Three keys can help you safely treat dementia’s difficult behavioral and psychological symptoms:

• Differentiate medical from psychiatric causes of patients’ distress.
• Use antipsychotics and other drugs as adjuncts to psychosocial treatments.
• Start low and go slow when titrating dosages.

Although no treatment reverses the pathophysiology of progressive neurodegenerative disorders, managing agitation and other behaviors can alleviate patient suffering and reduce caregiver stress. Based on the evidence and our experience, this article describes a practical approach, including a treatment algorithm and evidence of atypical antipsychotics’ efficacy and side effects in this patient population.

continued
Using antipsychotics in patients with dementia

**DEMENTIA’S BEHAVIORAL SYMPTOMS**
An International Psychogeriatric Association consensus statement grouped dementia’s behavioral and psychological symptoms into two types:
- those usually assessed by interviewing patients and relatives—anxiety, depressed mood, hallucinations, and delusions
- those usually identified by observing patient behavior—aggression, screaming, restlessness, agitation, wandering, culturally inappropriate behaviors, sexual disinhibition, hoarding, cursing, and shadowing.

These behaviors in community-living patients are distressing to family members and increase the risk for caregiver burnout—the most common reason for placing older patients in long-term care. In the nursing home, dementia’s symptoms reduce patients’ quality of life; interfere with feeding, bathing, and dressing; and—when violent—may endanger staff and other patients.

**RULE OUT A MEDICAL CAUSE**
Differential diagnosis. Behavioral symptoms in dementia tend to be unpredictable, which makes diagnosis and treatment challenging. The first step is to determine if a medical or psychiatric condition might account for the behavior. For instance:

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**Algorithm: Treating behavioral symptoms in patients with dementia**

- **Patient exhibits psychotic or agitated behavior**
  - Yes: Cause by medical problem (pain, delirium) or psychiatric disorder (bipolar mania, schizophrenia)?
    - Yes: Begin psychosocial interventions
    - No: Begin psychosocial interventions
  - No: Treat and monitor behavior
  - Continue

- **Adequate symptom reduction?**
  - Yes: Consider adding atypical antipsychotic:
    - risperidone, 0.5 to 1.5 mg/d
    - olanzapine, 5 to 10 mg/d
    - quetiapine, 25 to 350 mg/d
    - Titrate slowly, and monitor for side effects
  - No: Continue

- **Caused by medical problem (pain, delirium) or psychiatric disorder (bipolar mania, schizophrenia)?**
  - Yes: Treat and monitor behavior
  - No: Begin psychosocial interventions

- **Valproate, 125 mg bid, or carbamazepine, 100 mg bid, and titrate to effect, and/or a cholinesterase inhibitor:**
  - donepezil, 5 mg qd; increase to 10 mg qd after 4 to 6 weeks
  - rivastigmine, 1.5 mg bid; increase to 9 to 12 mg/d in divided doses
  - galantamine, 4 mg bid; increase to 8 mg bid after 1 month
A patient with dementia may be agitated because of a distended bladder or arthritis but unable to communicate his or her pain in words.

In mild dementia, a pre-existing psychiatric disorder such as schizophrenia might be causing a patient’s hallucinations or delusions.

Pacing and restlessness may be drug side effects and might be controlled by reducing dosages or switching to less-activating agents.

Delirium is also a risk for older patients—especially those with degenerative neurologic disorders. Common triggers in older patients include acute illness such as a urinary tract infection or pneumonia, alcohol or benzodiazepine withdrawal, anticholinergic agents, medication changes, and dehydration.

Delirium is characterized by acute onset and fluctuating neuropsychiatric symptoms, including disturbed consciousness and changes in attention and cognition. Taking a careful history to learn the course of treatment and the patient’s baseline cognitive function can help you differentiate dementia from delirium. Family members, physicians, and nursing staff are valuable sources of this information.

**USE ANTIPSYCHOTICS AS ADJUNCTS**

**Psychosocial interventions.** After medical causes have been ruled out, consensus guidelines recommend psychosocial interventions as first-line treatment of dementia’s behavioral symptoms (Algorithm). Suggested interventions for patients and caregivers are listed in Table 1.

**Antipsychotics.** For patients who respond inadequately to psychosocial measures, the next step is to add an atypical antipsychotic. Because of side effects, conventional antipsychotics are not recommended for patients with dementia.

When prescribing atypicals, remember that older adults:

- are more sensitive to side effects than younger adults
- require lower starting and target dosages
- exhibit heterogeneity of response.

Older patients’ medical status can range from “fit” to “frail,” which influences individual response to medications. Generally, age-related changes in the way their bodies metabolize drugs account for older patients’ increased sensitivity to drug side effects (Box, page 61).

Atypical antipsychotics and dosages that have been shown benefit for managing behavioral symptoms in older patients with dementia include:

- risperidone, 0.5 to 1.5 mg/d
- olanzapine, 5 to 10 mg/d
- quetiapine, 25 to 350 mg/d

Start with low dosages, and titrate slowly. Increase once or twice a week until the lowest effective dosage is reached.

**Augmenting agents.** If antipsychotic monotherapy fails to achieve an adequate response or if side effects limit dosing, adjunctive agents may be added with caution. Augmenting agents that have shown benefit in some patients with dementia include:

- mood stabilizers such as divalproex or carbamazepine
- cholinesterase inhibitors, such as donepezil, rivastigmine, or galantamine

Start divalproex at 125 mg bid or carbamazepine at 100 mg bid and titrate to effect. Concomitant carbamazepine will decrease blood levels of risperidone, olanzapine, and quetiapine because of hepatic enzyme induction.

Start donepezil at 5 mg once daily and increase after 4 to 6 weeks to 10 mg qd. When using rivastigmine, start with 1.5 mg bid and titrate to 9 to 12 mg/d in divided doses. Start galantamine at 4 mg bid and increase after 1 month to 8 mg bid.
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Table 1

Suggested psychosocial interventions for older patients with dementia

Communicate clearly
• Validate patients’ statements, then redirect any that may be inappropriate

Minimize the impact of sensory deficits
• Decrease risk of disorientation by providing needed corrective eyeglasses and hearing aids

Modify environment when necessary
• Install adequate daytime lighting to improve sleep patterns in patients with disturbed sleep/wake cycles

Encourage consistent daily routines
• Schedule times for meals and for arising in the morning and going to bed at night to minimize disruptions and distress

Optimize social/physical stimulation
• Display photos and names of family and friends in the patient’s living area
• Help the patient do daily stretching exercises to music

Encourage caregiver to:
• Make use of support groups and caregiver resources
• Consult with attending psychiatrist or physician when psychosocial interventions do not adequately manage a patient’s problem behaviors

ANTIPSYCHOTIC SIDE EFFECTS

Atypical antipsychotics are more effective than conventional agents in treating negative symptoms and are associated with lower rates of extrapyramidal symptoms (EPS) and tardive dyskinesia (TD). Tardive dyskinesia. All antipsychotics can cause TD, although the risk is about 10 times greater with conventional than atypicals. With conventional, the annual cumulative TD incidence for young adults is 4 to 5%, and rates are much higher for middle-aged and older adults receiving chronic therapy:

• 29% after 1 year
• 50% after 2 years
• 63% after 3 years.31

In older patients, use atypical rather than conventional antipsychotics to minimize TD risk. Observe carefully; if TD symptoms occur, cautiously withdraw the antipsychotic and consider trying another agent.

Other risks. Atypical antipsychotics may cause sedation, orthostatic hypotension (with an increased risk for falls), increased serum prolactin, and weight gain (Table 2).

Weight gain from atypical antipsychotics has been associated with adverse effects on glucose metabolism and increased risk for type 2 diabetes. Some might argue that weight gain associated with olanzapine and other atypicals might benefit low-weight older patients. The frail elderly need to increase muscle mass, however, and the atypicals are associated with increases in fat mass.

Increased serum prolactin with risperidone theoretically could lead to loss of bone density, but evidence of this effect in older patients does not exist.

START LOW, GO SLOW

Clozapine may help control treatment-resistant psychosis in patients with schizophrenia and manage patients with severe TD. However, clozapine’s increased risk of agranulocytosis, neurologic side effects (seizures, sedation, confusion), and anticholinergic effects limit its use in older patients, particularly those with neurodegenerative disorders (Table 2).

Dosing. In rare cases when using clozapine in older patients, start with 6.25 to 12.5 mg/d. Increase by 6.25 to 12.5 mg once or twice a week to 50 to 100 mg/d.

Risperidone has been used to treat agitation in older patients with dementia in two small studies:

In a 9-week, open-label trial, 15 patients (mean age 78) with dementia were given risperidone...
done, 0.5 to 3 mg/d. Agitation improved significantly, as measured by the Cohen-Mansfield Agitation Inventory (CMAI)—a 29-item questionnaire completed by caregivers. CMAI scores at study’s end averaged 49.5, compared with 70.5 at baseline.27

A 12-week, placebo-controlled, double-blind study examined risperidone—0.5, 1, or 2 mg/d—in 625 institutionalized patients (mean age 83) with dementia and agitation. Ninety-six patients had Functional Assessment Staging Rating Scale scores of 6A, indicating moderate to severe dementia. In patients receiving risperidone, these behavioral measures were significantly reduced:

- Behavior Pathology in Alzheimer’s Disease Rating Scale (BEHAVE-AD) total scores, which measure behavior severity
- BEHAVE-AD psychosis subscale scores
- BEHAVE-AD aggressiveness scores
- CMAI verbal and aggression scores

Adverse effects were reported at ≥ 82% for all three risperidone dosages and 85% for placebo. Side effects including somnolence, EPS, and peripheral edema were dose-related.12

Another trial compared risperidone or haloperidol, 0.5 to 4 mg/d, with placebo in treating 344 patients with behavioral symptoms of dementia. After 12 weeks of risperidone, mean dosage 1.1 mg/d:

- mean total BEHAVE-AD score decreased by 53%, compared with 37% in the placebo group
- CMAI score decreased by 32%, compared with 18% in the placebo group.

EPS were more severe with haloperidol than with risperidone or placebo.28

Risk of stroke. A small but significantly increased incidence of stroke and stroke-like events was recently reported in older patients with dementia when treated with risperidone. These events occurred in double-blind, placebo-controlled trials in patients (mean age 82) with Alzheimer’s, vascular, and mixed dementias.

Most patients who experienced cerebrovascular events had one or more stroke risk factors, including diabetes, hypertension, atrial fibrillation, heart arrhythmia, atherosclerosis, or heart failure. They did not show a pattern of reduced blood pressure or orthostatic changes.12,29

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**Age-related changes affect how older patients metabolize psychotropics**

**Pharmacokinetic changes** can influence concentrations of drugs in tissue compartments over time. Drug absorption declines with normal aging, but a clinically significant decrease in total absorption of psychotropics appears not to occur.13

In the liver, lipid-soluble psychotropics are metabolized into pharmacologically active or inactive metabolites. Some metabolic pathways, such as demethylation, may be influenced by age, leading to increased plasma concentrations of drugs and their metabolites.14,15 However, hydroxylation tends not to be affected by age.16

The ratio of body fat to water increases with aging,17 increasing the volume of distribution for lipid-soluble psychotropics. An age-related decrease in glomerular filtration accounts in part for increased accumulation of hydrophilic metabolites in some older patients.17,18

**Pharmacodynamic changes** with aging occur in neurotransmitter systems within cellular processing, such as at receptor or reuptake levels.19 These changes may exaggerate drug-drug interactions or affect complex neurotransmitter interactions.

The number of neurons in nigrostriatal pathways declines with age. Decreases are also seen in tyrosine hydroxylase activity, presynaptic dopamine D2 receptors, and dopamine levels—which may be particularly relevant to a discussion of antipsychotic medications.20

The net effect of these changes is the need to prescribe lower-than-usual starting and target dosages of many medications, including antipsychotics.

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Dosing. For older patients with dementia and psychosis, start risperidone at 0.25 to 0.5 mg/d and increase by no more than 0.25 to 0.5 mg once or twice per week. Do not exceed 3 mg/d. For agitation, a 1998 Expert Consensus Guideline Series panel recommended starting risperidone at 0.25 to 0.5 mg/d and increasing to an average of 0.5 to 1.5 mg/d.

Olanzapine. Two double-blind, placebo-controlled studies have examined olanzapine in treating agitation associated with dementia.

Saterlee et al compared olanzapine, mean 2.4 mg/d, with placebo in outpatients (mean age 79) with Alzheimer’s disease and psychosis. No significant differences were noted in hepatic transaminase levels, leukopenia, EPS, or orthostatic changes.

In a later study, nursing home patients (mean age 83) with Alzheimer’s disease, psychosis, and agitation were randomly assigned to receive olanzapine—5, 10, or 15 mg/d—or placebo. After 6 weeks, patients receiving olanzapine, 5 or 10 mg/d, showed significant improvement in Neuropsychiatric Inventory (NPI) total core scores. Olanzapine, 15 mg/d, was not significantly more effective than placebo.

Adverse events such as somnolence and abnormal gait occurred more often with olanzapine than placebo. The somnolence rate with olanzapine was 14% for 5 mg/d and 13% for 10 mg/d, compared with 3% for placebo. For abnormal gait, the rate with olanzapine was 11% for 5 mg/d and 7% for 10 mg/d, compared with 1% for placebo.

Dosing. Start olanzapine at 2.5 mg/d, and increase after 1 to 3 days to 5 mg/d. If symptoms are not adequately controlled, titrate by 2.5-mg increments to 10 mg/d.

Quetiapine. One open-label study examined using quetiapine in older patients with psychotic disor-
Aripiprazole. As with ziprasidone, little data exist to guide the use of aripiprazole in older patients. In a randomized preliminary trial, 32 192 noninstitutionalized patients with Alzheimer’s disease and psychosis were treated for 10 weeks with aripiprazole, mean 10 mg/d, or placebo.

At 8 and 10 weeks, BPRS psychosis subscale scores improved significantly in patients taking aripiprazole, compared with placebo. EPS and akathisia improved, and somnolence was the most common side effect. Although this study enrolled noninstitutionalized patients with dementia, the results suggest that aripiprazole may help treat long-term care residents with neurodegenerative disorders and behavioral disturbances.

References


Atypical antipsychotics can help manage psychosis and agitation in patients with dementia. Keys to safe prescribing are to differentiate medical from psychiatric causes of behavioral symptoms, use atypicals as adjuncts to psychosocial treatments, and start low and go slow when titrating dosages.

Ziprasidone and aripiprazole appear safe and effective in older patients with dementia.

**Dosing.** Start quetiapine at 25 mg once at bedtime or bid; increase in 25-mg increments until the lowest effective dosage is achieved.

**Ziprasidone.** Little data exist on using ziprasidone in long-term care. In one recent study, 31 ziprasidone (mean 100 mg/d) was given to 62 patients ages 64 to 92 with medical illnesses plus major depression, bipolar disorder, schizoaffective disorder, Alzheimer’s disease, or multi-infarct dementia. A retrospective chart review of 10 patients showed decreased agitation, as mean NPI scores declined from 76 to 33.

Sedation was the most common side effect. QTc findings, postural hypotension, and syncope rates did not change. Despite its limitations, this study suggests that ziprasidone is safe and effective in treating psychosis associated with dementia or other disorders.
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Related resources


**DISCLOSURE**

Dr. Kasckow receives research support from, is a consultant to, or is a speaker for Eli Lilly & Co., Forest Laboratories, Solvay Pharmaceuticals, AstraZeneca Pharmaceuticals, Organon, Janssen Pharmaceutica, and Pfizer Inc.

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**DRUG BRAND NAMES**

- Aripiprazole • Abilify
- Carbamazepine • Tegeotol
- Clozapine • Clozaril
- Donepezil • Aricept
- Galantamine • Reminyl
- Haloperidol • Haldol
- Ziprasidone • Geodon
- Olanzapine • Zyprexa
- Quetiapine • Serquel
- Risperidone • Risperdal
- Valproate • Depakote

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Related resources


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