Review Article

Behavioral and Psychological Symptoms in Alzheimer’s Disease

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Neuropsychiatric symptoms (NPS) such as depression, apathy, aggression, and psychosis are now recognized as core features of Alzheimer’s disease (AD), and there is a general consensus that greater symptom severity is predictive of faster cognitive decline, loss of independence, and even shorter survival. Whether these symptoms result from the same pathogenic processes responsible for cognitive decline or have unique etiologies independent of AD-associated neurodegeneration is unclear. Many structural and metabolic features of the AD brain are associated with individual neuropsychiatric symptoms or symptom clusters. In addition, many genes have been identified and confirmed that are associated with symptom risk in a few cases. However, there is no single gene strongly predictive of individual neuropsychiatric syndromes, while functional and structural brain changes unique to specific symptoms may reflect variability in progression of the same pathological processes. Unfortunately, treatment success for these psychiatric symptoms may be lower when comorbid with AD, underscoring the importance of future research on their pathobiology and treatment. This review summarizes some of the most salient aspects of NPS pathogenesis.

1. Introduction

The recent establishment of a professional interest area (PIA) within the International Society to Advance Alzheimer’s Research and Treatment (ISTAART) devoted to the neuropsychiatric symptoms (NPS) of Alzheimer’s is a sign of the emerging consensus among researchers and clinicians alike that these symptoms are major components of Alzheimer’s disease (AD) and significant influences on both patient and caregiver quality of life (QOL) [1]. Indeed, neuropsychiatric symptoms such as apathy, depression, aggression, agitation, sleep disruption, and psychosis are now recognized as core symptoms of AD that are expressed to varying degrees throughout the course of the illness [2]. In addition to providing insight into AD pathology, specific neuropsychiatric and behavioral anomalies during the early prodromal phase of mild cognitive impairment (MCI) may have prognostic values. For example, late-life depression increases AD risk by 2-fold [3]. In this review, the major neuropsychiatric and behavioral symptoms of AD are reviewed with emphasis on how these symptoms may illuminate disease pathogenesis or provide prognostic information. Alzheimer’s dementia is the end result of multiple pathogenic processes including aberrant amyloid processing [4, 5], changes in lipid metabolism due to apolipoprotein E (APOE) risk alleles [6, 7], tau hyperphosphorylation [8], protein misfolding and endoplasmic reticulum (ER) stress [9], vascular dysfunction [10], oxidative stress and mitochondrial dysfunction [11, 12], neurotrophic factor dysregulation [13], disrupted leptin signaling [14], fibrin clots [15], and processes mediated by a myriad of other AD-associated gene [16], and the pathogenic processes also occurred in major neuropsychiatric symptoms (Figure 1). It is likely that these processes target nonoverlapping neural networks, accounting for difference in disease progression and the variability in neuropsychiatric symptoms.

The neuropsychiatric symptoms and behavioral anomalies of AD have a significant impact on patient QOL and are thought to be predictive of eventual (or more severe) dementia [17, 18], more extensive neurodegeneration [19], loss of functional independence and institutionalization [20], and early death [21]. Thus, there is general agreement that these neuropsychiatric symptoms and behavioral anomalies
Figure 1: Possible mechanism linking the neuropsychiatric symptoms (NPS) with AD. NPS such as depression, apathy, aggression, and psychosis shared some pathogenic processes (in red color) with AD, while they also have their unique pathogenic processes.

Based on studies in depressed non-AD populations, early studies on the pathophysiology of depression in AD focused on serotonergic transmission. One of the earliest studies reported an association between major depression in AD at baseline and 5-HT (2A) and 5-HT (2C) receptor polymorphisms, with CC carriers of the 5-HT (2A) C102 allele five times more likely than heterozygotes and 5-HT (2C) Ser allele carriers 12 times more likely than 5-HT (2C) Cys allele carriers to develop depression [29]. Moreover, reduced 5-HT (1A) receptor expression was specifically correlated with depressive symptoms [30]. In contrast, Pritchard and colleagues found no significant association between depression in AD and either the C allele/CC genotype of the T102C variant of 5HT (2A) or the cys23ser variant of 5HT (2C) receptor, although these alleles were associated with psychosis and aberrant motor behavior [31]. Moreover, no association was found between depression in AD and alleles of the serotonin transporter (SERT) [32]. Similarly, although SERT expression was reduced in the frontal cortex of AD patients, there was no difference in expression in patients with or without comorbid depression [33]. It is possible that serotonergic dysfunction may be heterogeneous among brain regions across patients, accounting for these differences in association. In addition to 5-HT signaling, elderly subjects destined to exhibit signs of major depression were more likely to harbor the GG genotype of the tumor necrosis factor (TNF)-alpha 308 (G/A) SNP variant, implicating inflammation in late-onset MD [34].

Early studies also examined the relationship between depression and molecules implicated in general AD pathology, particularly Aβ and APOE 4, the strongest risk allele for AD. Early onset depression was associated with a higher serum Aβ40/Aβ42 ratio [19], suggesting that depression may
be associated with AD pathogenesis. One early small sample study found no association between APOE genotype and depression in AD [35], although subsequent studies have demonstrated that APOE genotype can modify the effects of other genes associated with the neuropsychiatric symptoms of AD (see below). A higher serum concentration of Aβ at baseline predicted both depression and AD over 5 years suggesting shared etiology [36]. Plasma GABA was positively correlated with depression and apathy scores on the NPI in AD patients [37].

In addition to gene association studies, the pathogenesis of AD has also been examined by various neuroimaging modalities, which have revealed morphological and metabolic signs of neurodegeneration in the AD brain specifically associated with depression. Compared to nondepressed AD patients, those with depression exhibited hypoperfusion in the left frontal lobe on single-photon emission computed tomography (SPECT) images [38] and reduced glucose metabolism in the dorsolateral prefrontal regions as revealed by (18)F-fluorodeoxyglucose PET [39]. Correlation analysis of brain SPECT and NPI score revealed a region in the left middle frontal gyrus (Brodmann area 9) specifically associated with depressive symptoms [40]. Depression in AD has also been associated with specific neurochemical changes; GDS scores but not agitation scores were correlated with choline/creatine ratio in left dorsolateral prefrontal cortex [41]. Cortical atrophy associated with depression was observed in wide regions of the prefrontal cortex and temporal cortex [42] and decreased gray matter volume in the left inferior temporal gyrus was confirmed in an independent study [43]. Depressed AD patients also exhibited greater white matter atrophy in frontal, temporal, and parietal lobes than AD patients without depressive symptoms [44]. One study also reported lesions in the caudate nucleus and lentiform nucleus of AD patients with late-onset depression [45]. Expansion of the lateral ventricles was also correlated with depression, general cognitive decline, and poor outcome [46]. Thus, depression is associated with both gray and white matter atrophy, particularly in specific regions of the prefrontal cortex.

However, it remains unclear if depression results from AD or conversely if geriatric depression is a risk factor for AD. In the first case, depression may be a psychological response to AD or result from the same pathogenic processes that lead to the other symptoms of AD (e.g., aberrant amyloid Aβ processing, tau hyperphosphorylation, etc.) [47]. Depression in AD is associated with accelerated cortical regression and white matter atrophy, particularly in frontal and temporal areas. It has been proposed that AD-associated degeneration may eventually damage regions involved in regulation of mood, a finding consistent with the high rates of depression in severe AD. Nonetheless, several genetic risk factors for major depression appear to increase the risk of depression in AD but not AD without depression, so the emergence of depression may not be entirely dependent on AD pathogenesis. For example, the tryptophan hydroxylase-1 (TPH1) A218C allele, monoamine oxidase A (MAOA) variable number tandem repeat (VNTR), and BDNF Val66Met allele were associated with depression in females with AD, with significantly increased likelihood of comorbid AD and depression in homozygous TPH1 A-allele and MAOA VNTR carriers [48]. In this same study, there was also a significant association between the chaperone FK506 binding protein 5 (FKBP5) rs1360780 SNP and depression in all AD patients. In addition, homozygous carriers of the rs10410544T allele of the SIRT2 gene (encoding an NAD-dependent deacetylase possibly involved in cell cycle regulation) may have reduced depression risk in AD [49]. Aside from the Val66Met allele of BDNF, the C allele of the SNP GI1757C and the A allele of GI96A were also more common in AD patients with depression [50]. One of the strongest associations with late-onset AD and depression is that with the transforming growth factor 1 (TGF-1) gene the CC genotype of the +10T/C SNP was associated with AD and conferred a 5-fold increase in depression in AD as well as an increase in depression severity [51]. Finally, the presence of the APOE 4 allele increased depression in women with AD by 4-fold [52]. In contrast, another study reported that APOE 4 was associated with anxiety but not depression [53], while others have found no association between APOE 4 and neuropsychiatric symptoms [35].

Whether depression increases AD risk in premorbid or MCI patients is still a matter of debate. In an Italian study, newly diagnosed AD patients with persistent depression exhibited a greater cognitive decline over one year, and patients with incident depression demonstrated the greatest drop in cognitive function, while cognitive decline in cases with resolved depression was not different from nondepressed AD patients [54]. Late-onset depression does increase the risk of progression to MCI, but chronic depression was associated with only a modest increase in the risk of MCI-to-AD transition. Another Italian study reported that apathy but not depression was associated with MCI to AD transition [55]. In contrast, the Honolulu-Asia Aging Study using the Center for Epidemiological Studies depression scale (CES-D) reported that depression was an independent risk factor for cognitive decline in AD. Moreover, the effect was independent of pathological progression, such as increases in the number/density of neurofibrillary tangles (NTs), Lewy bodies, or ischemic lesions [56]. These differences in the reported prognostic value of depression may depend on diagnostic criteria; for example, the Vienna Transdanube Aging study did report an association with AD emergence over a 5-year period in 75-year-old individuals with no history of depression, but only 1 of 9 depression subsyndromes, “loss of interest,” was associated with AD risk [57]. Another report concluded that depression does appear to increase the risk of transition from MCI to dementia, but this effect was stronger for all-cause dementia and vascular dementia than AD [3] or exclusive to vascular dementia [58].

Regardless of this etiological relationship, it is clear that AD-associated depression markedly reduces cognitive capacity, QOL, and activities of daily function (ADF). Thus, treatment of depressive symptoms is expected to benefit AD patients. However, there have been relatively few controlled clinical trials on antidepressant therapy for depression in AD and clinical response is generally poor to modest [59–64]. The uncertain relationship between AD and depression...
undoubtedly arises in part from diagnostic uncertainty. As mentioned, AD is only confirmed at autopsy while estimates of depression vary marked depending on the instruments used. Furthermore, only certain depressive symptoms may be associated with AD [57]. In sum, depression may be a modest risk factor in premorbid patients [59–64] for additional reviews but when present it markedly reduces cognition, QOL, and ADL in AD patients.

3. Apathy

Apathy is defined by a cluster of motivational deficits such as loss of goal-directed cognition, action, and emotion [65]. Like other neuropsychiatric symptoms associated with AD, persistent apathy is predictive of more rapid cognitive decline compared to AD without apathy. Apathy and depression are often comorbid. In one relatively large cohort (255 patients), 47.9% of the study group had depression, 41.6% apathy, and 32.4% both, with smaller prevalence of depression and apathy alone (15.4% and 9.4% resp.) [23]. A similar pattern has been reported in other studies (e.g., 23% depression only, 23% depression + apathy, and 20% apathy only) [66, 67]. This frequent comorbidity suggests shared etiology. Indeed, like depression, apathy is generally associated with hypofrontality as well as serum GABA [37]. However, apathy was specifically correlated with hypometabolism in left orbitofrontal areas while depression was associated with hypometabolism in left dorsolateral prefrontal regions [39]. Scores on the frontal assessment battery (FAB) for executive function are decreased by both apathy and depression alone, but the largest decrease was observed in comorbid patients [68]. This hypofrontality has been correlated with AD-associated pathogenesis [68]. Retention of the (11C) Pittsburgh compound-B (PIB) under PET to reveal Aβ plaques was higher in the bilateral frontal cortex of patients with apathy as determined by the NPI apathy subscale than in AD patients without apathy, and apathy scores were positively correlated with PIB signal in bilateral frontal and right anterior cingulate cortices. No correlations were found between PIB and any other NPI subscale, including depression, while depression alone did not increase risk of transition [72]. Also distinct from depression, apathy tends to stabilize during AD progression while apathy tends to increase [73].

4. Agitation and Aggression

Agitation and aggression are significant dangers both to patients and caregivers. Like other behavioral and neuropsychiatric abnormalities, rates of agitation and aggression correlate with cognitive decline, loss of independence, and other metrics of poor outcome [74]. Aggression and agitation are more common in male patients [75]. Among nursing home residences, the severity of cognitive decline/dementia was correlated with physical agitation and verbal aggression as measured by the Cohen-Mansfield Agitation Inventory (CMAI). The intensity of dementia disorders is associated most strongly with physical agitation and verbal aggression. Aggression- and (or) agitation-specific changes in neurochemistry and neuropathology have been observed. Based on the well-established correlation between frontal lobe serotonergic dysfunction and aggression, many studies have investigated the correlations between aggression presence/severity and 5-HT signaling molecules. Among the first such studies reported a significant association between aggression in AD and the more highly expressed serotonin transporter long variant (5-HTTPL) [76]. Male AD patients with a history of agitation/aggression were also more likely to harbor the C allele of the 5-HT synthetic enzyme tryptophan hydroxylase A2I8C intronic polymorphism [77]. Aggressive patients also exhibited an altered 5-HT6 receptor to choline acetyltransferase (ChAT) ratio in frontal and temporal cortices [78] and reduced 5-HT1A binding in temporal cortex after controlling for dementia severity [79]. However, others have found a much more complex interaction among gender, dementia severity, aggression, and serotonergic function [80]. Serotonin transporter polymorphisms have also been linked to a combined aggressive/psychotic phenotype [81]. Polymorphisms of dopamine receptor alleles also confer complex associations among psychotic symptoms and aggressiveness. Psychosis and aggression were more frequent in DRD1 B2/B2 allele carriers [82]. In a subsequent study, carriers of the DI B allele were more prone to aggression and (or) to experience hallucinations, while carriers of the DRD3 I allele were more likely to experience delusions [83]. In fact, there is a strong correlation between psychosis and aggression. Delusions are among the strongest predictors of aggression [84], possibly because aggression is driven by specific delusions of persecution [85, 86].

The CMAI scores, but not GDS scores, were negatively correlated with the N-acetylaspartate (NAA) to creatine ratio
in the left posterior cingulate gyrus [41], indicating neurochemical changes specific to the aggressive AD phenotype. Moreover, only agitation/irritability scores were correlated with Aβ42 accumulation [87] and only Behavioral Pathology in Alzheimer’s Disease (Behave-AD) aggressiveness subscale scores correlated with serum BDNF [70] in AD and MCI patients. One study demonstrated a correlation between aggressive symptoms and the magnitude of locus coeruleus damage, suggesting a role for reduced norepinephrine or compensatory changes in adrenergic receptors in aggression [88]. In fact, a test of β2 receptor function, the growth hormone (GH) response to clonidine, was blunted in aggressive patients compared to nonaggressive AD patients [89]. Higher NPI Agitation and Aggression subscale scores were associated with greater atrophy in a large number of regions of interest in the frontal and cingulated cortices as well as the insula, amygdala, and hippocampus [90]. The phospho-tau to tau ratio was also higher postmortem in the frontal lobe of AD patients with aggression compared to those without [91]. Similarly, hippocampal NTs and hyperperfusion in the mesial temporal lobe [92] were associated with symptoms in AD. The frequency of the APOE 4 allele was also associated with aggression [93].

Neuroleptics and other antipsychotics may be modestly effective in treating aggression and agitation, but longer-term treatment is associated with a significant increase in adverse events [94].

5. Psychosis

Psychotic symptoms, including delusions and hallucinations, are obviously the most salient and serious neuropsychiatric symptoms associated with AD, also generally the least frequent symptoms during the early stages of AD [80]. Like depression, the emergence of psychotic symptoms predicts greater or more rapid cognitive decline [44, 95]. Hallucinations are among those noncognitive AD symptoms, also including cognitive impairment level, physical aggression, and depressive symptoms, strongly predictive of institutionalization [96]. A longitudinal study found that psychosis in AD (observed in 7.8% of patients) was associated with greater initial cognitive dysfunction, more accelerated cognitive decline, and greater mortality [97]. Like aggression, to which it is strongly correlated, psychosis is associated with difference in the 5-HT system, such as a higher frequency of the C allele and CC genotype of the T102C variant of 5HT (2A) receptors in patients with hallucinations and delusions [31]. There may also be an association between the SERT long form and psychosis in AD [96]. Moreover, psychosis is associated with CSF tau, suggesting more severe tauopathy in psychotic patients [98] and with greater intracellular accumulation of hyperphosphorylated tau [99]. The APOE 4 allele also increases psychosis risk [100]. In general, AD-associated psychosis follows the severity of AD-associated neurodegeneration and cognitive dysfunction [101]. 18F-Fluorodeoxyglucose positron emission tomography revealed reduced metabolic activity in the right lateral frontal cortex, orbitofrontal cortex, and bilateral temporal cortex in patients with delusions, overlapping with areas associated with loss of memory and insight [102]. Atrophy of the supramarginal cortex of the parietal lobe was predictive of increasing hallucinations over time. Active psychosis is associated with hypofrontality, particularly in orbitofrontal regions [102], and thus is strongly correlated with disruption in executive function, particularly working memory [103]. Another study found lateral frontal, lateral parietal, and anterior cingulate atrophy in AD patients with psychosis, with the lateral frontal region most severely degenerated [104]. Individual delusions may be associated with specific abnormalities in neural processing as evidence by PET imaging [105], with delusions of persecution associated with hyperperfusion in the precuneus and hyperperfusion in the insula and thalamus. Other delusional forms are associated with distinct changes in perfusion, metabolism, receptor binding, and structural alternations [106]. Persecutory delusions occur early during the progression of AD and associate with disruption of frontostriatal circuits [107]. One study found delusions in 27.4% of AD patients, with paranoid delusions being the most common (60.3%), followed by misidentification delusions (19.0%), and then mixed delusions (17.5%), the latter appearing later and associated with greater cognitive impairment [108]. Psychotic patients show greater Aβ1-42 at autopsy [109]. A recent genome wide association study identified an intergenic region on chromosome 4 (rs753129), SNPs upstream of SLC2A9 (rs6834555), and within the neuronal Ca2+ sensor (NCS) proteins VSNL1 (rs4038131) as promising regions for specific associations with psychosis [110]. Note, however, that VSNLs are associated with Aβ and tau and so may reflect the overall severity of AD neurodegeneration rather than psychosis per se. Other possible associations specific to AD-associated psychosis include that with the dopamine oxidase A (DOA) gene and muscarinic receptors in orbital frontal cortex [110].

6. Conclusion

The neuropsychiatric symptoms in the early stages of AD are predictive of more rapid deterioration of cognitive function. These symptoms may simply reflect more rapid and extensive neuronal death caused by the myriad of primary and secondary degenerative processes associated with AD. In this case, the appearance of different neuropsychiatric symptoms reflects variability in the progression of neurodegeneration across neural systems. There are genes associated with specific symptoms or symptom clusters, but none appear to exert strong and specific effects. Moreover, many of these association studies have not been followed up, suggesting some publication bias for positive results. Regardless of the dependence of these symptoms on AD pathogenesis and cognitive decline, the deleterious effect of these symptoms on both patient and caregiver quality of life warrant further studies on more effective treatments.

Conflict of Interests

We declare that we have no conflicts of interest.
Acknowledgments

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