Assessment and Management of Psychiatric Symptoms in Alzheimer's Disease

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ABSTRACT

Neuropsychiatric symptoms are near universal among patients with neurodegenerative disorders, thus almost all patients experiencing at least one of them during the course of their illness. Currently, there is no FDA-approved medication for the treatment of neuropsychiatric symptoms in the course of neurodegenerative disorders with the exception of pimavanserin approved for the treatment of psychosis in the course of Parkinson's disease. Pimavanserin shows benefit in addressing psychosis in Alzheimer's disease. Although used "off label", a large number of antidepressants, mood stabilizers, typical and atypical antipsychotics are prescribed for behavioral disturbances in persons with neurodegenerative disorders. In this review, the terminology, clinical features, demographic and epidemiologic facts, diagnostic criteria and management strategies for selected neuropsychiatric symptoms like depression, psychosis with a special attention to apathy and agitation in the course of Alzheimer's are discussed. Data on the suspected pathophysiologic mechanisms are also reviewed. Several issues relating to future therapies that may impact patients are briefly mentioned.

Abbreviations: AChEI: Acetylcholinesterase Inhibitor; AD: Alzheimer's Disease; ADMET: Alzheimer’s Disease Methylphenidate Trial; AE: Adverse Effect; AES: Apathy Evaluation Scale; APA: The American Psychiatric Association; BPRS: Brief Psychotic Rating Scale; BPSD: Behavioral and Psychological Symptoms of Dementia; CBT: Cognitive Behavioral Therapy; CMAI: Cohen-Mansfield Agitation Inventory; CGI-C: Clinical Global Impression of Change; CGI-S: Clinical Global Impression of Severity of Illness; CGI-I: Clinical Global Impression of Improvement; CSDD: Cornell Scale for Depression in Dementia; DPA: Dopaminergic Agonist; DMAS: Dementia Mood Assessment Scale; NBRSA: Neurobehavioral Rating Scale-Agitation Subscale; NDD: Neurodegenerative Disorders; NPS: Neuropsychiatric Symptoms; MADRS: Montgomery-Asberg Depression Rating Scale; MAOI: Monoamine Oxidase Inhibitor; MMSE: Mini Mental State Examination; MRI: Magnetic Resonance Imaging; NIMH-dAD: The National Institute of Mental Health Depression of Alzheimer’s Disease; NPI: Neuropsychiatric Inventory; PT: Physical Therapy; RCT: Randomized Clinical Trial; SNRI: Serotonin-Norepinephrine Reuptake Inhibitor; SSRI: Selective Serotonin Reuptake Inhibitor; TCA: Tricyclic Antidepressant; rTMS: Repetitive Transcranial Magnetic Stimulation; UPDRS: Unified Parkinson's Disease Rating Scale

Introduction

Behavioral and psychological symptoms classified as neuropsychiatric symptoms (NPS) commonly accompany dementia syndromes in the course of neurodegenerative diseases (NDD) [1,2]. Cardinal psychiatric symptoms such as depression, apathy, psychosis or agitation complicate the clinical presentation of Alzheimer’s disease (AD) and Parkinson’s diseases (PD) and some are considered core symptoms of frontotemporal dementia and dementia with Lewy bodies (DLB) [1]. NPS are near universal among patients with neurodegenerative disorders (NDD), thus almost all patients with AD experiencing at least one of them...
During the course of their illness [3], NPS are usually assessed by comprehensive psychiatric evaluation but more structured tools like the Neuropsychiatric Inventory (NPI) commonly used in the setting of trials can be of help in clinical practice [2]. The group of common and debilitating NPS include apathy, irritability, agitation, depression, delusions, hallucinations, anxiety, disinhibition, aberrant motor behavior, sleep disturbances and eating abnormalities [2]. NPS can manifest themselves at all types and stages of NDD (especially when a dementia syndrome is present), often cluster, tend to be persistent and are associated with excess morbidity and mortality, contribute to patients’ distress and caregiver burden.

They are NPS are linked to increased healthcare use, healthcare costs, and institutionalization [4]. The neurobiology of NPS is extremely complex. Dysfunction in frontal-subcortical and corticocortical networks was proposed as a model of NPS [5]. In terms of neurochemistry dysfunction in ascending monoaminergic systems involving serotonin, norepinephrine and dopamine, glutamate-mediated excitatory neurotoxicity, tau-mediated pathology, inflammation play a role in the occurrence of NPS in NDD (Murai et al.); [6,7]. Despite the significant burden of these symptoms, there are few recommended, evidence-based treatments including pharmaceuticals [8,9]. Pharmacotherapy in the geriatric population in general is challenging due to age-associated changes in pharmacodynamics and pharmacokinetics as well as high rates of medical comorbidities and concomitant medications, which increases risk for polypharmacy, drug-drug interactions, and adverse drug effects [9]. Additionally, there are risks attributed to use of antipsychotics, antidepressants, anxiolytics and /or mood stabilizers in elderly, and specific risks to patients with dementia which makes a decision about treatment complicated both for providers and patients and caregivers [8].

Psychotropics, especially antipsychotic medications, may alleviate certain NPS, but may have severe adverse effects including increased risk of involuntary movements, cerebrovascular events, falls, and death [9]. Effectiveness of psychopharmacologicals in the treatment of NPS is a subject of ongoing debate. AD treatments may have effects on NPS and can affect decisions regarding treatments including psychotropic agents. Cholinesterase inhibitors may reduce the emergence of NPS and have a role in their treatment. These agents may delay initiation of or reduce the need for other drugs such as antipsychotics thus ACHI’s should be initiated, optimized and maintained for the management of both cognitive symptoms as well as NPS [7]. Currently, there is no FDA-approved medication for the treatment of NPS in NDD with the exception of pimavanserin which was recently approved for the treatment of psychosis in the course of PD [10]. Although used “off label”, a large number of antidepressants, mood stabilizers, typical and atypical antipsychotics are prescribed for behavioral disturbances in persons with NDD [8,11]. Parallel to robust efforts in pharmacological trials there is an ongoing need to assess and verify existing, as well as create new behavioral strategies.

Cohen-Mansfield [12] conceptualizes behavioral disorders in the course of NDD as representing unmet personal needs such as pain and any other somatic discomfort, need for social contact, and stimulation to alleviate boredom. Those needs should be recognized and successfully addressed by nonpharmacologic interventions. Therapeutic approaches should be individually tailored to each patient with NPS using behavioral management techniques, caregiver education and support, problem solving and communication skills training, music therapy, aromatherapy and modified cognitive behavioral and interpersonal therapies. With few exceptions initiating pharmacotherapy should occur only after treating or eliminating underlying medical or environmental factors and should be limited to cases where nonpharmacological measures have failed [12]. All patients should be carefully monitored for development of adverse events and side effects during a time-limited treatment course; symptoms often resolve over time regardless of medication use [11]. An ongoing assessment of benefit versus harm should continue throughout the course of treatment with periodic consideration of withdrawing the medication [8,13].

**Depression of Alzheimer’s Disease**

Depression which develops during the course of AD is conceptualized as one of NPS, “depression of AD” [14]. The concept was defined as a depressive syndrome with prominent decreased affect, irritability, agitation, and anxiety, diminished attention and fatigue but less evidence of guilt and suicidality than major depressive episode. Depression of AD is relatively common (affecting up to 50% of persons with AD) and persistent with 50%-60% of untreated depressed patients with AD remaining depressed at 1-year follow-up [14]. A National Institute of Mental Health Work Group developed a set of diagnostic criteria for depression of AD (NIMH-dAD) (Table 1). These criteria were derived from DSM-IV criteria for major depression, with some distinctions. The number of symptoms required for a diagnosis of depression was decreased from five to three. The duration and frequency of depressive symptoms was also decreased; symptoms need only be present together within the same 2-week period. The decreased ability to think and concentrate was eliminated. The criteria for anhedonia were modified to focus on decreased affect and pleasure associated with social and other activities. Social isolation/withdrawal and irritability were added as new symptoms [15]. So far neither DSM 5 related version nor correction of NIMH-AD had been published.
Table 1: Criteria for depression of Alzheimer’s disease.

| Criteria for Depression of Alzheimer’s Disease |
|-----------------------------------------------|
| Three (or more) of the following symptoms must be present during the same 2-week period and represent a change from previous functioning. At least one of the symptoms must either be 1) Depressed mood or 2) Decreased positive affect or pleasure |
| A | 1) Clinically significant depressed mood |
| 2) Decreased positive affect or pleasure in response to social contacts and usual activities |
| 3) Social isolation or withdrawal |
| 4) Disruption in appetite |
| 5) Disruption in sleep |
| 6) Psychomotor changes |
| 7) Irritability |
| 8) Fatigue or loss of energy |
| 9) Feelings of worthlessness, hopelessness, or excessive or inappropriate guilt |
| 10) Recurrent thoughts of death, suicidal ideation, plan or attempt |
| B | All criteria are met for Dementia of the Alzheimer Type (DSM-IV) |
| C | The symptoms cause clinically significant distress or disruption in functioning |
| D | The symptoms do not occur exclusively in the course of delirium |
| E | The symptoms are not due to the direct physiological effects of a substance |
| F | The symptoms are not better accounted for by other conditions such as major depressive disorder, bipolar disorder, bereavement, schizophrenia, schizoaffective disorder, psychosis of Alzheimer disease, anxiety disorders, or substance-related disorders |

Note: Adapted from: Teng E, Ringman JM, Ross LK, Mulnard RA, Dick MB, Bartzikis G, Davies HD, Galasko D, Hewett L, Mungas D, Reed BR, Schneider LS, Segal-Gidan F, Yaffe K, Cummings JL; Alzheimer’s Disease Research Centers of California-Depression in Alzheimer’s Disease Investigators. Diagnosing depression in Alzheimer disease with the national institute of mental health provisional criteria. Am J Geriatr Psychiatry 2008 Jun;16(6):469-77.

Rationale for Interventions and Treatment

Despite the widespread antidepressant use (almost 50% of patient with dementia are on antidepressants) there is mixed evidence regarding the benefits from the use in AD depressed patients. Many trials have been carried out on small numbers of patient and were underpowered to detect differences. Variable trial methods, comorbid conditions and differences in the administered antidepressants further confound findings. Use of antidepressants in AD and other NDD is based on their use in major depressive disorder pending better evidence for or against their use in depression of Alzheimer’s disease. Acetylcholinesterase inhibitors (AChEIs) studies provide some evidence of benefit from their use. Among the NPS in the course of AD most likely to improve appear to be apathy and depression followed by aberrant motor behavior [16]; (Howard et al.). A secondary analysis of patients with severe behavioral disturbances previously treated with donepezil and sertraline, Cummings et al. [17] suggest that donepezil reduces behavioral symptoms, particularly mood disturbances and delusions, in patients with AD. A withdrawal study by Holmes et al. [18] provided additional evidence to support the use of donepezil in the treatment of neuropsychiatric symptoms (including depression) in patients with mild to moderate AD with marked NPS.

Following open-label treatment with donepezil patients randomized to placebo showed a significant worsening of neuropsychiatric symptoms and a worsening of caregiver distress compared with a continued improvement in those who remained on donepezil treatment. Optimization as well as maintenance treatment with AChEIs should be considered as a step in the management of depression of AD. The American Psychiatric Association (APA) recommends a trial of an antidepressant to treat clinically significant, persistent depressed mood in patients with dementia. Selective serotonin uptake inhibitors (SSRIs) are preferred because of their favorable safety profile [19]. Sertraline, citalopram or escitalopram in low doses are the most appropriate first-line agents. Other SSRIs like fluoxetine and paroxetine are not recommended as a first line SSRI’s due to debatable efficacy and unfavorable mostly anticholinergic effects [20]. This dose might be increased weekly, if tolerated, to a maximum of 150 mg of sertraline or 40 mg of citalopram per day with close monitoring of side effects. Although improvement should occur within 4 to 6 weeks at the target dose, a longer period may be required to reach full effect [20].

If patients do not respond to SSRI switching to a different agent or augmenting a treatment with second agent should be considered. Especially for patients who have psychotic symptoms or agitation along with depression, an atypical antipsychotic in a small dose might be considered [21,22]. An anticonvulsant in smaller doses (the best evidence is for carbamazepine) might be considered as additional therapy to an antidepressant if there is moderate or severe agitation [23]. Switching to an antidepressant from a different class (as opposed to augmentation) is recommended in cases of severe side
effects induced by initial medication. Preferred second-line agents are selectove norepinephrine reuptake inhibitors properties (SNRIs) such as venlafaxine or duloxetine, or antidepressants with a mixed pharmacology (mirtazapine, bupropion). Evidence for benefit from use of non-SSRI antidepressants specifically for depression in AD is lacking. Tricyclic antidepressants are not recommended due to lack of convincing evidence and the occurrence of various side effects including anticholinergic ones. Psychiatric hospitalization should be considered an option in complex severe cases.

For patients with severe, refractory depression, electroconvulsive therapy (ECT) might be considered, especially if there is risk of self-harm or harm to others [20]. There is evidence for ECT as an effective and well-tolerated option for treating depression in people with dementia [24]. The situation of limited effectiveness of pharmaceuticals supports nonpharmacological interventions in depression of AD. Interpersonal psychotherapy (IPT) especially versions modified to address the needs of older adults with mild cognitive deficits (IPT-CI) [25]. Cognitive behavioral therapy (CBT) has been adapted for depressed older adults with mild stages of dementia [26]. Methods specifically developed to treat geriatric depression such as home delivered problem adaptation therapy (PATH) or problems solving skills and caregiver training show benefit [27].

Table 2: Criteria for apathy in neurodegenerative disorders.

| Criteria for Apathy in Neurodegenerative Disorders |
|--------------------------------------------------|
| **A** Loss of or diminished motivation in comparison to the patient’s previous level of functioning and which is not consistent with his age or culture. These changes in motivation may be reported by the patient himself or by the observations of others. |
| **B** Presence of at least one symptom in at least two of the three following domains for a period of at least four weeks and present most of the time. |
| **Domain B1-Behaviour:** Loss of, or diminished, goal-directed behavior as evidenced by at least one of the following: |
| **Initiation Symptom:** loss of self-initiated behavior (for example: starting conversation, doing basic tasks of day-to-day living, seeking social activities, communicating choices) |
| **Responsiveness Symptom:** loss of environment-stimulated behavior (for example: responding to conversation, participating in social activities) |
| **Domain B2-Cognition:** Loss of, or diminished, goal-directed cognitive activity as evidenced by at least one of the following: |
| **Initiation Symptom:** loss of spontaneous ideas and curiosity for routine and new events (i.e., challenging tasks, recent news, social opportunities, personal/family and social affairs). |
| **Responsiveness Symptom:** loss of environment-stimulated ideas and curiosity for routine and new events (i.e., in the person’s residence, neighborhood or community). |
| **Domain B3-Emotion:** Loss of, or diminished, emotion as evidenced by at least one of the following: Initiation Symptom: loss of spontaneous emotion, observed or self-reported (for example, subjective feeling of weak or absent emotions, or observation by others of a blunted affect). |
| **Responsiveness Symptom:** loss of emotional responsiveness to positive or negative stimuli or events (for example, observer-reports of unchanged affect, or of little emotional reaction to exciting events, personal loss, serious illness, emotional-laden news). |
| **C** These symptoms (A - B) cause clinically significant impairment in personal, social, occupational, or other important areas of functioning. |
| **D** The symptoms (A - B) are not exclusively explained or due to physical disabilities (e.g. blindness and loss of hearing), to motor disabilities, to diminished level of consciousness or to the direct physiological effects of a substance (e.g. drug of abuse, a medication). |

Note: Adapted from: Mulin E, Leone E, Dujardin K, Dellaix M, Leentjens A, Nobili F, Dessi B, Tible O, Agüera-Ortiz L, Osorio RS, Yessavage J, Dachevsky D, Verhey FR, Cruz Jentoft AJ, Blanc O, Llorca PM, Robert PH. Diagnostic criteria for apathy in clinical practice. Int J Geriatr Psychiatry 2011 Feb; 26(2):138-65.
Rationale for Interventions and Treatment

Psychostimulants including methylphenidate and modafinil have been used to treat apathy in AD. Methylphenidate acts by blocking the dopamine transporter and norepinephrine transporter, leading to increased concentrations of dopamine and norepinephrine within the synaptic cleft. Methylphenidate but not modafinil proved to be effective in reducing apathy in AD in a small cross-over trial [30] and was further assessed in the larger, multicenter, double-blind controlled trial Alzheimer’s Disease Methylphenidate Trial (ADMET) [31]. In ADMET, methylphenidate (20 mg daily for 6 weeks) was associated with a significant reduction in apathy symptoms. Two study outcomes measures CGI-C and NPI apathy score, showed diminished apathy symptoms with methylphenidate treatment. Adverse events and side effects were modest. The results suggest that methylphenidate treatment may have clinical utility in treating apathy of AD with a potential for improving cognition as well. Besides promising results from dopaminergic agents, AChEIs appear to be successful in improving symptoms of apathy in AD [16]. Symptoms may worsen after discontinuation of AChEIs [18]. Optimization and maintenance treatment with AChEIs should thus be considered as a crucial step in the management of apathy of AD.

Psychosis of Alzheimer’s Disease

Extent of the Problem, Pathogenesis and Overall Impact

Estimates of the incidence of psychosis in AD range widely from 10% to 75% [32]. The common psychotic symptoms reported in the AD patients are delusions and hallucinations followed by misidentification phenomena [1]. Hallucinations are predominantly visual. Auditory phenomena especially of a schizophrenic quality are rare in AD [33]. Delusions in the course of AD are typically paranoid, non-bizarre, and simple. Delusions tend to recur or persist for several years in AD patients, but active and vivid perception and delusions have a tendency to diminish in intensity in the course of cognitive decline, shallow insight and decreasing ability for verbal expression [33]. In a study which compared AD delusional and non-delusional subjects, those with delusions were significantly older, with higher age at onset and cognitive impairment, a more severe stage of dementia, and more depressive symptoms than AD patients with no delusional symptoms.

Disease duration was slightly higher in AD delusional patients than in those without. Delusional patients showed a higher grade of disability in basic and instrumental activities of daily living [34]. Delusions to cluster with hallucinations, agitation/aggression, depression mood, apathy, irritability, aberrant motor activity, sleep disturbances, and eating disorders as assessed by NPI. More severe cognitive impairment and faster rate of cognitive decline are associated with and predictive of hallucinations and delusions in patients with AD. Parkinsonism also is predictive of imminent psychotic symptoms in AD. Psychosis of AD is believed to be the result of dysfunction of frontal lobe circuitry with contributions from neurofibrillary tangles in limbic structures as well as neurochemical abnormalities including the cholinergic deficit and dopaminergic dysfunctions.

Rationale for Interventions and Treatment

AChEIs may reduce or postpone the need for the use psychotropics with worse safety profile. In a community dwelling studies (completed by twenty-four patients with AD) donepezil was proved to significantly reduce delusions, hallucinations and agitation in the majority of subjects. Memantine appears to provide modest benefit for the management of AD psychosis and has a favorable safety profile. In a pooled, retrospective analysis of data from three placebo-controlled trials in moderate to severe AD, memantine was linked to significant reduction in psychosis, agitation, and aggression [35]. In more severe psychotic symptoms, which doesn’t respond to AChEIs or memantine antipsychotic medications are used. Atypical antipsychotic drugs are more effective than placebo, although adverse effects may limit their overall effectiveness [36]. Antipsychotic medications have been associated with a small but significant decrease in caregiver burden [37]. Olanzapine (up to 10 mg daily) has shown benefit in managing delusions and hallucinations, anxiety and agitation in AD patients [38].

Aripiprazole (5 to 10 mg/day) was efficacious and relatively safe for psychosis associated with AD, significantly improving psychotic symptoms, agitation, and NPS as assessed by NPI, BPRS and CMAI scores [39]. Risperidone (mean doses of 1.5 mg daily) treatment was proved efficacious for psychosis of AD in couple of studies [21]. Quetiapine is also commonly used in off-label indications such as NPS of AD. It is believed to have a lower incidence of serious side effects such as extrapyramidal symptoms and tardive dyskinesia when compared with other antipsychotics. Quetiapine can be used in a wide range of doses. Sedating properties are of some use in certain clinical situations in NDD [40]. Results may vary across studies but most show modest benefit for psychosis and agitation with an acceptable side effect profile. Atypical antipsychotic in general appear to have some impact in reducing psychosis as well as agitation in AD with the best evidence base for risperidone. Carefully monitored and relative brief courses of antipsychotics are recommended. Recent studies with pimavanserin addressing psychosis in the course of different types of dementia (including Alzheimer’s disease) are showing promising results [41].

Agitation in the Course of AD

Agitation as a symptom of AD is common (prevalence ranges from 20% to 60% of) and highly disruptive. It is considered the most problematic symptoms among NPS [42]. Agitation commonly clusters with aggressive behavior and tends to co-occur with sleep disorders, delusions, hallucinations, anxiety and dysphoria [12,42]. Table 3 provides the International Psychogeriatric Association (IPA) criteria for the definition of agitation in cognitive impairment [43] (Table 3). Frontal-subcortical and corticocortical networks dysfunction is proposed as the basis for the agitation syndrome [6].
Table 3: International Psychogeriatric Association definition of agitation in cognitive impairment.

| International Psychogeriatric Association Definition of Agitation in Cognitive Impairment |
|---|
| **A** | The patient meets criteria for a cognitive impairment or dementia syndrome (e.g. AD, FTD, DLB, vascular dementia, other dementias, a pre-dementia cognitive impairment syndrome such as mild cognitive impairment or other cognitive disorder). |
| **B** | The patient exhibits at least one of the following behaviors that are associated with observed or inferred evidence of emotional distress (e.g. rapid changes in mood, irritability, outbursts). The behavior has been persistent or frequently recurrent for a minimum of two weeks and represents a change from the patient's usual behavior. |
| (a) Excessive motor activity (examples include pacing, rocking, gesturing, pointing fingers, restlessness, performing repetitive mannerisms). |
| (b) Verbal aggression (e.g. yelling, speaking in an excessively loud voice, using profanity, screaming, shouting). |
| (c) Physical aggression (e.g. grabbing, shoving, pushing, resisting, hitting others, kicking objects or people, scratching, biting, throwing objects, hitting self, slamming doors, tearing things, and destroying property). |
| **C** | Behaviors are severe enough to produce excess disability, which in the clinician’s opinion is beyond that due to the cognitive impairment and including at least one of the following: |
| (a) Significant impairment in interpersonal relationships. |
| (b) Significant impairment in other aspects of social functioning. |
| (c) Significant impairment in ability to perform or participate in daily living activities. |
| **D** | While co-morbid conditions may be present, the agitation is not attributable solely to another psychiatric disorder, suboptimal care, and placement. |

Note: Adapted from: Cummings J, Mintzer J, Brodaty H, Agitation in cognitive disorders: International Psychogeriatric Association provisional consensus clinical and research definition. Int Psychogeriatr 2015; 27:7-17.

Rationale for Interventions and Treatment

The symptom of agitation may be caused by pain or any other discomfort, medical comorbidities, environmental factors or drug effects. The optimal approach to treating agitation in AD thus requires assessing the individual patient's medical circumstances with careful verification of medications. This step should be followed by thorough assessment of symptom severity, cognitive function, and presence of other NPS. Additionally, vulnerability to adverse effects of pharmaceuticals should be carefully assessed and included to the treatment plan. Nonpharmacological, behavioral interventions are crucial in the management of agitation in AD and are considered first-line treatments. Meta-analysis of various psychological approaches found that behavior management therapies, care by specific types of caregiver and residential care, and staff education had the most lasting benefits for the management of agitation in dementia patients. Music therapy and sensory stimulation had positive but short-lived effects [44]. In cases when psychosocial approaches are inadequate, antipsychotics, antidepressants, anticonvulsants, and other classes of drugs are being used off-label.

AChEIs studies provide only modest evidence to support benefit from their use in managing agitation [15,45]; (Howard et al.). In clinical practice AChEIs are not helpful when immediate intervention is required [6]. A study comparing galantamine, (an AChEI) with risperidone, showed that the levels of agitation decreased in both treatment groups, but the improvement was significantly greater in the risperidone group [46]. Memantine provides modest benefit in the treatment of agitation and aggression in dementia and is well tolerated. In pooled, retrospective analysis of data from three placebo-controlled trials in moderate to severe AD, memantine showed significant reduction in agitation, aggression or psychosis [35]. Substantial atypical antipsychotics are used off label despite modest clinical benefits and side-effect burden and risk of mortality. Atypical antipsychotic were more beneficial than placebo and were associated with decreases in caregiver burden, but adverse effects limit their overall effectiveness [36].

There is some evidence to support the use of typical antipsychotics to manage aggression and agitation in the acute clinical setting. Haloperidol is useful in treatment of aggression with agitation (but not general agitation behaviors, such as wandering or verbal agitation) [47]. The use of typical antipsychotics in NDD even in acute situations is considered high risk. Typical antipsychotics are not recommended in non-emergent treatment of agitation in dementia [48]. Experts recommend that risperidone, olanzapine and aripiprazole be used for severe agitation, aggression and psychosis associated with AD where there is risk of harm to the patient and/or others [13]. The potential benefit of all antipsychotics must be weighed against the significant risks, such as cerebrovascular adverse events and mortality. A metaanalysis of four large placebo-controlled clinical trials proved risperidone’s efficacy in the management of agitation and aggression even in severely impaired AD patients [49]. Risperidone may be considered as an option for short term intervention in cases of acute, treatment resistant cases of agitation in AD. In AD patients with psychosis or agitation who had responded to risperidone therapy for 4 - 8 months, discontinuation of risperidone was associated with an increased risk of relapse [50].

If there is no clinically significant response after a 4-week trial of an adequate dose of an antipsychotic drug, the medication should be tapered and withdrawn. In cases of adequate response
an attempt to taper and withdraw the drug should be made within 4 months of initiation, unless the patient experienced a recurrence of symptoms with prior attempts at dose reduction. There is modest evidence to support effectiveness of carbamazepine in targeting agitation and aggression in AD [16]. In practice it’s use is limited by the risk of common side effects such as dizziness, sedation, ataxia, confusion, headaches, nausea, vomiting, diarrhea, blurred vision; and the more rare but significant adverse effects of inappropriate antidiuretic hormone with hyponatremia, cardiac and hepatotoxicity, and increased risk of suicidal behavior and ideation [8]. Patients should also be explicitly informed of warnings for aplastic anemia, agranulocytosis, and rare but sometimes fatal dermatologic adverse reactions. The evidence for valproate in management of cognition in AD is mixed, with a meta-analysis of pooled results concluding valproate is and is associated with unacceptable rates of adverse events, notably sedation and urinary tract infections [13].

Among other pharmaceuticals from this category topiramate has some efficacy: gabapentin and lamotrigine, oxcarbazepine and levetiracetam have been the subject of observational or uncontrolled studies and are considered as low priority agents. Trazodone, a hypnotic and antidepressant (pharmacologically serotonin antagonist and reuptake inhibitor), is used for management of irritability, agitation and aggression in AD. Trazodone has sedating properties with minimal anticholinergic activity. Trazodone has a favorable safety profile if administered in small doses and appears to produce a stabilization of the circadian rhythms in individuals with AD [51]. A few retrospective or observational studies suggest that trazodone may be effective for the treatment of aggression or agitation in AD [52]. The most promising potential pharmacological alternatives to antipsychotics and anti-epileptic agents include citalopram, dextromethorphan/quinidine, and prazosin [53]. Comparator studies indicate sertraline and citalopram are probably as effective as risperidone in treating agitation in dementia. A recent study had shown that dextromethorphan/quinidine significantly improved AD-associated agitation, reduced caregiver burden, and was generally well tolerated [42,54-60].

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