improvement in mood (particularly apathy) and social interaction has been observed with cholinesterase inhibitors treatment [38]. Recent studies revealed that caffeine shows an inhibitory effect on Phosphodiesterase (PDE) enzyme and it also inhibits GSK3b. Microtubule stabilization criteria can be adopted in which the administration of paclitaxel (i.e. microtubule stabilizer) to tau-transgenic mice significantly improves fast axonal transport, microtubule density, and motor function. Epothilone D, is a potent inhibitor of active transporter which is also used as microtubule stabilizer is known for its good blood-brain barrier elimination [39]. Rojo, et al., has suggested that the lansoprazole and astemizole (both benzimidazole derivatives) bind with tau protein and in reducing the formation of neurofibrillary tangles. Enhancing tau degradation helps to augment the degradation of polymerized tau, thus declining its toxicity. Another, heat shock protein 90 (Hsp 90), also help to the denatured proteins, seems to play a role in preventing tau degradation [40]. Tau immunotherapy is successful for passive immunization with monoclonal antibodies decreases the phosphorylated tau molecules to reduce disease instances [41] (Figures 2&3).
Role of Pericyte in Alzheimer’s disease: Pericytes are individually situated in the neurovascular unit between endothelial cells of brain capillaries, astrocytes and neurons. Pericytes are cells in the Blood-Brain Barrier (BBB) which elevates the amyloid β-peptide (Aβ), cause tau pathology and neuronal loss that leads to progressive cognitive decline and dementia [42]. Most of the studies have shown that pericytes regulate key neurovascular functions including BBB formation and maintenance, regulation of capillary blood flow and clearance of toxic cellular by products necessary for normal function formation and maintenance, regulation of capillary blood flow and clearance of toxic cellular by products necessary for normal function [43]. Oxidative damage caused by these negative charge species can further target intracellular structures decrease mitochondrial metabolism which act as causative factors of abnormal function and cause cell death [44]. In the cells, mitochondria produce ATP, which is essential for the neurons survival and functional impairment results in neurodegeneration. Interestingly, vascular hypoperfusion is responsible for the dysfunction of mitochondria in AD with subsequent RNA oxidation, lipid per-oxidation or mitochondrial DNA deletion [45]. Mitochondria are the major site involved in the generation of free radicals like O_2− and H_2O_2 respectively. It is well reported that increases in mitochondrial DNA in the cell soma of AD susceptible neurons can itself cause increased oxidative potential [46].

Increased oxidative stress and low cellular defense directly move the cells toward decay which can be reversed by enhancing antioxidant levels, such as glutathione. The detrimental effects of oxidative stress cause structural damage, cellular apoptosis which finally leads to necrosis. Production of ROS is a particularly destructive aspect of the cellular growth [47]. Some of the less reactive species (such as superoxide) can be converted into more destructive radical species by redox reactions with transition metals or other redox cycling compounds (including quinones) that can cause extensive cellular damage.

Similarly, the long-term effects are caused by damage to DNA which may be induced by ionizing radiation that is similar to oxidative stress and these lesions are mostly implicated in aging and cancer [48]. Some time the high-intensity radiation can cause single-base dents like 8-oxoguanine and thymine glycol and their biological effects have been studied extensively. Most of the oxygen-derived species responsible for oxidative stress cause necrosis are formed during normal cellular aerobic metabolism. The antioxidants are capable of destroying most of these oxygen derived species and any damage to cells is continually repaired. But under some harmful, conditions these species can cause necrosis leads to ATP depletion and finally delay the regeneration of the cells. Xanthine oxidase, NADPH oxidases, and cytochromes P450 are some of the other enzymes which are capable of producing superoxides [49]. Thus, to improve the cellular survival a balance must be necessary between production and consumption of free radical species. Disrupted mitochondrial respiratory ATP synthesis caused by oxidative stress leads to neurodegeneration followed by AD, Parkinson’s disease and other neurodegenerative diseases [50].

Hypoxia exercise induces overproduction of (ROS) and RNS and alters redox balance. Infact, both acute and long-term hypoxic exposures have been shown to enhance the oxidative stress whereas increased oxidative stress has been observed in hypobaric as well as normobaric hypoxia (i.e., simulated altitude). Debevec, et al., has indicated that hypobaric hypoxia might induce higher oxidative stress levels than normobaric hypoxia [51] (Figure 5).
Role of Caspase in Alzheimer’s Disease

Caspases also called cysteine-aspartic proteases are consist of a large number of cysteine proteases that play crucial roles in programmed cell death and initiate inflammatory responses [52]. Caspases are also termed “executioner” molecules because of their roles in the cellular signaling. Some caspases are reported to have their useful function in the immune response regulation because they help in the maturation of lymphocytes. Stoppage of apoptosis can contribute to tumor progression and autoimmune diseases that can initiate unwanted apoptosis commonly occur with ischemia or Alzheimer’s disease [53]. There are two types of apoptotic caspases: initiator (apical) caspases and effector (executioner) caspases. Initiator caspases (e.g., CASP2, CASP8, CASP9, and CASP10) cleave inactive pro-forms of effector caspases, thereby activating them. Effector caspases (e.g., CASP3, CASP6 and CASP7) in turn cleave other protein substrates within the cell, to trigger the apoptotic process [54] (Figure 6).

It is well reported in various studies that the Amyloid Precursor Protein (APP) is a substrate for caspase-3-mediated cleavage, which may contribute to $\beta$ formation, synaptic loss, and the behavioral changes associated with AD. The principle hypothesis supporting this study was obtained using a site-directed Caspases-cleavage antibody to APP. In consideration to this report lot of studies have put forward and been able to demonstrate the caspases cleavage of fodrin in the AD brain as well as the activation of specific initiator and executioner caspases including caspase-3, 6, 8, and 9 [55] (Figure 7).

![Figure 5: Effect of oxidative stress and mitochondrial failure in AD.](image1)

![Figure 6: Process of activation of Executioner caspases.](image2)

![Figure 7: Role of Caspases in Alzheimer’s disease.](image3)
Lastly, these studies provide lot of information which supports the activation of apoptotic pathways during the course of the disease progression in AD. Moreover, the role of caspases in AD is too much wider because the production of Aβ is also associated with their activation. Hence the number of studies suggest that caspases may be showing their proximal role in the disease mechanisms underlying AD including promoting Aβ formation as well as linking plaques to NFTs [56] (Figure 8).

Role of sleep changes in Alzheimer’s disease

Alzheimer’s disease (AD) is a chronic neurodegenerative disorder that causes degeneration of CNS neurons thereby debilitating symptoms like loss of memory, interrupted speech and inability to carry out daily activities [57]. In AD the excessive loss of neuronal function leads to decrease mental abilities which may also disrupt the sleep/wake cycle, responsible for sleep problems, nighttime wandering, and agitation. In AD sleep the percentage of sleep disruption depends upon the severity of disease like those which are at early stages of AD may suffer more than usual [58]. As the disease is at initiation level, patients may begin to sleep during the day and awaken frequently throughout the night. Patients with more advanced in AD they sleep very rarely for long time periods. This also changes daily alterations in body temperature, sleep, wakefulness, and metabolism may also be sometimes disrupted in older adults.

Several reports have evidenced that sleep disruptions may be worse in patients with AD and due to this patient may lose the ability to stay asleep or keep alert during AD progression [59]. Sleep problems may also increase the incidence of agitation among AD patient according to the results provided by various reports [60]. Similarly, a term is used for this type of agitated behavior in AD that is “sundowning” mostly occurs in the evening schedule in various patients. This increased agitation behavior may be explained by sleepiness or may be more troublesome to caregivers during evening time [61]. But overall this creates a sleep apnea result in severe stress for caregivers and among AD patients which are on care therapy in nursing homes instead of at home. At initial stage this Sundown Syndrome is associated with increased stress, mental confusion and restlessness in patients due to which they are more prone to dementia that is commonly associated with AD [62].

The non-pharmacological factors which control the progression of AD are: firstly to make consistent sleeping, secondly change stressful lifestyle and lastly to do physical exercise. Physical activity is the best treatment for Alzheimer’s, which also encourage patient for night sleep. Decrease the use of CNS stimulants like caffeine at night if a night’s sleep is needed. Some evidence also supports the exogenous use of melatonin in combination with other antioxidants like lycopene or vitamin E can be used to induce sleep and preventing neurodegeneration [63]. Other pharmacological agents which are useful for the treatment of sleep deprivation induced or neurodegeneration induced AD are: Acetylcholinesterase inhibitor like donepezil which slows down the cognitive decline in some patients but its major adverse effect is it can cause nighttime stimulation and is reported to promote nightmares in AD patients. Others like atypical antipsychotics and selective serotonin reuptake inhibitors are proved to be useful therapeutic agents [64].

Conclusion

AD is extremely complicated neurodegenerative disorder occurs due to the accumulation and overproductions of β-amyloid, tau formation by stressful lifestyle but the exact mechanism is still unknown. There are a number of molecular targets described which reflect the pathophysiology of AD but neurodegeneration have been speculated to occur through number of other factors also including environmental, genetic predisposition but oxidative stress and free radical generation are a prominent one. At present for the treatment of AD number of drugs like antioxidants and other pharmacological agents has been used but all provide symptomatic relief. So, there is a need for treatments which reduces the β-amyloid plaque formation or increases their elimination from CNS thereby reducing the oxidative stress. In this review, we have combined a number of molecular pathways involved in the pathogenesis of AD. Therefore either the inhibition of these pathways or the use of their antagonist may divert or decreases the progress of neurodegeneration in AD.

References


