Getting to the Heart of Alzheimer Disease

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Abstract: In a somewhat narrow diagnostic lens, Alzheimer disease (AD) has been considered a brain-specific disease characterized by the presence of Aβ (β-amyloid) plaques and tau neural fibrillary tangles and neural inflammation; these pathologies lead to neuronal death and consequently clinical symptoms, such as memory loss, confusion, and impaired cognitive function. However, for decades, researchers have noticed a link between various cardiovascular abnormalities and AD—such as heart failure, coronary artery disease, atrial fibrillation, and vasculopathy. A considerable volume of work has pointed at this head to heart connection, focusing mainly on associations between cerebral hypoperfusion and neuronal degradation. However, new evidence of a possible systemic or metastatic profile to AD calls for further analysis of this connection. Aβ aggregations—biochemically and structurally akin to those found in the typical AD pathology—are now known to be present in the hearts of individuals with idiopathic dilated cardiomyopathy, as well as the hearts of patients with AD. These findings suggest a potential systemic profile of proteinopathies and a new hypothesis for the link between peripheral and central symptoms of heart failure and AD. Herein, we provide an overview of the cardiovascular links to Alzheimer disease. (Circ Res. 2019;124:142-149. DOI: 10.1161/CIRCRESAHA.118.313563.)

Key Words: Alzheimer disease ■ atrial fibrillation ■ heart failure ■ inflammation ■ reactive oxygen species

Alzheimer disease (AD) is a progressive neurodegenerative disorder which accounts for about 70% of all dementia cases1–3 and is projected to affect 13 million individuals in the United States by 2050.4 It is characterized by the formation of senile plaques composed of aggregated Aβ (β-amyloid) fibers and neurofibrillary tangles formed by hyperphosphorylated tau protein, with associated neuronal inflammation, oxidative stress, and widespread degeneration of neurons. These physiological disruptions are accompanied by physical symptoms: memory loss, impaired cognitive function, personality and judgment disorder, speech abnormalities, and apraxia.1

A similar epidemiological profile characterizes heart failure (HF), a widespread public health problem, affecting 5.8 million individuals in the United States alone and nearly 23 million worldwide.4 Obesity, sex, history of diabetes mellitus, and especially, age are risk factors for both HF and AD.1,5–7 Genetic factors, such as the presence of the Apo E4 (apolipoprotein Eε) allele and variants in the presenilin 1 (PSEN1) and presenilin 2 (PSEN2) genes are associated with the development of AD8–10 and the same, as well as novel genetic variants of the PSEN genes, are associated with dilated cardiomyopathy.11,12 Despite those similarities, much of the current understanding is centralized around the pathogenesis of AD via vascular factors and has failed to be extended to cardiac contributions. Vascular factors, such as macro and microinfarcts, white matter hyperintensities (WMH), atherosclerosis, and hypertension that lead to decreased cerebral blood flow (CBF) before Aβ deposition would accelerate AD progression.13

Here, we reviewed the current knowledge on the vascular component and explore the new head to heart connection that could help define cardiac abnormalities occurring in concert with the development of AD.

Vascular Role in AD

For years, the role of the vasculature in the progression of AD has prevailed as a main comorbidity contributing to AD. Vascular cognitive impairment remains a common risk factor of dementia as over half of all patients with vascular cognitive impairment will advance to dementia.14,15 Thus, when attempting to explain the pathophysiology of AD, a vascular hypothesis arose in opposition to the amyloid cascade hypothesis.16 The amyloid cascade hypothesis articulates that the pathology behind AD is driven by the deposition of the Aβ peptide in the brain.17 However, supporters of the vascular hypothesis propose that AD is a vascular disorder, in which the pathology is induced via cerebral microvascular abnormalities.18 In 1993, it was discovered that cerebral microvascular abnormalities lead to a decrease in CBF, glucose metabolism, and oxygen consumption in AD patients.19 de la Torre and Mussivand19 established an inversely proportional relationship between these vascular abnormality symptoms and the severity of AD. In this study, they noted multiple different risk factors, such as aging,
indicating a decrease in CBF, and consequently, a decrease in pulse pressure in AD patients, further supporting the hypothesis that hypoperfusion leads to brain hypoperfusion and dementia of AD. The same study demonstrated that cerebral blood flow (CBF) to be 20% lower in the AD group compared with the non-AD group, suggesting a decrease in cerebral perfusion. A decrease in brain perfusion is a direct contributor to the pathogenesis of AD.20–22 Roher et al22 observed total CBF (and thus oxygen) to maintain proper function.20 It has been proposed that hypoxia and ischemia resulting from cerebral hypoperfusion is a direct contributor to the pathogenesis and development of AD.20–22 Roher et al22 observed total CBF to be 20% lower in the AD group compared with the non-demented control group, suggesting an association between brain hypoperfusion and dementia of AD. The same study found a decrease in pulse pressure in AD patients, further indicating a decrease in CBF, and consequently, a decrease in cognitive measures.22 Nishimura et al23 more specifically observed decreased CBF to the frontal lobe in correlation to both reduced cognitive function and progression of AD. The reduced cerebral perfusion causes a metabolic energy crisis because of the reduced oxygen exposure. This hypoxic state induces oxidative stress and acidosis eventually culminating in neuronal degradation.24,25

Oxidative Stress in AD

Contrary to popular belief, it has been discussed that a balanced reactive oxygen species (ROS) response actually promotes tissue repair via disinfection of existing tissue and stimulation of healthy tissue turnover.26 However, the imbalanced metabolism of oxygen induced by this hypoxic state leads to excess ROS, resulting in deleterious effects.27 Furthermore, the mitochondria of the vascular wall cells have been identified as the primary target of oxidative stress before development and progression of AD.28 However, it is believed that the accumulation of ROS and thereby oxidative stress, on neural tissue is a result of hypoperfusion in combination with Aβ proteotoxicity.29 An increase in Aβ production has been linked to the progression of oxidative stress which can induce mitochondrial dysfunction. Work by Matsuoka et al20 using transgenic mice carrying mutant APP (amyloid precursor protein) and PSEN1 illustrates the key role of oxidative stress in the AD model. Additionally, 3-nitrotyrosine (protein oxidative stress marker) and 4-hydroxy-2-noneal (lipid oxidative stress marker) have been found to be increased with progressing levels of fibrillar Aβ.30 Although most Aβ aggregations are found in extracellular regions, some collections of Aβ have been found in the mitochondria31,32 of individuals with AD. This phenomenon may diminish mitochondrial function, specifically respiration and thereby increase the levels of ROS inside and potentially outside of the cell. Oxidative stress not only can induce further Aβ production but also triggers the production of tau protein, another critical component to the pathology of this disease.31 Furthermore, it has been demonstrated that individuals with AD have decreased antioxidant levels, allowing higher levels of ROS to accumulate in the local environment.32 This evidence, therefore, suggests a cyclic effect between Aβ aggregation and oxidative stress, driving the progression of AD and its physical symptoms.

Inflammation Contributes to Vascular Dysfunction and Neurodegeneration in AD

Another key aspect of the pathogenesis of AD as a result of reduced blood flow is inflammation. Changes in the vasculature either as a result of AD or part of an overall vasculopathy are associated with the release of multiple inflammatory factors, with cerebral microvessels secreting higher levels of TNF (tumor necrosis factor)-α, IL (interleukin)-1β, IL-6, and leukocyte adhesion molecules than non-AD controls.33 However, despite the vast amount of literature focused on neural inflammation and its role in AD, no clear evidence has illuminated whether inflammation is a contributing factor towards the cause of AD, part of its pathology, or a secondary phenomenon.34 However, it is widely accepted that the negative consequences of inflammation contribute towards the progression of AD, including neurodegeneration, and thereby diminished cognitive function. It is also understood that microglia, the resident macrophage in the brain, play a key role in the activation of the inflammatory response. Studies have illustrated that microglia are increased in individuals with AD and in transgenic mouse models. Although they exhibit heterogeneous phenotypes, it is understood that they are involved in Aβ maintenance.35 Studies in vitro suggest that cytokine signaling and secretion from microglia can greatly impact the cerebral microenvironment and function of neurons. These cytokines—specifically IL-1β, IL-6, TNF-α, IFN (interferon)-γ—and various chemokines can strengthen the inflammatory response.35 More recently, cytokine signaling in AD mouse models can have significant effects on amyloidosis, neurodegeneration, and cognition,36 potentially disrupting the blood-brain barrier (BBB) in neurodegenerative disorders.37 In AD pathology, the integrity of the BBB is reduced37,38 as well. Whether as a result of the chronic cerebral inflammation or part of a larger systemic pathology, this defective BBB is believed to play a role in the pathogenesis of AD.

Structural Consequences of Hypoperfusion in the AD Brain

Furthermore, this same cerebral hypoperfusion has been found to break down the neurovascular unit, enabling progression...
into neurodegenerative disorders like AD.39 The neurovascular unit encompasses many different types of brain cells (endothelial, pericytes, and vascular smooth muscle cells), which in turn control the BBB.39 Breakdown of the neurovascular unit would result in a dysfunctional BBB. Many recent studies have now correlated BBB dysfunction with the accrual of vasculotoxic molecules and hypoxia resulting from a decrease in CBF.39

**Amyloid Directly Contributes to Cerebral Angiopathy**

Changes in peripheral perfusion also have the potential to induce cerebral amyloid angiopathy—fibrillary amyloid deposition in small cerebral vessels. However, the exact mechanism whereby cerebral amyloid angiopathy contributes toward the pathogenesis of AD remains unknown. It has been reported that cerebral amyloid angiopathy can affect cell viability, induce apoptosis or oxidative stress, and trigger an inflammatory response.40 Furthermore, cerebral amyloid angiopathy can create vascular dysfunction through hemorrhagic complications or blocking blood flow (resulting in cerebral ischemia). These events, either alone or in combination, may contribute towards the progression of AD.40

**Atherosclerosis, Hypertension, and Ischemia**

Cascade Effects in AD

Additionally, for many decades, the role of decreased CBF in the pathogenesis of AD relied on the impairment of vasculature through the lens of atherosclerosis.10,14 Similarities between AD and atherosclerosis have sparked investigation considering that both are found in conjunction with vascular wall thickening and blood vessel occlusion.41 After examination of the cerebral arteries in AD patients, Roher et al42 found cases of significantly increased cerebral artery occlusion in comparison to control groups.41 Additionally, a positive correlation was determined between arterial stenosis and neurofibrillary tangles,42 which is a hallmark of AD. Other studies by Roher et al43 found more widespread intracranial atherosclerosis in AD patients versus nondemented patients. Cognitive dysfunction was also found to be exacerbated in AD patients with cerebral atherosclerosis compared with nonatherosclerotic AD patients, regardless of intracranial or extracranial localization.44 Furthermore, the Nun Study of Aging and AD indicated that patients with multiple brain infarctions have lower cognitive function and higher prevalence of dementia.45,46

Hypertension has been independently identified as a risk factor for AD, but many have reasoned that this is because of hypertension leading to other diseases, which then result in the onset of dementia.45 Throughout time, there have been multiple intuitions about the exact mechanism of hypertension leading to AD: (1) vascular alterations leading to infarcts in the brain, (2) adverse effects on neuronal health leading to deposition of Ab in the brain, and lastly (3) development of cardiovascular disease giving rise to AD.38 Hypertension has also been proposed to evoke AD via ischemia, oxidative stress, inflammation, and small-vessel disease.47,48 More recently, hypertension has been largely noted as the most detrimental vascular risk in the progression of AD.49 This designation is because of its causation of small-vessel disease, which later results in lacunar infarcts, white matter lesions/hyperintensities, and microinfarcts,50 all of which are indicated in AD and accelerate the reduction in CBF.51 Hypertension, however, is an independent risk factor for cardiovascular diseases, including HF,52 indirectly affecting AD through the old lens of cerebral hypoperfusion and the new one of the common pathogenesis of AD and HF as age-related proteinopathies.

One of the main hallmarks of cerebrovascular disease is found in the microvascular structural changes, namely WMH,53 a manifestation of small-vessel disease.54 WMH are a commonality with age progression and a causative agent behind normal cognitive decline.55 However, as imaging and other diagnostic techniques have progressed, it has been shown that small-vessel disease and WMHs in the brain both play a role in the development of AD.56 Population studies using magnetic resonance imaging have also designated a high occurrence of small-vessel disease being associated with higher risk of stroke and dementia.57 These WMH are indicative of vascular lesions and have been found to lower the threshold for the clinical diagnosis of AD.58 Tosto et al59 established that WMH distributions in the brain can progress to AD both directly via neurodegenerative changes and indirectly via aggravation of the tau effect on clinical transformation. This connection has enabled vascular risk factors to be common targets when testing novel therapeutic approaches against AD, such as hypertension and cholesterol treatments.57 However, these studies have been preliminary, and there is much more to be investigated.

In summary, the role of the vasculature in AD progression is one that is implicated primarily via decreased CBF. This remains the main talking point for the vascular hypothesis of AD, which, as seen in Figure 1, claims that many vascular risk factors exert their pathology through cerebral hypoperfusion.60 However, there is a multitude of contributing vascular risk factors, some of which include atherosclerosis, hypertension, small-vessel disease, and BBB dysfunction, all contributing to reduced CBF and the progression of AD, as shown in Figure 1. Much of the discussed literature has focused solely on the vasculature and the brain; however, it was recently proposed that there is a closer link between the brain and the heart than originally expected.39

**Newly Discovered Link Between HF and AD**

Until recently, the only clear link between HF and AD was based on epidemiological data indicating that both of these debilitating conditions have a high incidence of coexisting, especially among older patients. Furthermore, both of these conditions share risk factors; some of which include obesity, sex, high cholesterol, and more importantly age. As described above, a link to cognitive impairment in HF was also recognized in cerebral hypoperfusion and anoxic state.2,61 This theory follows a cascade effect: HF leads to decreased CBF, causing a metabolic energy crisis. This, in turn, causes acidosis and oxidative stress in multiple regions of the brain, eventually culminating in neuronal degradation.24,25,62 Notably, the severity of HF has been shown to positively correlate with the degree of cognitive decline.63,64 and neuroimaging studies have demonstrated a link between HF and structural changes to the brain.25,65 Individuals with HF often exhibit regional
brain atrophy and demyelination, as well as impaired axonal circuit functionality. However, cognitive impairment in HF may have a more complex pathogenic background. In fact, in a more general scope, misfolded protein disease is not limited to the brain. Amylin Cardiomyopathy: a Direct Link Between Heart and Brain Failure

Recent studies focusing on hyperamylinemia, a common disorder in diabetic patients, have revealed a more complex systemic pathogenesis of aggregating amylin. Previous work has found evidence of amylin/amyloid aggregations in the hearts of patients with diabetic cardiomyopathy. Work by Jackson et al sought to examine the cerebrovascular tissue of diabetic patients with vascular dementia of AD for amyloid deposits, wherein they found amylin oligomers and plaques in the temporal gray matter from diabetic patients. Furthermore, researchers found amylin deposition in the brain vasculature and parenchyma of individuals in a specific test group with late-onset AD and no apparent diabetes mellitus. Jackson et al also found some instances where amylin and Aβ depositions were mixed. Expanding on this finding, it has been suggested that amylin and Aβ are connected in terms of wider net pathophysiology. This hypothesis is supported by the fact that both Aβ and amylin can form similar toxic aggregates that can induce inflammation, oxidative stress, and changes in the microvasculature of brain parenchyma. These findings suggest that amylin deposition, and thereby its negative effects on the vasculature and parenchyma of the brain, may participate in the progression of AD.

Common Genetic Profiles Between AD and HF

The concept of AD affecting both the head and heart is reinforced when looking at genetic profiles between connected disease states. Research has shown that there are similar genetic profiles between HF and AD. Work by Li et al found that in the familial forms, these 2 conditions share variations in the PSEN1 or PSEN2 genes. In this specific study, a PSEN1 missense mutation (Asp333Gly) and a PSEN2 missense mutation (Ser130Leu) were associated with both DCM and HF.

Similarly, Gianni et al identified the same missense mutations in the PSEN1 and PSEN2 genes associated with AD in sporadic cases of idiopathic dilated cardiomyopathy (iDCM) and described new genetic variants of the promoter region of the genes affecting the expression levels of the protein. In this study, Gianni et al discovered plaque-like amyloid deposits in the hearts of patients with iDCM. These aggregations of proteins hold a considerable degree of proteotoxicity, inducing cell death. At the microscale, Demuro et al investigated the biochemical mechanisms by which aggregations of soluble amyloid proteins have a neurotoxic effect. Their findings suggest that oligomer amyloid aggregations can disrupt calcium flux homeostasis and thereby cell membrane function. Similar to this proteotoxic effect in neurons, oligomeric aggregates exercise the same effect on cardiomyocyte Ca homeostasis. The toxic role of amyloid deposition as a causal agent for AD was challenged by the failure of some clinical trials targeting amyloid plaques for efficacy of clinical symptoms (the immunoglobulin/albumin combination flebogamma/albutein, and small molecule targeting the β-secretase 1 cleaving enzyme verubecestat) or causing side effects, such as encephalopathy (the humanized anti-Aβ mAb gantenerumab). An initial explanation for the failure of the trials, and therefore the failure of the amyloid theory altogether, included late administration of the drug when the amyloid had triggered neuronal cell death and other terminal changes. However, more recent trials targeting multiple forms of Aβ, such as oligomers in addition to the insoluble fibrils (the mAb crenezumab, aducanumab), did not cause side effects and provided some evidence of clearance of plaques and decline of clinical symptoms. Although waiting for the ongoing phase III clinical trial results (eg, for crenezumab, aducanumab, the BACE (beta site amyloid precursor protein cleaving enzyme))...
inhibitors lanabecestat and elenbecestat) in vitro studies from the del Monte laboratory using atomic force microscopy suggested that a possible explanation for the failure of some of the drugs may reside in targeting fibers versus the more toxic oligomeric species. The study indicated that although dissolving fibers may result in increased release of toxic species, targeting the latter may at least slow the progression of the disease by removing the source of neuronal poison.74

Additional Common Protein Profiles Between AD and HF

Focusing on the heart, the del Monte laboratory analyzed the composition of the aggregates in the myocardium of iDCM patients and identified that cofilin-2—along with its substrate actin and competing protein MLCII (myosin regulatory light chain 2)—was sequestered in the aggregates.79 Cofilin is an actin-depolymerization protein that regulates the turnover of actin in contractile cells and the structural integrity of the cell. In a defective and sequestered state, this protein is also known to play a role in multiple neurodegenerative diseases, such as corticobasal degeneration, William syndrome, fragile X syndrome, and spinal muscular atrophy,12,59,75 as well as other cardiac disorders, such as myocardial ischemia. Recently, the Salloum laboratory described similar changes in cofilin activity, as described in iDCM patients with end-stage ischemic cardiomyopathy.76 This new evidence further supports the link between neurodegenerative disease and various cardiovascular diseases leading to HF.

After this work, del Monte’s group59 analyzed the complex pathology of proteinopathies and sought to discover a closer link between HF and AD by examining the hearts of individuals with AD diagnosis. In this work, Troncone et al9 analyzed the hearts and brains of patients with AD. On investigation, they discovered in the heart, Aβ (both Aβ40 and Aβ42) structurally akin to those found in the brain and, in a retrospective analysis, found that AD patients present with myocardial diastolic dysfunction. Thus, like in traditional cardiac amyloidosis, the pathophysiology of diastolic dysfunction in AD can be described by the accumulation of misfolded proteins in the heart.77

The concept of peripheral accumulation of Aβ in patients with AD is not restricted to the heart. In fact, an early study by Joachim et al78 analyzed Aβ deposition in nonneuronal tissue—most notably, skin and intestine. Eight of the samples from test subjects with AD showed clear and definite evidence of Aβ deposition in these peripheral tissues. Similarly, Aβ40 and Aβ42 aggregates were identified in skeletal muscle of AD individuals using fast performance liquid chromatographic (FPLC) size exclusion chromatography. Researchers discovered elevated levels of Aβ in the temporalis muscles of AD individuals, indicating a possible contributor to elevated concentrations of Aβ plasma levels and potentially indirectly contributing to Aβ deposits in cerebral blood vessels and brain parenchyma.79 Furthermore, these varying levels of Aβ suggest an alteration in APP or Aβ metabolism in peripheral tissues outside of the central nervous system.79

Viewing AD with a new lens as a potential multiorgan disease,59 as shown in Figure 2, it is possible that an inflamed...
and acidic cerebral microenvironment potentially contributes to a defective BBB (a common symptom of AD). This impaired BBB allows permeability of Aβ plaques into the bloodstream. This spreading event in combination with a defective production/clearance homeostasis of misfolded proteins would cause a combination of central nervous system and peripheral symptoms. In the case of HF, the involvement of the heart would accelerate the progression of the disease by contributing to the oxidative stress and acidosis via brain hypoperfusion. However, there might be a possible specific genetic profile—including a miRNA (micro RNA) signature—that accounts for the systemic link between the head and the peripheral organs (Table).

### Conclusions

With recent studies discovering pathogenic mechanisms and possible links between AD and iDCM, including genetic and environmental background, and that AD individuals had significant amounts of Aβ plaques in peripheral organs, it is critical to understand the contribution of the peripheral organs to the overall clinical picture of proteostatic diseases. The failing heart, by reducing CBF to peripheral organs, but also by sustaining the spread of the pathological fragments together with other peripheral organs, activates or aggravates aggregate pathology in the brain. Thus, within this new framework, understanding the disease in its entirety may help in discovering new approaches to delay or reverse proteinopathies involving these 2 vital organs.

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### Disclosures

None.

### References


It seems that there are strong evidences about the direct connection between heart failure (and related diseases) and AD. It appears from the review that soluble amyloid oligomers and insoluble fibrils might be the link to ostensibly connect the two diseases from the clinical perspective. These materials may be produced and circulated throughout the body, depositing in various tissues and organs, including the heart and brain (especially via compromised BBB).

There might be various reasons for the proteins (a-beta, alpha synuclein, tau, immunoglobulin light chain, etc.) to unfold, oligomerize, and aggregate. There have been various studies to look for biochemical causes, but it might be overlooked to examine the possible cause by hemodynamic and mechanical stress and disturbance. As we have been proposing, shear stress and/or fluid pressure could cause the proteins to (partially) unfold and thus become more prone to self-assembly.

Other possible reasons for directly triggering amyloid formation:

* local concentration (obviously; more input and less drainage)
* pH (acidic ?)
* ionic strength (salt)
* aldehyde (chemical linker or promoter for unfolding)
* temperature (lower temperature induces precipitation)
* nucleation seeding (structurally similar amyloids or fibrils)
*