Agitation Associated with Dementia (AAD) Management without Antipsychotics
A practical rough and tumble guide for family and professional care partners

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One Day Mindfulness Millionaire: A light-hearted primer for the uninitiated.
Abhilash Desai and Faith Galliano Desai
Book available at store.bookbaby.com

7-15-24

Each of us literally chooses, by his way of attending to things, what sort of a universe he shall appear to himself to inhabit.

- William James
### Table of Contents

<table>
<thead>
<tr>
<th>Page</th>
<th>Topic</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Title</td>
</tr>
<tr>
<td>2</td>
<td>Table of Contents</td>
</tr>
<tr>
<td>3</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td>Introduction</td>
</tr>
<tr>
<td>6</td>
<td>Basics of AAD management for Primary Care Providers (PCPs)</td>
</tr>
<tr>
<td>7</td>
<td>–</td>
</tr>
<tr>
<td>8</td>
<td>Collaborative de-escalation of behavioral emergencies in persons with dementia</td>
</tr>
<tr>
<td>9</td>
<td>14 common reversible medical causes of agitation in persons with dementia</td>
</tr>
<tr>
<td>10</td>
<td>MMSE checklist for biopsychosocial factors causing / contributing to AAD</td>
</tr>
<tr>
<td>11</td>
<td>CALM checklist</td>
</tr>
<tr>
<td>12</td>
<td>–</td>
</tr>
<tr>
<td>13</td>
<td>SHIELD plan for AAD prevention and treatment</td>
</tr>
<tr>
<td>14</td>
<td>A-Z Psychosocial Spiritual “Therapies”</td>
</tr>
<tr>
<td>15</td>
<td>Sundowning</td>
</tr>
<tr>
<td>16</td>
<td>Trauma-Informed Care for persons with dementia</td>
</tr>
<tr>
<td>17</td>
<td>Routine psychosocial spiritual / non-drug interventions for AAD</td>
</tr>
<tr>
<td>18</td>
<td>Relationship building communication techniques</td>
</tr>
<tr>
<td>19</td>
<td>–</td>
</tr>
<tr>
<td>20</td>
<td>STEPS – support system training education praise support</td>
</tr>
<tr>
<td>21</td>
<td>Caregiver Dementia Questionnaire – 9</td>
</tr>
<tr>
<td>22</td>
<td>CDQ-9</td>
</tr>
<tr>
<td>23</td>
<td>Don’t TRAC – Test, Reason, Argue, Correct</td>
</tr>
<tr>
<td>24</td>
<td>Don’t BITE – Blame, Insist on reality, Take things personally, Expect unrealistically</td>
</tr>
<tr>
<td>25</td>
<td>T-A-DA method to manage agitation</td>
</tr>
<tr>
<td>26</td>
<td>Mission Impossible</td>
</tr>
<tr>
<td>27</td>
<td>Mindful Improv – Caregiver handout</td>
</tr>
<tr>
<td>28</td>
<td>Bright light therapy</td>
</tr>
<tr>
<td>29</td>
<td>–</td>
</tr>
<tr>
<td>30</td>
<td>Feeding the squirrels therapy</td>
</tr>
<tr>
<td>31</td>
<td>Doll therapy (controversial)</td>
</tr>
<tr>
<td>32</td>
<td>Resistance to care and AAD</td>
</tr>
<tr>
<td>33</td>
<td>ABCD of spiritual assessment</td>
</tr>
<tr>
<td>34</td>
<td>Dignity conserving care</td>
</tr>
<tr>
<td>35</td>
<td>SPPEICE</td>
</tr>
<tr>
<td>36</td>
<td>–</td>
</tr>
<tr>
<td>37</td>
<td>SPPEICE case example Mr. MT</td>
</tr>
<tr>
<td>38</td>
<td>–</td>
</tr>
<tr>
<td>39</td>
<td>SPPEICE case example Mrs. B</td>
</tr>
</tbody>
</table>
SPPEICE case example Ms. Grace

Sexually “Inappropriate” Behaviors in Persons with Dementia Psychosocial Environmental Approaches

Ten Best Practices: Reducing Use of Antipsychotic Medications in individuals with Dementia

Alternatives to antipsychotics for management of AAD

Citalopram for AAD

Stages of Alzheimer’s Disease Dementia

4 Ms of Age-Friendly Health Systems and Long-Term Care Institutions (includes nursing homes and assisted living facilities)

AAD in hospitals: Tomorrow will bring good news: Dementia Friendly Hospital Practices

AAD in hospitals: 10 key interventions

Delirium Care Score

Personalized Multicomponent Delirium Prevention Interventions

Resources

Resources for Dementia Training

Bibliotherapy

Arts on Prescription for dementia care: Resources

Dr. Desai books and podcasts

Dementia Physician Experts in Idaho

Supporting caregivers – key to preventing / reducing AAD

The Healing Power of Mindful Care-Partnering

AAD Medical Poems - Sheer Terror (Wandering in dementia)

AAD Medical poems - Unthinkable future

AAD Medical Poems - Anything but dementia

AAD Medical Poems - My world

AAD Medical Poems - Long buried sorrow (Trauma and AAD)

AAD Medical Poems - Restraints

AAD Medical Poems - Thanx

AAD Medical Poems – Delirium on Dementia
<table>
<thead>
<tr>
<th>Page</th>
<th>Title</th>
</tr>
</thead>
<tbody>
<tr>
<td>80</td>
<td>AAD Medical Poems – An Angel (nurse’s aide, nursing home and AAD)</td>
</tr>
<tr>
<td>81</td>
<td>AAD Medical Poems - A blessing for caregivers</td>
</tr>
<tr>
<td>82</td>
<td>When Is Enough, Enough? Personal Reflections on the Threats and Ills Facing Persons Having Dementia</td>
</tr>
<tr>
<td>83</td>
<td>–</td>
</tr>
<tr>
<td>84</td>
<td>–</td>
</tr>
<tr>
<td>85</td>
<td>Holistic Dementia Care Medicine – 11 Domains</td>
</tr>
<tr>
<td>86</td>
<td>New Research</td>
</tr>
</tbody>
</table>
Introduction

Individuals with dementia experience agitation frequently and it is an expression of emotional distress caused by one or more factors that need to be looked for and addressed (e.g., pain, boredom). Agitation manifests as verbal aggression (e.g., yelling), physical aggression (e.g., hitting), physical hyperactivity (e.g., exit seeking, pacing). It is often accompanied by anxiety, depression and occasionally psychotic symptoms (delusions, hallucinations).

Agitation in persons with dementia is often treated with antipsychotics.

Antipsychotics are high risk medications that are often used inappropriately / unnecessarily and often cause more harm than good.

Risks of antipsychotics in persons with dementia include but not limited to:
- Black box warnings (strokes and death)
- Dysphagia
- Aspiration pneumonia
- Falls and injury (e.g., hip fracture, traumatic brain injury)
- Day time sedation
- Tiredness and fatigue
- Increased visits to emergency department
- Akathisia (motor restlessness)
- Insomnia (with certain antipsychotics like aripiprazole)
- Dysphoria, irritability, apathy

Our hope with this eBook is to provide education and guidance on all the things we can do to reduce agitation and improve experience of positive emotions in persons with dementia so that they can live the best life possible.
Basics of Agitation Associated with Dementia (AAD) Management for PCPs

1. STAR approach: Safety, Team input, Action plan, Response to treatment
   a. Safety: assess if the person is in delirium or a similar condition posing imminent danger to self and or others. If so, they may need ASAP assessment or emergency department assessment.
   b. Team input: just input from one family member is not enough, especially if the person with dementia appears calm during the visit.
   c. Action plan: Biopsychosocial spiritual care plan based on the strengths and interests of the person with dementia.
   d. Response to treatment using objective measures (behavioral tracking data) and modification of action plan based on response or lack of response.

2. MMSE (acronym – not the cognitive test 😊)
   a. Medical causes – pain, constipation, dehydration, skin infections, dental problems, metabolic-endocrine-nutritional problems.
      i. OT referral
      ii. PT referral
      iii. Community health worker
      iv. Memory center referral
   b. Medication induced – de-prescribing
   c. Social factors – family / staff directed interventions (e.g., powerful tools for caregivers, DICE training), adult day programs.
   d. Environmental – Occupational therapy, environmental modifications

3. Biopsychosocial spiritual wellness care plan
   a. Biological: chamomile extract, melatonin, citalopram
   b. Psychological: engagement in fun activities, communication
   c. Social: family support and intervention
   d. Spiritual: purpose and meaning in daily life, spiritual rituals

4. STEPS Support system training education praise support
   a. Community resources education
   b. Communication
   c. DICE Approach Online Training Program: https://diceapproach.com
   d. TADA approach
   e. Don’t TRAC – Test, Reason, Argue, Correct
   f. Don’t BITE – Blame, Insist on reality, Take things personally, Expect unrealistically
   g. Individual counseling
   h. Support groups
   i. Respite
   j. Bibliotherapy
Basics continued: Five-Step Approach

Step one: Medical assessment
Thorough initial assessment and workup for medical causes of psychiatric problems. Example: pain, dehydration, constipation, infection, delirium. Consider CBC, CMP, TSH, B12, D, Mag levels and for chronic insomnia, ferritin levels (iron deficiency can cause RLS and restless sleep). Address hearing and vision impairment.

Step two: Rational Deprescribing
Review medications and look for medication induced psychiatric problems. This includes medication adverse effects, toxic levels (may need blood levels), and medication withdrawal. Consider rational deprescribing of unnecessary medications, medications not in keeping with goals of care, and medications on AGS 2023 Beers list of medications potentially inappropriate in older adults. Consider checking for total anticholinergic burden of medications using the www.acbcalc.com and reduce anticholinergic burden. Consider pharmacy, geriatrician or geriatric psychiatry eConsult for review of meds and suggestions for deprescribing.

Step three: Psychosocial environmental interventions
Identify social and environmental factors causing or contributing to agitation. Examples include lack of structure, boredom, loneliness, lack of respect and inclusion in decision making. The nursing and social work team can take leadership role in addressing this. Family / staff training, education, praise and support are included here. Create an individualized, strengths-based psychosocial spiritual care plan. Create an IPAS: individualized pleasant activity schedule. Examples: continuous fun and meaningful activities programming, lavender lotion several times a day, bright light therapy, exercise therapy, soothing bedtime rituals.

Step four: Benign psychopharmacological interventions
Consider relatively benign medications first psychopharmacological interventions are needed.
Anxiety: chamomile extract 500mg three times daily
Insomnia: melatonin 3mg at bedtime daily, chamomile 500mg-1500mg at bedtime daily

Step five: Traditional psychopharmacological interventions
This step is fraught with ethical challenges, especially if antipsychotics, benzodiazepines and trazodone (all high risk and often cause more harm than good) are being considered. Comprehensive psychiatric assessment and traditional psychiatric medications may help in certain situations (e.g., severe depression, severe persistent aggression, suicidal ideas, distressing psychotic symptoms). This may be done by primary care providers with psychiatrist available via telemedicine for guidance.

P.S. Depending on the context, all five steps may need to be done simultaneously with different team members tackling different steps. Family members can have PCP help with these steps or seek help via telemedicine from one of the Dementia Physician Experts (page 53)
Collaborative De-escalation for Behavioral Emergencies in persons with Dementia

Guidance for first responders, family and professional caregivers.

REACH acronym – Florida State University School of Medicine Department of Geriatrics: https://reach.med.fsu.edu/be-prepared-for-emergent-situations-involving-persons-with-dementia/

R – Risk perception (assess risk of the person with dementia injuring themselves or harming others)
E – Establish trust (includes keeping personal energy calm and friendly, do not challenge what they believe is true)
A – Assess immediate needs (includes providing information and handouts on community resources to family caregivers)
C – Calm environment (includes reducing distractions or stressors, turning off TV, interacting in as calm and quiet environment as is feasible)
H – Healthy partnership (collaboration with the person having dementia and their family members; this includes getting the person with dementia to an Emergency Department for assessment)

The same website has a 4-minute video for first responders.

Examples of common behavioral emergencies: suicidal behaviors, violent behaviors, physical aggression towards family members or others, severe agitation combined with high risk of falls, involvement of guns.

Five most common causes of behavioral emergencies:
- Delirium
- Psychotic symptoms
- Pain
- Medication adverse effects
- Alcohol and or cannabis intake

Namaste
14 Common Reversible Medical Causes of Agitation / Acute Psychosis in Persons with Dementia

1. Dehydration
2. Hyperglycemia / hypoglycemia
3. Infection (e.g., urinary tract infection, pneumonia, dental infection, cellulitis)
4. Medication adverse effects and toxicity (may need blood levels of certain medications) (common cause I have seen in my practice: medications used for Parkinson’s disease and restless leg syndrome [e.g., ropinirole, pramipexole, levodopa-carbidopa])
5. Alcohol, opioids, and cannabis withdrawal
6. Sedative hypnotic withdrawal (e.g., a benzodiazepine such as lorazepam)
7. Acute traumatic brain injury (typically after a fall – witnessed or unwitnessed)
8. Electrolyte imbalance (e.g., low sodium, high calcium)
9. Acute endocrine disorders (e.g., thyroid problems – took more than prescribed thyroid medication or not taking thyroid medication)
10. Acute cerebrovascular event / stroke
11. Substance use / intoxication (e.g., marijuana, alcohol)
12. Acute hypoxia (e.g., due to an acute respiratory condition)
13. Epilepsy / seizures
14. Multiple causes (more than one cause)
Checklist to Identify Biopsychosocial Environmental Factors causing “Behavioral” Problems

Checklists are an important tool to manage complexity. I use the acronym MMSE

M = Medical conditions  
M = Medications induced (includes substances such as caffeine and illegal drugs)  
S = Social factors  
E = Environmental factors

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<thead>
<tr>
<th>Questions</th>
<th>Yes</th>
<th>No</th>
<th>Don’t know</th>
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<tr>
<td>Was there any abrupt change in behaviors?</td>
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<td>Are there any active medical problems going on?</td>
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<td>Is the person experiencing pain?</td>
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<td>Is the person having constipation?</td>
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<td>Is the person drinking adequate fluids?</td>
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<td>Is the person eating too little or excessively?</td>
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<td>Is the person sleeping too much or too little?</td>
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<td>Is the person having any skin problems?</td>
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<td>Are there any sensory sensitivities, hearing, vision problems?</td>
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<td>Is the person taking medications are prescribed?</td>
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<tr>
<td>Is the person having any adverse effects from medications*?</td>
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<td>Is the person taking substances that alter mood?</td>
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<td>Is support team creating a positive validating environment?</td>
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<td>Has there been a significant change in support system?</td>
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<td>Are there physical environmental factors causing stress?</td>
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<tr>
<td>Does the care team know what brings the person joy**?</td>
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*headache, stomach discomfort, drowsiness, insomnia, vivid dreams, nausea, vomiting, loss of appetite, increased appetite, tremors, agitation, mood instability, tearfulness, anxiety, dry mouth, constipation (onset correlates with when medication was started or dose increased). Adverse effects from medications includes discontinuation and withdrawal symptoms and signs.

**aka what matters to the person

Feel free to modify these questions or add questions that reflect the common MMSE factors for that particular person. I use this to help individuals with advanced dementia, ASD, and IDD.

Namaste
CALM-Checklist
Controlling Agitation with Less Medications (CALM) – Checklist for Older Adults with Dementia

To be completed by family members and or staff (nurse, social worker, certified nursing assistant) in collaboration with primary care providers prior to Psychiatry consult for management of agitation in persons with dementia.

Check if done  Intervention

______ Mr./Ms. __ is adequately hydrated

______ Mr./Ms. __ is having regular bowel movements

______ Mr./Ms. __ is not in pain / pain optimally controlled

______ Mr./Ms. __ is sleeping well regularly

______ Mr./Ms. __ has exposure to sunlight or BLT* at least 20 minutes a day

______ Mr./Ms. __ has some physical activity every 1-2 hours

______ Team has created an Individualized Pleasant Activity Schedule (IPAS)**

______ Mr./Ms. __ is engaged in IPAS throughout the day

______ Mr./Ms. __’s MedRev*** done

______ Mr./Ms. __ support systems have been encouraged to be with resident

______ Mr./Ms. __ has access to his preferred food and activity

______ Sensory deficits (e.g., hearing, vision) have been addressed.

*BLT: Bright Light Therapy using a bright light therapy box (https://cet.org/product/light-therapy-lamp/)

**IPAS: It is a list of activities that the patient in the past has found pleasant/meaningful/fun (created using information obtained from interviews with patient, his family and friends, and observation by medical team) and includes spiritual activities and interaction with a chaplain.

***MedRev: Medication Review. Best done by PCP in collaboration with facility’s consultant pharmacist. There are two goals of MedRev: Rational Deprescribing and Identifying medications that may be causing psychiatric symptoms and distress. Rational deprescribing involves at least three of the following: anticholinergic load (if any) has been minimized; medications on Beers’
list have been minimized / discontinued; unnecessary medications and medications where risk is more than benefits have been discontinued through the process of *Rational Deprescribing***.

Note: Primary interventions for prevention and treatment of agitation involve nursing and psychosocial-environmental approaches and rational deprescribing.

The SHIELD Plan for prevention and treatment of agitation, exit seeking in persons with Dementia

SHIELD is an acronym for the following predisposing and precipitating factors of agitation and exit seeking behaviors seen in individuals with advanced dementia, especially those living in nursing homes and assisted living communities:

<table>
<thead>
<tr>
<th>Anticipate and Address</th>
<th>Interventions (examples)*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>S</strong> Stress (elevated stress hormones, elevated BP, HR)</td>
<td>Soothing music, Lavender lotion, cold compresses to eyes and face, address sources of stress (e.g., pain, constipation, dehydration) Encourage 1500 cc fluid daily or more to prevent dehydration</td>
</tr>
<tr>
<td><strong>H</strong> Hearing and vision deficits and other sensory sensitivities may cause frustration and agitation that may lead to increased exit seeking behaviors</td>
<td>Address these deficits or make accommodations</td>
</tr>
<tr>
<td><strong>I</strong> Insomnia causing agitation and exit-seeking behaviors</td>
<td>Discontinue medications and care practices that cause / worsen insomnia, sunlight – bright light therapy, avoid daytime naps, soothing bedtime rituals, melatonin</td>
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<tr>
<td><strong>E</strong> Exercise insufficiency: lack of adequate physical activity causing restlessness and exit seeking behaviors</td>
<td>Physical and occupational therapy, walking program</td>
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<tr>
<td><strong>L</strong> Loneliness and boredom causing exit seeking behaviors</td>
<td>Increase interaction with family, friends, pets, engage them in meaningful and strengths-based activities throughout the day</td>
</tr>
<tr>
<td><strong>D</strong> Drug-induced agitation manifesting as exit seeking</td>
<td>Discontinue medications that are unnecessary, that care highly anticholinergic and drugs that are causing / can cause agitation; involve PCP, pharmacist and psychiatrist in reviewing medications</td>
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</tbody>
</table>

*Need to personalize it (e.g., if they prefer a different aroma than lavender, switch to it)
A-Z Psychosocial Spiritual “Therapies” for prevention and treatment of AAD

Aromatherapy
Art therapy
Animal – Assisted therapy
Bright light therapy
Cognitive stimulation therapy [https://www.slu.edu/medicine/internal-medicine/geriatric-medicine/aging-successfully/cognitive-stimulation-therapy.php](https://www.slu.edu/medicine/internal-medicine/geriatric-medicine/aging-successfully/cognitive-stimulation-therapy.php)
Clown therapy
Cooking therapy
Doll therapy
Dignity therapy as described by Canadian psychiatrist Dr. Harvey Chochinov
Exercise therapy, includes walking therapy
Feeding the squirrels therapy, includes feeding fish
Fun activities therapy
Garden / horticulture / plant therapy
Group activities therapy
Gratitude based therapy
Humor-based therapy
Humanitude therapy [https://humanitudesingapore.com/#](https://humanitudesingapore.com/#)
Individualized pleasant activities schedule therapy
IN2L (It’s Never 2 Late) now called Life Loop [https://lifeloop.com](https://lifeloop.com)
Laughter therapy
Massage therapy, includes hand massage, scalp massage
Music therapy, includes personalized music, singing, dancing, spiritual music
Nature therapy, includes farm therapy, river and ocean therapy, sky-time therapy, stars therapy
One-to-one therapy, companionship and fun + meaningful activities done together
Pet therapy
Relaxation exercises therapy
Reminiscence therapy
Robotic pets’ therapy
Simulated presence therapy (basically audio video recordings of family members, pets)
Spiritual therapy, practices, rituals
Sun therapy
Tai Chi therapy
Tech therapy – IN2L, virtual reality, guided meditation / relaxation with high quality free apps (e.g., Healthy Minds Program, UCLA Mindful, Cleveland Clinic Wellness CCW Mindful Moments)
Timeslips therapy, creative engagement therapy [https://www.timeslips.org](https://www.timeslips.org)
Touch therapy (includes hugs – 20 seconds or more)
Validation therapy, [https://vfvalidation.org](https://vfvalidation.org)
Virtual reality (VR) therapy
Weighted blanket therapy
Yoga therapy, includes chair Yoga
Zen garden therapy
**Sundowning / Sundowner syndrome in Persons with Dementia**

Symptoms: Increased anxiety and restlessness in afternoons and evenings.

Management: Address the potential cause(s) – typically more than one cause. Create a “Do’s” and “Don’t” list that is individualized, strengths based. No meds have been found to be useful.

<table>
<thead>
<tr>
<th>Potential cause</th>
<th>Intervention</th>
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<tbody>
<tr>
<td>Brain fatigue</td>
<td>Short afternoon nap</td>
</tr>
<tr>
<td>Elevated stress hormone causing severe anxiety</td>
<td>Relaxing / Calming activities</td>
</tr>
<tr>
<td></td>
<td>- Lavender lotion massage three times in afternoon</td>
</tr>
<tr>
<td></td>
<td>- Back rub / massage</td>
</tr>
<tr>
<td></td>
<td>- Soothing music</td>
</tr>
<tr>
<td></td>
<td>- Soothing multisensory room</td>
</tr>
<tr>
<td></td>
<td>- Going through photo album</td>
</tr>
<tr>
<td></td>
<td>- Favorite laughter programs on TV</td>
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<td>- Watching recording of children / grandchildren</td>
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<td>- Meaningful calming activities (e.g., folding towels)</td>
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<td></td>
<td>- Pets / Robotic pets</td>
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<tr>
<td>Overstimulation (excessive noise, too many people, too much movements of people)</td>
<td>Taking the person to a quieter place with less stimulation, reduce visits by family friends in the afternoon (can visit in the morning)</td>
</tr>
<tr>
<td>Under stimulation / boredom</td>
<td>Walking / other exercise program in the morning and individualized meaningful and pleasant activity schedule</td>
</tr>
<tr>
<td>Insomnia</td>
<td>Avoid excessive daytime napping, remove meds causing insomnia, soothing nighttime rituals (e.g., bedtime prayers) and other interventions to address insomnia effectively</td>
</tr>
<tr>
<td>Low lighting</td>
<td>Exposure to sunlight, bright light therapy, open blinds and curtains in the daytime, reduce shadows</td>
</tr>
<tr>
<td>Diet</td>
<td>Reduce / eliminate caffeine, excessive sugar with dramatic blood sugar fluctuations</td>
</tr>
<tr>
<td>Medication(s)</td>
<td>Request provider / consultant psychiatrist / pharmacist to review meds and suggest rational deprescribing</td>
</tr>
<tr>
<td>Medical condition(s)</td>
<td>Request provider to assess for potential reversible causes (e.g., pain, dehydration, infection, vitamin deficiencies, thyroid dysfunction, hypo/hyperglycemia)</td>
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**Trauma-informed care in person with dementia who cannot remember traumatic event**

**Step 1:** Try to find out as much details from collateral sources of the nature of trauma (including TBI). Keep in mind that trauma maybe from past events as well as present experiences. For example: use of physical restraints in a hospital setting; holding the person to give intramuscular injection.

**Step 2:** Try to avoid situations that would mimic the trauma experienced.

**Step 3:** Monitor for the person’s statements that may indicate the type of trauma they may have experienced. For example, they may say, “go away...get out of the room” when she sees a male caregiver would indicate a male may have caused the trauma.

**Step 4:** Monitor using observation scales for depression as depression is often caused by trauma. Cornell Scale for Depression in Dementia is one such scale.

**Step 5:** Educate staff that this person has experienced trauma and that their approach should be slow, gentle, closely observing for any triggers, communicating any identified triggers first-hand to other team members, documenting triggers if observed, and then minimizing the person’s exposure to triggers.

**Step 6:** Create a psychosocial wellness care plan that includes focus on the following:
- Trust building – the caregivers reassure that they are there to help and not harm
- Identifying and avoiding triggers (e.g., avoiding male caregivers if trauma was caused by a male)
- Reducing sympathetic nervous system overdrive – calming rituals and routines, simple breathing exercises, lavender lotion three times a day, chamomile tea or extract twice a day, soothing music, calm and predictable environment are examples of achieving this. These same strategies can be employed when the person is having trauma-related symptoms (e.g., nightmares, flashbacks, panic attacks)

**Step 7:** Consult a clinician for possible need for psychiatric medications if symptoms are moderate to severe and not responding to psychosocial spiritual, behavioral and environmental approaches.

**Note:** The order of the steps can be changed to better meet the needs of the person.

Namaste
Routine Psychosocial Spiritual / Non-Drug Interventions (NDI) Orders for AAD

1. Lavender lotion (or another scent that the person prefers) three times daily and as needed
2. Personalized music / soothing music 20 min three times daily and as needed
3. Walking (with assistance if necessary) 15 min three times daily and as needed
4. Outdoor sitting and sipping chamomile / lavender tea twice daily and as needed
5. Bright light therapy 20 min – 2 hours once a day (morning or afternoon) and as needed

Namaste
**Relationship Building Communication Techniques**

Relationship building communication approaches are an essential skill for all healthcare professionals. Many persons having dementia (PhD) are looking to be part of warm loving relationships when they express wish to “go home” and relationship building communication approaches are key to making PhDs feel emotionally secure. “Agitation” in PhDs is also frequently related to feelings of insecurity, loneliness, lack of feeling connected and feeling “lost” (in the interpersonal world).

There are at least six key relationship-building communication techniques that all care-partners / caregivers (family and professional) of PHDs should become familiar with and utilize in their daily interactions with the PhD.

*Reflective listening:* This technique involves making eye contact, adjusting body posture (e.g., leaning forward, avoiding hands folded across one’s chest), and providing nonverbal and verbal encouragement for PhD to express themselves (e.g. nodding, stating verbal continuers [e.g. “I see,” “Go on,” “Sounds like…”]). It is important to avoid expressing judgment, getting distracted, or redirecting the PhD. Repeating what the PhD has stated and seeking clarification is also important. *Humanitude training* may facilitate development of this skill.

*Elicit storyline:* Positive (e.g. joy) as well as negative emotions (e.g. anger) are embedded in stories that are unique to each PhD. A PhD’s narrative can be brought out through use of open-ended questions and continuers (e.g., “tell me more,” “what next?”) as well as Improv techniques such as “Yes and”. Understanding the PhD’s narrative is essential to seeing the PhD as a unique person.

*Explore the PhD’s perspective (point of view):* When another person understands our perspective, we feel heard. Specific questions can help elicit PhD’s perspective (e.g. what is your opinion? How do you feel about the care you are getting? Is anyone bothering you? How are they bothering you?).

*Validate PhD’s experiences:* When our experiences are validated, healing occurs. Validation can be done by genuinely felt and expressed statements such as, “I would also feel scared if I was in a strange place...I would feel sad too if my family was not with me.” Giving advice, opinion and correcting or interrupting the PhD is avoided. Gentle touch is important.

*Mirroring:* This technique involves mirroring PhD’s emotions such as a smile or sadness. Mirroring conveys to the person that we understand their emotional state.

*Touch:* Gentle touch can promote bonding through release of oxytocin (and other mechanisms) and help build trust in relationship.
**Case Example: When do I go home?**

Ms. G has been very upset, anxious, and tearful almost every day from the day a few weeks ago when she was admitted to a secure memory-care unit of an assisted living community. She has been repeatedly stating “when do I go home.... did I do something to be here...just kill me...” Staff listens to Ms. G using reflective listening skills mentioned above (e.g., “sounds like you are unhappy here”). Staff then make efforts to bring out the PhD’s narrative (e.g., where would you like to go? Tell me more about your home?). Staff then try to explore Ms. G’s perspective (e.g., “what is worrying you? What would you like us to do? If you could have anything, what would you want? Why is it so terrible over here?) and validate her feelings of abandonment and loneliness. Staff also mirrors PhD’s sadness and frequently holds PhD’s hands.

Relationship building communication approaches over time will give Ms. G a sense that staff care, that they understand her perspective and this will help Ms. G slowly but surely feel less anxious and lonely. Ms. G will feel part of several loving relationships with different staff members.

**Resources:**


Namaste
**STEPS: Support system / Staff Training Education Praise and Support**

STEPS is a comprehensive approach to improving quality of care provided by family members (patient’s support system) and professional caregivers (e.g., staff to our long-term care residents). It involves three components:

1. **Education and Training**: Two key areas:
   b. Mindfulness training. Goals of this training: Improved skills to provide care with mindfulness (with awareness and compassion) and Increased Joy at work (besides reduced burnout)

2. **Praise**: This includes case-based in-time praise for specific behavior of the staff that lead to specific positive outcome. Sharing success stories routinely during meetings and at other times is one of the many ways to praise staff regularly.

3. **Support**: This includes direct one-on-one support by leadership team members, one-to-one support by colleagues and group support strategies.


Two essential trainings:

1. DICE – Describe Investigate Create Evaluate. [www.diceapproach.com](http://www.diceapproach.com)
2. Mindful Care-Partnering (see page
**Caregiver Dementia - CD-9 screening tool**

1. Do you forget that TRACing is counterproductive?
2. Do you forget that BITEing is counterproductive?
3. Do you forget that your loved one has dementia?
4. Do you repeat the same mistakes?
5. Do you forget to take time to maintain your health?
6. Do you forget what you have learned in support groups and caregiver education and training workshops?
7. Do you get confused about how to cope, what to say?
8. Do you have emotional outbursts?
9. Do you forget to give yourself a break?

TRACing = Testing, Reasoning, Arguing, Correcting

BITEing = Blaming, Insisting on reality, Taking symptoms personally, Expecting unrealistically

Resource Book: Travelers to Unimaginable Lands. Stories of dementia, caregiver, and the human brain. Dasha Kiper. Psychologist. Caregivers are provided education and training to minimize TRACing and BITEing in their interactions with their loved one with dementia but this book beautifully and with great compassion illustrates just how hard it is.

Scored like Patient Health Questionnaire – 9 (PHQ-9) used for depression; 0-3 points for each. Max score 27. 5-12 mild; 13-20 moderate; 20-27 severe

Rx:
normalize (most caregivers develop mild caregiver dementia)
empathize
expand / improve utilization of their support network
respite
support group
individual counseling
bibliotherapy - Dasha Kiper’s book
mindfulness skills training
creative engagement (e.g., journaling, Improv, poems, art)
other practical help
family interventions
participate in dementia friendly community programs like Memory Cafes

*Caregiver Dementia* 100% reversible but if not addressed, can leave serious negative effects on mental, spiritual and physical health. Reversing *caregiver dementia* is key to reducing AAD. Note: I have made up this term *Caregiver Dementia*. No research to support this.
Caregiver Dementia Questionnaire – 9
CDQ-9

Name of the caregiver / care partner of the person living with dementia:

date:

over the past two weeks, how often:

<table>
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<tr>
<th>#</th>
<th>behavior</th>
<th>not at all</th>
<th>several days</th>
<th>more than half days</th>
<th>nearly every day</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>do you forget that TRACing is counterproductive?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>do you forget that BITEing is counterproductive?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>do you forget that your loved one has dementia?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>do you repeat the same mistakes?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5</td>
<td>do you forget to take time to maintain your health?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6</td>
<td>do you forget what you have learned in support groups and caregiver education and training workshops?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7</td>
<td>do you get confused about how to cope, what to say?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>8</td>
<td>do you have emotional outbursts?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>9</td>
<td>do you forget to give yourself a break?</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

score 0 + + + +

total score (max 27):

TRACing: Testing, Reasoning, Arguing, Correcting the person living with dementia.
BITEing: Blaming, Insisting on Reality, Taking things personally, Expecting unrealistically

interpretation:
mild caregiver dementia: 5-12
moderate caregiver dementia: 13-20
severe caregiver dementia: 21-27

source: Abhilash Desai MD. Geriatric Psychiatrist. Dr.abhilashdesai@icloud.com

Note: This tool has NOT been researched / systematically studied. Dr. Desai has made up the term Caregiver Dementia. There is no research to support this concept.
Don’t TRAC: Test Reason Argue Correct

A common cause of agitation is when family members (especially spouses and children) test their loved one with dementia, try to reason with them, argue with them or correct them.

It is important for family members to take a deep breath and go back to minimizing TRACing.

Don’t:

- Test: avoid asking questions that require memory such as “what did you do today, what did you eat.”
- Reason: avoid trying to reason with them and trying to use logic to get them to change their beliefs or behavior based on their perception of reality
- Argue: avoid going back and forth stating your point and perspective again and again
- Correct: avoid correcting them

We have seen agitation escalate to verbal abuse and even paranoia and physical aggression if the family caregiver continues to TRAC.

It is very difficult not to TRAC so family caregivers need lot of support and guidance in their efforts to master this important communication skill. In many situations, individual counseling is needed for family caregivers to tackle the trauma, grief, guilt, fear, and shame they are struggling to cope with.
Don’t BITE: Blame, Insist on Reality, Take things personally, Expect unrealistically

A common cause of agitation is when family members (especially spouses and children) start either blaming the person with dementia or themselves about mistakes or setbacks; when family members insist on reality, take what the person with dementia says in anger, personally; and when family members have unrealistic expectations from their loved one with dementia.

It is important for family members to take a deep breath and go back to minimizing BITEing.

Don’t:

- Blame: avoid blaming the person with dementia (e.g., “you make me feel bad about myself”) or oneself (e.g., “see what you did, you are the reason she is unhappy”).
- Insist on reality: avoid trying to insist that the loved one with dementia live in reality, especially your (family caregiver’s) reality
- Take things personally: avoid believing every negative statement and accusation your loved one with dementia makes regarding you (e.g., “you are a terrible daughter for leaving me here” [meaning in the assisted living community])
- Expect unrealistically: avoid unrealistic expectations (e.g., the loved one with dementia will be reasonable or grateful for all the sacrifices you – the family caregiver – have made).

We have seen agitation escalate to verbal abuse and even paranoia and physical aggression if the family caregiver continues to BITE

It is very difficult not to BITE so family caregivers need lot of support and guidance in their efforts to master this important communication skill. In many situations, individual counseling is needed for family caregivers to tackle the trauma, grief, guilt, fear, and shame they are struggling to cope with.
T-A-DA method of managing agitation

Dr. Joseph Flaherty, geriatrician at Saint Louis University School of Medicine has a YouTube video on this. Please check it out. Although he describes this method to manage agitation in the context of delirium in the hospital setting, all the principles apply also to management of agitation in individuals with dementia.

T-A-DA is an acronym. It stands for:

T: Tolerate - increase your tolerance for negative behaviors of the loved one with dementia as often times, they are transient [like bad weather] and pass on their own).

A: Anticipate – anticipate the unmet needs and triggers so that agitation can be aborted. For example, anticipate that they will get agitated if not toileted frequently, if they don’t drink adequately throughout the day, if they get constipated or are in pain. Best to see all agitation as a way of communication of unmet needs (biological needs like hunger; psychological needs like having a choice; social needs like companionship; spiritual needs like having a purpose).

DA: Don’t Agitate. TRAC and BITE mentioned in earlier sections are common ways that family caregivers can worsen agitation. Other ways they can worsen agitation is by ignoring the persistent negative behaviors of the person with dementia (aka not investigating and meeting their unmet needs), and giving them psychiatric medications when the agitation is due to dehydration or pain or constipation or other biopsychosocial needs.
As we mentioned earlier, most family caregivers struggle to understand the negative emotions, statements, behaviors, verbal and physical abuse and paranoia expressed by their loved one with dementia. Family caregivers have told us that our suggestions on “Don’t TRAC” and “Don’t BITE” are “Mission Impossible” and that we are having unrealistic expectations from the family caregivers and we (my wife and I) are doing the same thing that we are telling them (the family caregivers) to stop doing. We are having unrealistic expectations from family caregivers.

We acknowledge everything that the family caregivers are saying and agree that it often is “Mission Impossible.” Both of us have been family caregivers to our elderly parents (one of them had vascular dementia) and it indeed was “Mission Impossible” at times.

With lot of support, guidance, role-playing (I take the role of the family caregiver and the family caregiver takes the role of their loved one with dementia), Improv, creative and compassionate engagement and communication, and time, we do feel that many family caregivers are able to dramatically reduce TRAC and BITE behaviors, increase T-A-DA approaches and will see amazing results in terms of not only reduction in agitation but even increase in frequency, duration and intensity of joyful and happy moments.
Mindful Caregiving / Care-partnering – Improv Workshop Handout

Mindfulness is our capacity to be in the present moment, with intention and non-judgmentally. The two key ways to be mindful while providing care to your loved one with dementia is by accepting the challenges involved in caring and engaging in creative and gentle ways.

This workshop will involve learning the two key skills of mindfulness – acceptance and creative engagement – through the use of Improv and one-minute guided meditations.

Improv is a form of theater in which much of the performance is created spontaneously, without any scripts. Improve, using a variety of techniques and strategies including comedy, has been shown to improve skills like empathy, teamwork, and creative problem solving.

During this three-day workshop, we will engage in at least two two-minute meditations, one at the beginning and one at the end of each day. The mindfulness teacher will guide you through the meditations and mind the time. There will be a brief lecture and discussion to explain mindfulness and mindful caregiving / care-partnering.

Instructions for guided meditation:

Get into a comfortable position. Gently close your eyes. Keeping eyes open and focused on something is okay. Put one hand on your belly and another on your chest and bring your attention to the movements of your hands as you breathe in and out. The mindfulness teacher will be repeating the aspirations “may you accept the challenges involved in caring; may you creatively overcome the challenges.” In between the aspirations, you will bring your attention back to the movements of your hands as you breathe in and out. As we do this meditation, your attention is likely to wander. No need to get upset. That is what our mind does. When you become aware that your attention has wandered, gently bring it back to your breathing and rejoin the meditation. Let’s begin the meditation

Improv training will involve a few interactive sessions to help you utilize Improve strategies and techniques to mindfully switch from negative TRAC to positive TRAC during caring.

Negative TRAC – Testing, Reasoning, Arguing, Correcting

Positive TRAC – Touching gently, Reassuring, Accepting and validating, Creative problem solving

One of the fundamental skills of Improv is “Yes And.” During this training, an Improv expert will teach us this fundamental skill and how it can be applied during periods of challenging behaviors experienced by your loved one with dementia in order bring them some relief.

If you are struggling, you can change the C for Creative problem solving to Compassion for self. This involves giving yourself a break and seeking emotional support for yourself as caring is hard. Self-compassion is another essential mindfulness skill and a core component of resilience.
Bright Light Therapy

Depression, insomnia and sun-downing related agitation are prevalent in persons with dementia and is typically treated with antidepressants and or sedating medications (e.g., trazodone). Bright light therapy (BLT) is a good non-pharmacological intervention that may help some residents with depression and or insomnia. It should generally be used along with other non-pharmacological interventions (e.g., behavioral activation, individualized pleasant activity schedule, exercise, music, gratitude-based interventions, dignity therapy) and / or pharmacological interventions.

If you prefer natural sunlight, try to get it early in the morning as damaging UV light is lower in the morning. Sitting next to the window for sunlight is also fine.

Usual indoor lighting is not strong enough to provide benefits.

BLT should be considered for all residents with seasonal affective disorder (especially winter depression) and may also help residents with non-seasonal Major depression and or Insomnia disorders (e.g., chronic insomnia). Research has also found it useful as an adjunct to pharmacological interventions for Bipolar depression (1).

BLT with sleep hygiene may improve sleep in individuals with dementia.

I recommend a BLT lamp (NLT bright light therapy lamp) available at the Center for Environmental Therapeutics (https://cet.org/shop/). It costs around $190 (includes shipping). I have no financial relationships with the makers of this BLT box. I recommend this because it was used in research settings. Other sources of BLT lamp are also fine.

A 10,000-lux light box is recommended. Intensity needs to be individualized and is effective in the range of 2,500 – 10,000. For most cases, 10,000 lux is needed for 30 minutes daily for beneficial effects. Light boxes with larger screens are preferred over smaller screens due to less eye strain. If lower intensity is used, extend the exposure to more than 30 min.

Benefits can be seen as early as 2 days but generally take 2-4 weeks. Many patients who show improvement in the first four weeks continue to improve further over next 4 weeks (2).

The resident can begin exposure to bright light for 15 minutes initially and slowly increase (15 minutes per week) as tolerated to 60 minutes per day. The box should be above the resident’s head at a 45-degree angle. One can start with exposure in the morning (6am-9am) but some residents may respond to exposure in the afternoons (noon-3pm). The resident is generally around 30-80 cm (30cm = one foot) from the screen, not looking at the screen but engaged in another activity (e.g., reading, watching TV).

Adverse effects are typically minor and transient and include eye strain, nausea and headache. Using BLT for a shorter time or lowering the intensity may be needed to reduce these problems.
for some individuals. Although rarely BLT could trigger a switch to a manic / hypomanic episode, in the two studies (1,2), that did not happen. Both the studies had a small N (small sample size).

Reference:

Resource:
1. University of Wisconsin patient handout on BLT.

Namaste
Ms. Linda came to our center at the beginning of this year. In the beginning she was apprehensive about long term placement, she was frustrated at her family for leaving her and distressed about leaving her animals. Ms. Linda quickly found companionship with those around her and found she enjoyed sitting next to fellow residents, giving hugs, candy and kisses on the forehead. When COVID precautions began, Linda struggled. She became frustrated at staff reminders to social distance. After a few months Ms. Linda settled into her new way of life, visiting others from afar and waving at fellow residents as she walked through the hallway. As precautions increased due to facility exposure to COVID-19, Ms. Linda was encouraged to stay in her room to decrease her level of exposure. These increased precautions resulted in a significant increase in behaviors. Staff reported that Ms. Linda was throwing objects at the window to see if the glass would break, formulating plans to escape, threatening to kill staff if they wore the proper protective gear in her room, hallucinations of Jesus and her grandmother in her room. To help deescalate behaviors staff would visit with Ms. Linda throughout the day, taking her on walks outside, and facilitating socially distanced family visits. Staff quickly found out that Ms. Linda enjoyed feeding the squirrels in the courtyard and watching them eat from her window. Now, twice a day Ms. Linda will seek out social services or activity staff to help her feed her "babies." There has been a visible improvement in Ms. Linda's behaviors, she feels she has a purpose in life and that she is "God's helper."

Credit: Credit for this goes to a wonderful social worker, Ms. April Bench that I have the privilege to work with in a local nursing home.
Doll Therapy (DT) for Behavioral and Psychological Symptoms of Dementia (BPSD)
Abhilash Desai MD

Landmark study:

Intervention: 2 hours of DT twice daily (morning and afternoon) and prn for agitation / aggression / wandering. The patient’s interaction with the doll was observed for 7 days. Patients with positive attitude towards the doll were entered into the study.

Results: DT is more effective than standard treatment for reducing BPSD and may be effective in reducing the incidence of delirium. DT also reduced perceived professional caregiver burden.

Dr. Milta Little (one of our national and international leaders in geriatric medicine and dementia care) stated in her discussion of this study at the annual 2022 conference of the AMDA (American Medical Directors Association [same as PALTC Post-Acute and Long-Term Care society]) that “doll therapy is based on the principles of attachment…dolls may actually be used as a translational object where people with dementia may be able to translate BPSD into a caregiving role towards what they perceive as a baby” – especially in cases where BPSD is due to unmet attachment needs. Dr. Little also highlighted the finding in the study that in 28 out of 32 instances when DT was used as needed for agitation, it was effective (with no need for as needed psychiatric medications) and in the other 4 instances, as needed medications were needed.

I agree totally with Dr. Little and would add that many patients who respond to DT may be feeling better because taking care of the doll gives them purpose and makes them feel needed. I have seen benefits with DT when used in a respectful manner (after education of staff and family that our intention is NOT to infantilize the person but to enhance positive experiences).

Doll therapy is controversial as many would see this as infantilizing the person with dementia. It is important to talk about this concern with family and team.
Resistance to Care in persons with Dementia

Individuals with advanced dementia often resist personal care, and sometimes even become aggressive. Antipsychotics are often used to reduce aggression during personal care. The following strategies are best practices to reduce distress experienced by the person with dementia during personal care and improve their comfort during such care.

If done with love, patience, training and creativity, these times of personal care can become an important bonding experience for both the person with dementia and the caregiver. Both may experience positive emotions, beauty in relationship and deeper meaning in life and living.

The Seven Essential Strategies

1. Know the individual – really know them (read a brief personal statement that represents the individual’s unique attributes and essence). This would apply not only to professional caregivers (e.g., staff in nursing homes and assisted living communities) but also family members who are struggling to see the person behind dementia.
2. Identify and consistently use individualized effective communication strategies (read the strategies and even post the strategies on the bathroom door)
3. Involve the individual in their own care (allow them to do what they can or put their hand on yours)
4. Gently massage the individual’s hands and feet (lavender lotion can be soothing and relaxing, but any other aroma may also work)
5. Engage and distract with positive images (pictures that have been found to hook the individual in a positive way and have been identified in advance)
6. Address pain (scheduled acetaminophen if appropriate)
7. Soothing background music (preferably personalized)
ABCD of Spiritual Assessment

A = Ask. Ask permission to inquire about patient’s spirituality. Explain that spirituality is an important factor in wellness, resilience and healing. Involve family whenever possible.

B = Beliefs. Inquire about their spiritual beliefs so that you can better understand them.

C = Community. Inquire about the patient being part of a spiritual community.

D = Daily practice. Discuss what daily spiritual practices can be continued and new ones added to promote wellness, resilience and healing.

Common spiritual practices and rituals
- Prayers
- Meditation (Healthy Minds Program – Best app for learning to meditate and understanding mindfulness)
- Reading religious scriptures and spiritual books (including audio books)
- Attending church, mosque, temple events and events and gatherings of other spiritual communities
- Volunteering
- Yoga, Tai Chi, Qi Gong
- Spending time with nature
- Listening to spiritual / religious podcasts
- Singing religious / spiritual songs and dancing to it
- Gratitude and Awe – based practices and rituals

Suggested reading:
Dignity Conserving Care

Dignity conserving care can improve depression and anxiety in older persons and greatly enhance their functioning and quality of life. For persons with dementia, doing this with their loved one besides the person is key.

1. Tell me a little about your life history, especially the parts that are the most important / meaningful to you.
2. When did you feel most alive?
3. Are there specific things that you would want your family to know about you and particular things you would want them to remember?
4. What are the most important roles you have played?
5. Why were these roles so important to you and what did you accomplish in those roles?
6. What are the most important accomplishments and what do you feel most proud of?
7. Are there particular things that you feel still need to be said to your loved ones or things that you would want to take the time to say again?
8. What are your hopes and dreams for your loved ones?
9. What have you learned about life that you would want to pass along to others?
10. What advice or words of guidance you would wish to pass along to your (son, daughter, husband, wife, others)?
11. Are there words or perhaps instructions that you would like to offer your family to help prepare them for the future?
12. In creating this permanent record, are there other things that you would like included?
SPPEICE: Strengths-based Personalized Psychosocial behavioral nutritional sensory spiritual Environmental Initiatives and Creative Engagement (aka Non-Drug Interventions) to improve psychosocial emotional cognitive spiritual wellbeing of individuals with advanced dementia.

In-depth information on SPPEICE is in chapter 13 of the book titled *Psychiatric Consultation in Long-Term Care: A guide for healthcare professionals* by Cambridge University Press, 2017.

- Relationship building staff approach (includes getting to know the resident well [using Life Story page posted outside the door or just inside the door, Life Story picture book, Life Story bin], validation, AwareCare approach)
- Spend time with friends and family
- Touch (includes massage)
- Exercise (includes postural exercises, walking with a song, chair Yoga)
- Music therapy
- Multisensory approaches in a multisensory room
- Individualized pleasant activity schedule with continuous activities throughout the day
- Pet / Animal assisted therapy
- Aromatherapy
- One on one
- Simulated presence therapy (e.g., video and audio recording a family event for 1-5 minutes and showing it to the resident; family may need some guidance and coaching and “multiple takes” to get the most emotionally strong and positive recording)
- Robotic pets (see Amazon video: Hasbro’s Joy for All Pet interactive puppy and cat; available on Amazon [around $120])
- Fidget blankets (available on [www.etsy.com](http://www.etsy.com) cost $15 upwards)
- Avoid / minimize intake of caffeine and chocolate
- Avoid alcohol
- Preferred food and drinks
Resources for SPPEICE:

1. AwareCare approach by Linda Clare PhD (http://psychology.exeter.ac.uk/media/universityofexeter/schoolofpsychology/reach/documents/AwareCare.pdf). A must for every staff working in long-term care and with persons having dementia.

2. TimeSlips (https://www.timeslips.org) creative storytelling. Online training course available. At least one long-term care staff should be trained in TimeSlips.


4. Cognitive Stimulation Therapy (http://www.cstdementia.com). Online training course available. At least one staff member should become trained in CST.

5. *Bathing Without A Battle* DVD (http://bathingwithoutabattle.unc.edu). All staff should undergo free online training.

6. *Alive Inside* movie (http://www.aliveinside.us). All staff should watch this movie and long-term care leadership should consider this work-education time.

7. Idahoan Dave Potter’s website https://palousemindfulness.com provides opportunity for FREE self-paced online training on Mindfulness and Meditation (eight-week Mindfulness Based Stress Reduction [MBSR] course). At least one long-term care staff should be trained in MBSR.

8. Multisensory room. Watch the video on Healing Spaces at https://news.usc.edu/147930/healing-spaces-video-game-targets-alzheimers-dementia-patients/

9. Educational programs by the Occupational Therapist and a celebrity in the dementia training world, Teepa Snow http://teepasnow.com
Strengths-based Personalized Psychosocial sensory spiritual and Environmental Approaches and Creative Engagement (SPPEICE) for Mr. MT.

When Mr. MT becomes agitated and calls out and reaches for things on the floor, provide him with things to do and the behaviors may quickly go away. Use the same activities to prevent agitation from happening in the first place. In general, give him only one activity at a time and put away the one before. This is to prevent him from getting overwhelmed or distracted. Rotate activities after 30-45 minutes or when Mr. MT loses interest.

Suggested additional activities and approaches (besides coloring and puzzles):

1. Use cookies to encourage resident to move to a different area in the common room.
2. Staff to start playing with the blocks for a few minutes and then encourage resident to try messing around with the blocks.
3. Encourage resident to play deck of playing cards with the bright circus pictures. With the recreational therapist (RT), for almost 45 minutes Mr. MT contently played various ways with the cards. He was counting them. He was putting them in piles. He was talking to them. Twice RT approached Mr. MT to help him sit up better but then without further direction left him to himself again.
4. Staff to approach Mr. MT respectfully to ask if they could show him something interesting. If he says “yes,” give him a National Geographic Magazine. He may flip through it, stop and observe many pages. This activity may engage him for 15 minutes or more. Keep the cards around as he may put the magazine aside and pick up the cards again.
5. Encourage resident to stand, stretch and walk (with assistance as necessary) around for a few minutes (or more) every hour.

Create a plastic bin with the following:

- Cards of several kinds – regular playing cards, Wild Kingdom Animal Playing Cards, playing cards that have pretty scenery or interesting pictures on them.
- Several interesting magazines like National Geographic, gardening, travelling, Architectural Digest (for his engineering background). Often used magazines can come from libraries or family members and friends if the word is given that they would like them.
- Many Nuts, Bolts, Washers and Screws of various kinds in a plastic container and 2-3 disposable muffin tins for sorting. The family may purchase the hardware at Second Chances or Goodwill at relatively low cost. Have Mr. MT sort the hardware and/or match up the nuts and washers. Staff may watch Mr. MT the first time for 5 minutes to make sure he isn’t trying to eat them.
- Historical picture books that have very little words but interesting pictures. Other hardback picture books of things Mr. MT used to love to do (e.g., gardening, outdoors), and other hard back books.
- Easy word searches, mazes, and easy brain builders.
- A Matching game with only give 20 matching cards at a time.
- Blocks (multicolored Jenga game)– especially wooden blocks instead of the plastic large Legos which Mr. MT might view as childish. Encourage resident to build a tower all one color or different colors. Resident can sort according to color. Staff can make a pattern and encourage resident to copy the pattern.  

Note: Keep the bin in a place that is accessible to Mr. MT. The content of the bin can be modified over time so that activities that are of interest are kept, activities that Mr. MT did not find interesting can be removed and new activities may be tried.

**Address Posture:**
- To encourage better posture and reduce pain due to abnormal posture, the staff may push Mr. MT’s wheelchair forward for him as he leans too much to move his own wheelchair. The staff may prop Mr. MT up and to the side (that is leaning) with a pillow to straighten his body so he sits almost upright. Mr. MT may also be able to handle the items on the table much better after the posture is corrected.
- Staff to transfer Mr. MT to a regular sitting chair at the table so that it doesn’t slide as easily. Staff may lock the wheelchair in place if the resident wants to be at the table and not constantly roll away from the table?

**Therapeutic Communication:**
- Staff need to get to the eye level of the resident and make eye contact when interacting rather than standing over Mr. MT or answering from across the room.
- Staff encouraged to avoid statements such as:
  - Let go of the knob. You don’t need to hold it. Let go.
  - You said you needed the restroom. That’s what we are here for. Let’s do it then.
  - Mr. MT, sit down, sit down. Just wait one minute and we will go.
- Staff are encouraged to make statements such as:
  - thank you for letting me help you
  - Please sit down for a moment
  - Mr. MT, I see your frustrated. I’m sorry. I’d like to help. Please let go of the knob;
  - Mr. MT, please let go of the knob. That’s great, thank you!
**Strengths-based Personalized Psychosocial sensory spiritual and Environmental Approaches and Creative Engagement (SPPEICE) for Mrs. B.**

Brief Personal History: Her daughter reports that Mrs. B loved sewing and crafting. She was involved in the church as a Nazarene and took her religion very serious. She enjoys Bible reading. Beaches were one of her favorite places to visit. Daughter reports she can still read.

1. **Addressing spiritual needs**
   a. Mrs. B may find a large print simple devotional beneficial (www.lifeway.com an online Christian bookstore [there are also stores in Boise] may be a good place to find a large print devotional).
   b. A subscription to a conservative Christian magazine may also be worthwhile as a way to meet Mrs. B’s spiritual needs. Looking at the pictures, advertisements and perusing the articles may remind her of her days in the Nazarene Church. Magazines such as “Today’s Christian Living” or “Christianity Today” may be good options.
   c. **Environmental enrichment initiatives**
   d. Consider putting a couple of large beach pictures that have scriptures on them to be place on the walls. Mrs. B spends considerable time looking around her room and seeing those might add warmth and encouragement.
   e. Please consider removing the red bag hanging in the room since, sometimes it causes her distress.
   f. Mrs. B may enjoy listening to music, reading her devotional or enjoying her fabric squares in the nicely lit front living area of the Orchards. People come and go and it is a bright cheery room. This might help her disposition and reduce weepiness if she has a warm drink and something to do in front of the large windows.

2. **Tapping into the Magic of Music**
   a. Christian music may be particularly beneficial because of Mrs. B’s church background. Suggest soft inspirational worship music and instrumental worship music. A CD player could be added to her room since she spends quite a bit of time in her room. I would also suggest some soothing beach music. If the resident enjoyed other music (oldies, jazz, etc.) an additional CD may be added.

3. **Relationship Building Approaches (Social enrichment initiatives)**
   a. Regular interactions with 1-2 other resident ladies Mrs. B could chat with if they have similar interests (magazines or fabric squares) may provide socialization and comradery. Mrs. B was quick to enjoy conversational as the recreational therapist was there and seemed to enjoy conversing with her. Since Mrs. B spends a lot of time in her room, interacting with (or even being around) other pleasant ladies in a common area may be very beneficial.

4. **Individualized Pleasant Activities**
a. Mrs. B loved crafts and sewing and often talks to herself about knitting. A small bin of fabric squares would bring her joy. 5-6 pieces of 5-6 different kinds of fabric cut into 4” squares may allow her to sort them, arrange them in patterns or just look at them. Suggest brightly colored prints and different textures.

b. A lap mechanical pet may bring great comfort since Mrs. B spends much time in her room. This animal could sit on her bed or lap. Suggest the www.joyforall.com Hasbro mechanical pets because they are so life-like that most residents feel they are real. They take 4 C Batteries that only need to be replaced every 4-6 weeks depending on the extent of use. Depending on the resident’s preference Hasbro offers dogs or cats.

c. Touch: Hand massages with lavender lotion for 5 minutes four times a day and as needed. Especially if resident Mrs. B is having anxiety due to delusions or hallucinations, it can help calm her mind and body.

d. Physical Activity: Suggest 5-10 minutes of physical activity where daughter or a caregiver plays bean bag toss or ring toss game with Mrs. B. Such bags and rings are relatively inexpensive and may be found on www.orientaltradingcompany.com or at a local toy store. Mrs. B has mentioned that one time she played a game of bean bags and liked it. For this activity the resident could have a set in her room and when her daughter visits they could play and staff can also play before or after any care-giving tasks.
Strengths-based Personalized Psychosocial sensory spiritual and Environmental Approaches and Creative Engagement (SPPEICE) for Grace.

The following is a list of ideas on how to better manage interactions with Grace. When she is “acting up” she can be difficult to work with, but here are some ways to avoid these episodes or calm her down during them. The information comes from people who have worked closely with her during her time on our unit, such as nursing staff and family.

1. **Grace loves praise.** When you need Grace to do something, or even if you would like her to calm down, she responds well if you focus on positive things she is doing. For example, if you would like her to walk a certain distance, tell her she’s doing a great job and she will walk twice as far as your initial goal.

2. **1-on-1 communication.** When Grace is “acting up,” the environment around her is key to getting her to calm down. If possible, make sure she is in a room with just her and one other person. She will respond much better if she feels that one person is talking to her at her level (and speaking loudly) than if she has 3 nurses surrounding her to calm her down.

3. **Grace loves animals.** Cats especially – at the nursing home we have heard that there are several cats and that when they are around she is much less likely to act out.

4. **She loves Bingo.** This is Grace’s favorite activity and she has played it every day for years up until recently, when she has had trouble with her cataract on her ‘good’ eye. If there is a way to have someone play with her as her partner and help her with the visual aspect of the game, this could be a great way to keep her calm or distract her during fits.

5. **Play music.** Having music playing is a good way to relax her. She enjoys all genres.

6. **Avoid CNN.** Try to avoid upsetting or serious television programs. These tend to get her worked up and anxious and appear to contribute to her paranoia.

7. **Get her walking.** She often complains that she is in her chair too long or in her bed for too long, but needs encouragement to get up and get moving. She also currently has a pressure ulcer on her coccyx and the pain could increase her agitation.

8. **During an episode, give her time alone.** When Grace acts out, sometimes it is best to let her spend time alone in her room. She needs an environment with low simulation to calm herself down.

Thank you for reading this and hopefully it helps Grace enjoy more of her time and makes her caretaker’s job easier.

Credit for this goes to an amazing medical student Matthew Colburn, MSIII at Saint Louis University many moons ago.
Sexually “Inappropriate” Behaviors in Persons with Dementia

Psychosocial Environmental Approaches

Words matter: Let’s call them socially inappropriate behaviors. Masturbation and other forms of self-pleasure per se are not sexually inappropriate behavior as long as they are done in private.

Behaviors that are most problematic: Masturbation / self-pleasure in public places, touching others (typically caregivers) in private parts without their consent.

Assessment: PCP assesses the person with dementia for medications (e.g., drugs that increase dopamine [drugs used for Parkinson’s disease]), medical-psychiatric conditions (e.g., mania, psychosis) that may cause / contribute to hypersexuality and impulsive sexually “inappropriate” behaviors.

Key Psychosocial Environmental Approaches:

- Try to understand the behavior as an unmet intimacy need and address the need directly.
- Limit the changes in the person’s life. These can make them confused and keyed up and lead to new or different sexual behavior.
- Provide physical touch. Everyone has a need for loving touch and physical contact. This need is generally unmet in individuals with dementia, especially in residents living in nursing homes. Find a way to touch them as part of your everyday routine. Hold their hand or give them a hug or back rub.
- Spend time with them. Keep them entertained: Look at photo albums, play board games, or go for a walk. These activities can prevent boredom that can lead to sexual behaviors.
- Avoid things that trigger the behavior. If it happens regularly, pay attention to what happens right before and try to avoid it.
- Allow certain behaviors in private. Masturbation and other forms of self-pleasure may be one of the few ways someone with dementia can feel pleasure or relieve sexual desires. If they do it in private and don’t hurt themselves, it’s often best to support it.
- Educate the family / staff:
  - That sexual needs, need for physical touch and sexual pleasure is normal, even in the old-old and in presence of advanced dementia. The sex drive is seen in many older adults for as long as they live.
  - Dementia often impairs frontal lobe function – the seat for socially appropriate behaviors (disinhibition) – and so the normal expression of sexual needs gets manifested in socially inappropriate ways. There is no reason for the caregiver (family / professional) to take offense or take it personally.
  - Dementia often causes the person to get “stuck” in one area and it is hard for them to get out of on their own.
  - It is important to stay calm, be patient and respectful and avoid using words or tone of voice that would make the person feel ashamed. The person is just
looking for human connection, love, affection, freedom from boredom and or looking for excitement and fun.

- Watch body position and avoid reaching over the resident.
- Create a safe distance between you and the person with dementia (or between the person with dementia and another resident in nursing home) so that the person cannot “grab” private parts.
- Guide the person to a private area so that they can engage in self-pleasure.
- Use distraction – redirection techniques if the resident does not want to self-pleasure in private. Change the subject, turn on the television or music. Goal is to engage the person in fun and meaningful activities.
- Remove triggers.
  - It may be necessary in some situations to say respectfully but firmly (slightly raised voice), “stop” or “no” to the person with dementia if they are touching you inappropriately. Then back away and re-approach after some time. It is important for all staff to be consistent in indicating that this behavior is not appropriate. If even a few staff condone such behavior, laugh and tease back, the behavior may become harder to address.
  - Understand that the behaviors are not the fault of the person with dementia but also, the caregivers (family, professional) have a right to and obligation to protect themselves and other residents from inappropriate verbal and physical behaviors (which may even be traumatizing).
  - Caregivers (family, professional) should not be arguing, scolding, reasoning, punishing, or shaming the person with dementia with these behaviors.

- **Tracking (for nursing home residents)** – behavior log (number of episodes per 8 hours):
  - verbal towards staff, physical towards staff, verbal towards other residents, physical towards other residents
- **Goals (for nursing home residents) - examples**: the person with dementia will engage in at least one socially appropriate behavior to meet their sexual needs (e.g., self-pleasure in private, physical touch – hugs as often as possible); the person with dementia will have less socially inappropriate behaviors (e.g., less than one episode per 8 hours).

- **Interventions (examples) for nursing home population**:
  - Monitor the resident’s problematic behaviors using the behavior log (accurate logging of behaviors is an absolute essential – interventions generally help modestly, and successful interventions need not be stopped even if benefits are modest as long as there are no / minimal negative effects)
  - Assessment (as mentioned above)
  - Provide resident privacy for self-pleasure (sign outside the door stating “do not disturb”; knocking on the door twice, calling the resident’s name and asking for permission to enter)
  - Promote holding hands, hugging, back rub and kissing (on the cheek or forehead) as safer and appropriate ways to meet some of the person’s intimacy and affection needs.
  - Educate and train staff (especially new staff and staff new to dementia care) of appropriate person-centered communication techniques
- Distract resident with fun and meaningful activities
- Avoid triggers (e.g., boredom, loneliness)
- Staff to wear professional attires such as scrubs

Opinion and Guidance from other Experts:

1. Dr. Jim Wright, MD, PhD, CMD, medical director at Westminster Canterbury Richmond in Virginia has talked about “the empowerment of impermanence” in his presentation on “Intimacy and dementia in the long-term care setting” at the PALTC22 in Baltimore, MD (annual AMDA conference). He said, “to identify the personhood of someone is to say ‘I give you permission to change at every stage of life.’ Being whole person means being allowed to express preferences today that are different from those in the past....letting them make choices inconsistent with their previous choices and values (or even their advance directives), and allowing their sexual preference to change.” He shared the story of Justice Sandra Day O’Connor, whose husband had dementia, was in a LTC home, and had formed a romantic relationship with another woman. Justice O’Connor approved of the relationship because she wanted her husband to be happy, even as his dementia advanced.

2. Dr. Peter Jaggard, MD, CMD, medical director at Presbyterian Homes in Evanston, IL also spoke along with Dr. Wright. He recommended that nursing assessments for ISB (inappropriate sexual behaviors) should be conducted to assess for triggers, and strategies for redirection and maintaining the dignity of the person with dementia should become part of the care plan.

Resource:

[https://www.uclahealth.org/dementia/sexually-inappropriate-behaviors](https://www.uclahealth.org/dementia/sexually-inappropriate-behaviors)
Ten Best Practices: Reducing Use of Antipsychotic Medications in individuals with Dementia

Inappropriate use of antipsychotics to manage behavioral disturbances in individuals with dementia is prevalent. Antipsychotic use is associated with serious risks such as strokes, death, falls and fractures, and accelerated cognitive decline. Antipsychotics also may severely impair daytime functioning and quality of life by causing sedation, tiredness, unsteady gait and swallowing problems.

Best Practices:

1. Spend time to get facts and diagnoses correct. Use the DICE/STAR model
   a. 78-year-old Japanese widowed male with COPD, DM, insomnia, “anxiety attacks and paranoia” at night. Etiology: hypoxia, hypoglycemia, beta-agonist inhalers, wants to reconcile with his estranged son
   b. 70-year-old with diagnosis of “Parkinson’s disease (PD)”, Dementia, “Schizophrenia” on Carbidopa-levodopa and pramipexole, olanzapine and sertraline. PD psychosis misdiagnosed as schizophrenia or drug-induced parkinsonism misdiagnosed as PD.
   c. Alcohol, marijuana, delirium and polypharmacy induced psychosis common
2. Geriatric Scalpel (rational deprescribing) and Medical Deintensification (includes DNH)
   a. 80-years old Caucasian widowed female in advanced stages of dementia, erratic eating, severe agitation, visual hallucinations. Rx: reduce and discontinue insulin, discontinue donepezil and memantine, discontinue metformin, reduce anticholinergic burden of meds, palliative care
3. STEPS (Support system [family, staff]) Training Empowerment Praise and Support in relationship building communication, environmental initiatives, creative engagement and other aspects of SPPEICE, and Snoezelen.
4. SPPEICE – strengths-based personalized psychosocial environmental initiatives and creative engagement) – includes comprehensive exercise program
5. ATMAN approach to managing pain effectively without opioids and tramadol
6. Judicious short-term non-antipsychotic psychotropic medication trial if symptoms severe persistent: Citalopram, Trazodone, Pimavanserin, Dextromethorphan-quinidine
7. Collaborative care using telemedicine (collaborate with PCP and pharmacist)
8. Recreational therapist consultation routine
9. Music therapist involvement routine
10. Fulltime advanced practice nurse or physician assistant supervised by a geriatric psychiatrist rounding frequently
Alternatives to Antipsychotics for Management of Agitation in Individuals with Dementia

Before prescribing medications, please ensure the following two steps:

1. A thorough assessment-work up and treatment of reversible factors for agitation. Examples of these factors: medication adverse effects, pain, infection, dehydration, constipation, nutritional deficiencies, caregiver depression.

2. Psychosocial spiritual interventions. Examples: family/staff education on DICE* approach, caregiver respite, recreational therapist guided strengths-based personalized activity schedule involving meaningful and or fun activities throughout the day, exercise therapy / walking, music, lavender (or alternative lotion) lotion three times daily and as needed, pets, adult day program, community care program (SW Area Agency on Aging).

<table>
<thead>
<tr>
<th>Medication</th>
<th>Clinical tips</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chamomile extract 500 mg three times daily</td>
<td>May help with anxiety, safer than other prescription medications</td>
</tr>
<tr>
<td>Citalopram 10mg daily, may increase to 20mg after a week</td>
<td>May help with agitation, anxiety, irritability, physical aggression and even psychotic symptoms; takes 4-8 weeks, even longer sometimes to work; carries all the risks and benefits of SSRIs</td>
</tr>
<tr>
<td>Escitalopram 5mg daily, may increase to 10mg after a week</td>
<td>Has less QTc prolongation than citalopram, otherwise benefits and risks similar to citalopram</td>
</tr>
<tr>
<td>Sertraline 25mg daily, may increase to 50mg after a week</td>
<td>Preferred over citalopram and escitalopram if there are concerns about QTc prolongation, otherwise benefits and risks similar to citalopram and escitalopram</td>
</tr>
<tr>
<td>Suvorexant 5mg, may increase to 10mg</td>
<td>Approved by the FDA for insomnia Rx in AD</td>
</tr>
<tr>
<td>Mirtazapine 7.5mg at bedtime, may increase to 15mg</td>
<td>May be considered for treatment of insomnia due to major depression causing agitation</td>
</tr>
<tr>
<td>Trazodone 25mg at bedtime for insomnia</td>
<td>Risk of falls, orthostasis, tiredness in daytime</td>
</tr>
<tr>
<td>Melatonin 0.5-3mg at bedtime for insomnia</td>
<td>May also help REM sleep behavior disorder</td>
</tr>
</tbody>
</table>

*DICE (Describe Investigate Create Evaluate) – involves training family / staff in being a detective, seeing agitation as expression of unmet need, addressing the need and evaluating response to intervention: https://diceapproach.com

Resource:
Citalopram for Treatment of Agitation in persons with Dementia

Antipsychotics are often used for management of agitation in persons with dementia. Antipsychotics carry serious risks and have very modest benefits and should be reserved for situations when there is severe danger to self or others due to agitation and nonpharmacological interventions have not been effective and there are no reversible medical conditions that are causing the agitation.

In my opinion, citalopram can and should be used before antipsychotics in majority of the situations where antipsychotics are being given to treat agitation or psychosis in patients with dementia. Citalopram is eminently safer than antipsychotics, even at 30mg dose (the FDA guidelines suggest avoiding more than 20mg in older adults due to risk of prolonged QTc interval).

The following are the three key research studies that support my recommendation.

1. CitAD study: A 9-week randomized placebo-controlled study (published in JAMA, Feb 19, 2014) found citalopram (30mg) did significantly better than placebo in reducing agitation and caregiver distress in patients with Alzheimer’s dementia.

2. Citalopram vs risperidone: A 12-week double blind controlled study did not find any statistical difference between citalopram and risperidone for the treatment of agitation or psychotic symptoms in patients with dementia. Agitation and psychotic symptoms decreased in both groups (published in American Journal of Geriatric Psychiatry 2007).

3. Citalopram vs perphenazine (an older antipsychotic) vs placebo: This was a 17-day double-blind placebo-controlled study in hospitalized patients. Citalopram did better than placebo in reducing agitation in dementia patients. Perphenazine also did better than placebo but citalopram did better than perphenazine (published in American Journal of Psychiatry 2002).

In all patients with dementia where antipsychotics are being considered to treat severe persistent agitation, I recommend we use citalopram first. The benefits do take time (compared to antipsychotics) but risks are way less. Antipsychotics carry risk of stroke, premature death, aspiration pneumonia, increased risk of falls and injury, increased risk of hospitalization, antipsychotic-induced parkinsonism, sedation, dysphagia and other serious risks.

Check EKG whenever feasible to look for prolonged QTc interval. In majority of cases, you will be able to safely start citalopram.

Start at 10mg daily for Dementia with behavioral disturbances, increase it after a week to 20mg daily and monitor for response to treatment and adverse effects. In some cases, I have increased it further to 30mg and found benefits without significant adverse effects and I have felt that benefits outweigh risks in all cases where the response has been significant.
Note: Citalopram does increase QTc but this increase has not been linked to cardiac outcomes. FDA needs to revisit this safety advisory (McCarrell et al. Mental Health Clinician 2019; Texas Tech University School of Pharmacy).
Staging of Alzheimer’s Disease Dementia

Name of the patient: [Missing]

<table>
<thead>
<tr>
<th>MOCA score (date)</th>
<th>FAQ score (date)</th>
<th>Katz index (date)</th>
<th>FAST* Stage (clinical judgment)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-30</td>
<td>0-30</td>
<td>0-6</td>
<td></td>
</tr>
</tbody>
</table>

MOCA: Montreal Cognitive Assessment measures cognitive function (high score = high function).

SLUMS: Saint Louis University Mental Status (recommended if one is not certified in MOCA administration, certification [online training offered at their website] for MOCA costs $125). SLUMS is not as well researched as MOCA but the score of MOCA similar to SLUMS.

FAQ: Functional activities questionnaire measures instrumental activities of daily living (IADL) (high score = low functioning)

Katz index: Measures basic activities of daily living (BADL) (high score = high functioning)

4 stages (brief description):

Mild cognitive impairment: Decreased performance in demanding employment and social settings, decreased job function evident to co-workers, difficulty traveling to new locations, decreased organizational capacity. Can pay bills, manage dinner for guests, manage finances, travel to familiar places. MOCA generally 23-25 but not normal (26 and above). FAQ low. Katz index: 6

Mild dementia: Difficulty paying bills, managing dinner for guests, managing finances, traveling to familiar places. Can recall their own address and telephone number, know where they are and what day it is. MOCA generally between 18-22. FAQ generally six or higher. Katz index 6.

Moderate dementia: unable to recall their address and telephone number, get confused about where they are and what day it is, can dress themselves but may wear the same clothes unless supervised, requires assistance in wearing proper clothing for the weather or event. Can remember significant details about themselves and their family. MOCA generally between 10-18. FAQ 12 or higher. Katz index 4-5

Severe dementia: can remember their own name but difficulty with names of close family members, can recognize familiar faces even if they have trouble naming the person or identifying their relationship. Need help dressing properly, have major changes in sleep patterns (e.g., day night reversal), trouble controlling bladder and or bowels. MOCA generally below 10, FAQ close to 30; Katz index 0-3

*FAST dementia staging: Functional Assessment Staging Test

AAD and stages of Dementia: AAD increases in prevalence with advancing of dementia. Palliative care and hospice may be appropriate for agitation in patients with severe dementia.
Age-Friendly Hospital System (AFHS)

4Ms of AFHS: 4Ms framework can be used to provide comprehensive and person-centered delirium care to older adults

- What Matters (aligning intensity of AAD management with patient goals, engaging family and friends in AAD care)
- Medications (especially avoiding deliriogenic medications, rational deprescribing)
- Mentation (routine depression, delirium and trauma screening across all settings)
- Mobility (avoid out of bed with assist, minimize catheters and restraints)

Institute for Healthcare Improvement: https://www.ihi.org/initiatives/age-friendly-health-systems/resources-and-news

Namaste
A stab went through her
so sharp that my patient
sobbed and clutched
at her belly.
“Just sleep,
tomorrow will bring good news,“ she kept repeating
like a mantra,
coordinating the words
with her breath.
She slept fitfully.
When she woke,
pale light of morning
was slanting
through her window.
She felt
as sick and achy
as if she had
not slept at all.
Her tummy
was still tied
in a knot.
“Just breathe slowly,
today will bring good news,”
She kept repeating
like a mantra.

🙏
This is my patient’s daughter. My 69-year-old patient with mild dementia was in the ICU for cardiac arrhythmia. She recovered beautifully to the surprise of all of us and was discharged home. I was teaching the daughter mantra meditation and mindful breathing.

Many individuals with dementia come out of the ICU in more advanced stage due to brain injury caused by underlying medical issues that required ICU plus trauma of being in a hospital and ICU - especially if the hospital doesn’t have HELP program and dementia friendly hospital practices. They often need to be discharged to a nursing home rather than home and may never return home 😞

HELP - Hospital Elder Life Program: For Delirium prevention through the American Geriatrics Society

Top 5 dementia friendly hospital practices:
• routine consult with dementia team - dementia physician / nurse practitioner expert (e.g., consultant psychiatrist) plus at least one of the following: chaplain, nurse, social worker, neuropsychologist, occupational therapist (addressing ageism and dementia-ism being a top priority)
• one to one with a trained staff
• supportive family educated and involved as much as the family can
• individualized strengths based routine bio-psychosocial spiritual culturally appropriate care plan that includes early mobilization with intensive physical therapy, continuous daily meaningful activities schedule and night time sleep enhancement
• close collaboration with Hospitalist team and HELP program team to prevent delirium, falls, pain, constipation, dehydration, de-conditioning and frailty
AAD in hospitals / Dementia Friendly Hospital: 10 Key Interventions to Reduce Stress of Hospitalization for Patients with Dementia

- Ambulate them frequently (preferably a few minutes every hour)
- Take positive risks and take persons with dementia outdoors, especially to spend time in natural settings. Get them out of the hospital / facility (e.g., to a garden or a labyrinth).
- Keep soothing instrumental background music (preferably personalized) in the room
- Use robotic pets when appropriate (e.g., for patients with advanced dementia who may not be able to recognize that these are robotic pets)
- Have (warm) fidget blankets available to calm them if they are restless
- Use pictures that have been identified as being of interest to the patient to help distract them during personal care or when they are agitated.
- Sing to the patients during personal care to ease their stress
- Encourage family members, friends, volunteers and chaplains to visit them daily (and preferably stagger the visits so that, ideally, only one person is visiting at a time) in person or virtually (e.g., skype)
- Engage them in continuous pleasant activities (ideally, create an individualized strengths-based daily pleasant activity program)
- Frequently (at least three times a day) provide soothing hand, neck and back massages with soothing lotions (e.g., lavender lotion).
Delirium Care Score*

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<thead>
<tr>
<th>Category</th>
<th>Criteria</th>
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<tbody>
<tr>
<td>Physical activity</td>
<td>Intensive physical therapy</td>
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<tr>
<td></td>
<td>Regular physical therapy</td>
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</tr>
<tr>
<td></td>
<td>Walking 1000 steps daily</td>
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<tr>
<td></td>
<td>Less than 1000 steps daily</td>
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<tr>
<td>Occupational therapy (OT)</td>
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<td></td>
<td>Regular OT</td>
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<td></td>
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<td>Sleep</td>
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<td>Less than 5 hours at night</td>
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<td>Intravenous lines</td>
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<td></td>
<td>Still in</td>
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<tr>
<td>Foley</td>
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<tr>
<td></td>
<td>Still in</td>
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<tr>
<td>Social interactions – family,</td>
<td>3 or more hours a day</td>
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</tr>
<tr>
<td>volunteers, chaplain, other</td>
<td>1-3 hours</td>
<td>1</td>
</tr>
<tr>
<td>staff</td>
<td>Less than one hour</td>
<td>0</td>
</tr>
<tr>
<td>Sunlight / bright light</td>
<td>2 or more hours a day</td>
<td>2</td>
</tr>
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<td></td>
<td>1-2 hours a day</td>
<td>1</td>
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<td></td>
<td>Less than one hour a day</td>
<td>0</td>
</tr>
<tr>
<td>Meaningful strengths-based</td>
<td>2 or more hours a day</td>
<td>2</td>
</tr>
<tr>
<td>activity schedule</td>
<td>1-2 hours a day</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Less than one hour a day</td>
<td>0</td>
</tr>
</tbody>
</table>

*Created by the author – evidence-informed. It has not been formally researched. Max score 21; Shoot for 21 for every patient. The better the score, the more likely one could prevent delirium. Hospitalized patients with dementia are at very high risk of delirium. Delirium increases risk of death, length of hospital stay, decline in functioning requiring discharge to a nursing home and low likelihood of returning back home.

Inspired by McCance Brain Care Score, McCance Center for Brain Health, MGH [https://www.massgeneral.org/neurology/mccance-center](https://www.massgeneral.org/neurology/mccance-center)

References and Resources

Personalized Multicomponent Nondrug Rx for Delirium Prevention

• Collaborative de-escalation
• Early mobilization with intensive PT / OT
• Ensure adequate hydration and nutrition
• Remove restraints, in-dwelling catheters, IV as soon as is feasible
• Hearing and vision aids
• Calm, low stimulation environment
• Lavender or other soothing lotion to arms four times a day and as needed
• Soothing personalized music in the background
• Assisted walking three times a day
• Bright light and or sunlight 20 minutes a day
• Family engaging patient in meaningful activities
• Gentle care*

*American College of Surgeons Best Practices for Geriatric Trauma 2023
Resources

To assess for medical conditions causing agitation including medication-induced agitation:

Saint Alphonsus Memory Center
Boise Nampa Eagle
Telehealth appointment available
No referral needed. Self-referral accepted.
208-302-5400 Boise
208-302-6000 Nampa Eagle
Free eBook titled *Improving Brain Health* available at the website. Links to podcasts on dementia topics available at the website
[https://www.saintalphonsus.org/specialty/memory-center/](https://www.saintalphonsus.org/specialty/memory-center/)

Primary care provider of the person with dementia can also do this assessment.

**Individual counseling (for family caregiver) and guidance regarding behavior management strategies**
Laura LaForte LCSW
206-953-3708
[https://www.lafortetherapy.com/](https://www.lafortetherapy.com/)

Connection Club Adult Daycare
Dick Eardley Boise senior center
208-608-7578

**Dementia and Creativity Sessions**
Erika Shaver-Nelson, MA
208-724-4754

**Family education on Behavior Management Strategies**
Family Caregiver Alliance Website: Behavior Management Strategies
High quality videos and quick reads: [https://www.caregiver.org/caregiver-resources/caring-for-another/behavior-management-strategies/?via=caregiver-resources,caring-for-another](https://www.caregiver.org/caregiver-resources/caring-for-another/behavior-management-strategies/?via=caregiver-resources,caring-for-another)

**Telephone support and guidance of community resources**
- 988 suicide and crisis line
- 211 Idaho care line
- 800-898-9626 Alzheimer Association 24/7 helpline

**Case Management Respite Education regarding Dementia**
Community Care Program
Regarding dementia
208-898-9626
Family caregiver wellness and resilience education and training
Powerful tools for Caregivers workshop
https://www.agingstrong.org/powerful-tools-for-caregivers
208-947-4283

Family support and respite
Family Caregiver Navigator
https://caregivernavigator.org/
208-426-5899

Family assistance for community resources (e.g., transportation)
Saint Alphonsus Community Health Worker program
208-367-4482

Websites

The DICE approach online training program. https://diceapproach.com/

International cognitive therapy stimulation (CST) center
Saint Louis University’s Geriatric Education Program offers CST
https://www.slu.edu/medicine/internal-medicine/geriatric-medicine/aging-successfully/cognitive-stimulation-therapy.php

Music and Memory Program. https://musicandmemory.org/

https://healthandwelfare.idaho.gov/dementia-friends

Community of Dementia Education. http://community.abcdementia.org/

Book for patients and their families
3. Agitation associated with Dementia management. eBook by Abhilash Desai MD. Available upon email request.
Other resources

1. Movement for Memory program at Tomlinson South Meridian YMCA (group exercise program for individuals with MCI): (208)-344-5502. Ext 524.
2. Blue Mood Senior Counseling – Teletherapy: https://bluemoonseniorcounseling.com
3. Grannie on the Move: (208)-288-2220
4. Area Agency on Aging: (208) 898-7060
6. Anticholinergic burden calculator. ACB calculator. Provides anticholinergic burden for many medications using 1-3 points, 3 being most anticholinergic. Reducing anticholinergic load of medications is a key intervention in prevention and treatment of AAD. https://www.acbcalc.com/
Resources for Dementia Training

**Free**
1. Health Resources and Service Administration HRSA Dementia modules (free): [https://bhw.hrsa.gov/alzheimers-dementia-training](https://bhw.hrsa.gov/alzheimers-dementia-training)
4. REACH certified caregiver support coaches. [https://www.caregiver.va.gov/REACH_VA_Program.asp](https://www.caregiver.va.gov/REACH_VA_Program.asp)

**Not Free: Certification available**
3. National Certification Board for Alzheimer’s Care [https://ncbac.net](https://ncbac.net)
5. Dementia Care Specialists. Crisis Prevention Institute [https://www.crisisprevention.com/Our-Programs/Dementia-Care-Specialists](https://www.crisisprevention.com/Our-Programs/Dementia-Care-Specialists)

**STAR-C VA Manual for Challenging Behavioral Management Strategies**
Dementia Bibliotherapy: Books recommended to improve lives of individuals with Alzheimer’s and other dementias and their family members, friends and carepartners

**Mayo Clinic on Alzheimer’s Disease and Other Dementias.** Jonathan Graff-Radford MD, Angela Lunde MA. 2020. Mayo Clinic Press. Excellent first book for individuals with mild memory problems who are worried that they may have dementia and family members of individuals with dementia. High-quality, relatively easy to understand information of complex topics as well as many practical tips and guidance to get early accurate diagnosis and live well.

**Day-to-Day: Living with Dementia: A Mayo Clinic guide for offering care and support.** Angela M. Lunde, M.A., offers essential caregiving guidance, including practical tips and resources, techniques for working through difficult emotions, and strategies for managing common dementia-related challenges.

**STAR-C VA Manual for Challenging Behavioral Management Strategies**

**Navigating Dementia: A workbook for family caregivers.** Free. Excellent resource for all caregivers, especially those living in Idaho.

**When your aging parent needs help: A geriatrician’s step-by-step guide to memory loss, resistance, safety worries, and more.** Leslie Kernisan, MD, MPH and Paula Spencer Scott. 2021

**Travelers to Unimaginable Lands.** Dasha Kiper. One of the best books for family caregivers to find deeper understanding about why caregiving and doing the right thing is SO HARD.

**Living with Memory Loss: A Basic Guide** by University of Washington Memory and Brain Wellness Center. This essential resource is downloadable (for free) from their website under the Resources tab, either the full handbook, or separate chapters. Excellent for patients with mild to moderate dementia and their family members.

**Before I Forget: Love, Hope, and Acceptance In Out Fight Against Alzheimer’s.** B. Smith & Dan Gasby. Excellent resource to promote positive approach to living with dementia.

**The Alzheimer’s Solution: A breakthrough program to prevent and reverse the symptoms of cognitive decline at every age.** Dean & Ayesha Sherzai, M.D. Codirectors of the Brain Health and
Alzheimer’s Prevention Program at Loma Linda University Medical Center. Useful for patients with mild dementia who wish to slow down cognitive decline.

*The Spectrum of Hope: An optimistic and new approach to Alzheimer’s Disease and other Dementias* 2017, by Gayatri Devi MD, neurologist and the Director of the New York Memory and Healthy Aging Services. Many cases with remarkably positive outcomes are discussed. Excellent for patients with mild dementia and their family members to learn about ways to slow cognitive decline.

*Seven Steps to Manage Your Memory* by Andrew Budson and Maureen O’Connor. Oxford University Press, 2017. Excellent resource to differentiate normal memory loss from memory decline due to dementia and many cases with positive outcomes are shared. Excellent for patients with mild dementia.


*Dancing with Elephants: Mindfulness Training for Those Living with Dementia, Chronic Illness, or an Aging Brain (How to Die Smiling)*. Jarem Sawatsky. 2017. Toronto, ON: Red Canoe Press.


With the End in Mind: Dying, Death and Wisdom in the Age of Denial. Kathryn Mannix


Dementia Medical Poems. Abhilash Desai. These poems are not for the faint of heart. I am happy to send this by email upon request.


One Day Mindfulness Millionaire: Living Mindfully – A light-hearted primer for the uninitiated. Abhilash Desai MD and Faith Galliano Desai Ph.D. eBook available at https://store.bookbaby.com. Excellent resource (my unbiased view LOL) to begin to develop skills to live mindfully and develop a simple daily meditation practice. Mindful living and meditation can help both the person with dementia and their family / friends live with less stress and more joy.

Namaste
Arts on Prescription for Dementia Care: Resources

Book

Websites
1. TimeSlips (https://www.timeslips.org) creative storytelling. Online training course available. At least one long-term care staff should be trained in TimeSlips
2. Center for Arts in Medicine. https://arts.ufl.edu/sites/creating-healthy-communities/resources/arts-on-prescription-a-field-guide-for-us-communities/
Dr. Desai Books, book chapters and Podcasts

   a. Chapter 3: Dementias / Neurocognitive disorders
   b. Chapter 12: Psychiatric aspects of rational deprescribing
   c. Chapter 13: Psychosocial spiritual wellness care plan
3. *Overcoming Anxiety, Depression and Anger.* Non-pharmacological interventions to manage stress, anxiety, depression and anger. eBook available upon email request.
4. *Mindfulness and Meditation.* eBook available upon email request.
5. *Fearless Strong Patient Kind - Short meditations to manage stress and become resilient.* eBook available upon email request.
7. *Delirium-related agitation in hospitalized older adults.* eBook available upon email request.
8. *Dementia Prevention.* eBook available upon email request.
Dementia Physician Experts in Idaho

**Eastern Idaho**
Scott Christensen MD, Geriatrician
Intermountain Medical Clinic
1951 Bench Road, Suite B Pocatello, ID 83201
Phone: (208) 238-1000
Dr. Christensen can do telemedicine visits

**Northern Idaho**
Susan Melchiore MD, Geriatrician
One Site for Seniors
296 W. Sunset Ave, #14-15
CDA, ID 83815
Phone: 208-967-4771
Dr. Melchiore can do telemedicine visits

**Central Idaho**
Jennifer Lewis MD, Geriatrician
211 Forest Street McCall Idaho.
Phone: 208-634-2225
Dr. Lewis can do telemedicine visits.

**Southwest Idaho**
Kara Kuntz MD, Geriatrician and team
Saint Alphonsus Memory Center, Boise, Eagle, Nampa
Phone: 208-302-5400; 208-302-6000
Memory Center team can do telemedicine visits
**Supporting Caregiver Wellbeing: Key to Prevention and Reduction of AAD**

If caregivers are well, they are able to interact with the person with dementia in positive ways that prevent and reduce AAD. Healthcare professionals can provide education and skills training in the following areas to promote caregiver wellness.

1. **Mindfulness:** It is important for family and friend caregivers to approach their loved one with dementia in a mindful way. Mindfulness means that the caregiver is paying attention to the loved one non-judgmentally. Mindfulness is a journey and memory of past negative interactions with the person with dementia and worries about the future make it very difficult for the caregiver to be in the present with the loved one. Non-judgmental presence by the caregiver helps the person with dementia to feel secure and valued. Being present also allows the caregiver to become aware of numerous small positive interactions and experiences with the loved one that they can cherish. I encourage family members to engage in mindfulness training.

2. **Validation:** It is important for the caregivers to validate the experiences and feelings of the person with dementia. Authentic (sincere) validation helps the loved one recognize that they are being heard and understood and not ignored. Caregiver burnout is the most common reason for failures on the part of the caregivers to provide validation. I encourage caregivers to track their own level of burnout and take steps to prevent it and address it head on.

3. **Trust:** It is important for caregivers to trust the perspectives of the person with dementia. Persons with dementia have brilliant perspectives and all of us (not just family caregivers) need to recognize that and allow them to lead us as often as being led in their journey to live well with dementia. Negative cultural stereotypes (e.g., persons with dementia don’t know what is going on so there is no reason to listen to them) is the key reason caregivers (both family and professional) fail to inquire their point of view or learn to listen to and see the creative ways the person with dementia is navigating challenges and seeking happiness.

4. **Embracing imperfection:** There are no road maps, only general guidelines that other caregivers and healthcare professionals can provide to the caregivers. Caregiving is complex, ever changing in its rules, and impossible to perfect. Caregivers need to be kind to themselves and not let their “numerous mistakes” and “frequent errors” get in the way of living and caring for self and loved one with kindness and forgiveness.

5. **Sense of humor:** It is not uncommon for the caregiver’s sense of humor to be the first casualty of stress related to caring for their loved one with dementia. It is crucial for the caregiver to appreciate the tremendous value a healthy sense of humor has in living well with dementia. Healthy sense of humor is the strongest expression of spiritually defeating dementia.
Are you (like us) a caregiver struggling with the responsibilities and stress of caregiving? Have you wondered how on earth do some caregivers find the role of caregiving a blessing? Are you amazed at how some caregivers tend to their responsibilities effortlessly and continue to be cheerful day in and day out? Yes, these remarkable individuals do exist, and, we have had the good fortune to have met them in our professional life (as a psychiatrist [AKD] treating patients from ages 5 and up, and child and adolescent psychologist [FDG]) as well as in our personal life. We have been looking for a path towards becoming better caregivers for some time now.

This long essay is aimed at helping caregivers (family [e.g., parents, siblings] and professional [e.g., staff in nursing homes]) of persons having dementia (PHD) improve their capacity for compassion, creativity during caregiving, effectiveness of their efforts, and at the same time improve caregiver’s own sense of wellbeing. Caregivers work hard to improve quality of life of PHDs but often experience frustration, helplessness, hopelessness, stress, and burnout. We have discovered that an approach that blends mindfulness with the concept of care-partnering can not only improve our skills in improving wellness and positive experiences of PHDs but also prevent burnout and promote our emotional and spiritual growth. For many caregivers, mindful care-partnering comes naturally, without any knowledge or training. For the rest of us, focus and effort in developing skills of mindful care-partnering are necessary.

Mindfulness is bringing one’s awareness to the present moment, with intention, to whatever one is experiencing (e.g., feelings, sensations, thoughts) with an attitude of kindness and curiosity. It is a skill and as such, can be developed much more easily than generally recognized. Mindfulness is an excellent way to become familiar with the workings on one’s mind and minds of others (especially of PHD). Intimate familiarity with the workings of one’s own mind and that of others opens up doors of healing and wellness that have otherwise remained obstinately closed.

Mindfulness is paying attention in a particular way – on purpose, in the present moment, and nonjudgmentally.
- Jon Kabat Zinn

Care-partnering includes all traditional aspects of caring for another person (e.g., helping them with daily activities) but also involves awareness of the PHD’s capacity to partner and collaborate with caregivers and the unique gifts caregivers receive from PHDs as well as from the very act of caring. Care-partnering also goes a step further in recognizing that the PHD is the expert in how best to have her/his needs met and thus, it is crucial to be led by the PHD at least as often as leading the PHD towards safety, better ways of adapting to life’s challenges and positive experiences.

Henceforth, all caregivers will be called care-partners.
Mindful Care-Partnering
It is easier to understand mindful care-partnering by understanding three common aspects of unmindful care-partnering. These are: autopilot, auto-reaction and premature cognitive commitments.

Autopilot
As you (the care-partner) embark on mindful care-partnering, you will quickly notice that your mind wanders to many different places/situations as you go through the task at hand (say for example, taking a walk with the PHD). It is typical for our minds to jump from paying attention to the task at hand to having imaginary conversations, plans, worries, replaying memories, and a host of other thoughts. One can say that our mind keeps wandering into the land of thoughts and thinking. You may finish the walk and may not be able to remember what areas of the park you walked and what conversation you have had with the PHD. This happens automatically because it is a natural tendency of our brain to go on autopilot. Autopilot requires much less energy and intentional activity takes up much more energy so given a choice, brain will always switch to autopilot to conserve energy (a strategy that improved survival of our hunter-gatherer ancestors). Mindful care-partnering involves gently bringing your wandering mind to the present moment, becoming aware that you are walking with the PHD, that you are holding their hand, and becoming aware of the green landscape around you. Mindfulness in such situations involves catching oneself having these thoughts, minimizing or withholding any negative judgment or criticism for being distracted and intentionally bringing our focus back to the present. You can be certain that your mind will wander again, and you can once again bring your mind gently to the present. Being present with the PHD in such a way will improve sense of value the PHD has for herself/himself and improve PHD’s sense of connectedness with you. Improved sense of one’s value and sense of connectedness are two most basic requirements for any significant and sustained healing and wellness to occur.

The faculty of voluntarily bringing back a wandering attention over and over again is the very root of judgment, character, and will. No one is compos sui if he have it not. An education which should improve this faculty would be the education par excellence.
– William James (Principles of Psychology 1950; p.424) Note: Compos Sui meanings master of oneself

Auto-reaction
As you become aware of your emotions during care-partnering, you will notice that you often react automatically from emotion first rather than pause, reflect and then respond. Once again, the brain under strong emotions (especially negative emotions such as fear or anger) will tend to respond immediately based on perceived threat. Brain’s initially perception of threat involves a part of the brain called amygdala and amygdala is biased towards over-reacting (guided by the goal of increasing survival). By being mindful of our brain’s (specifically amygdala’s) tendency to over-react, you can develop skills to inhibit the immediate auto-reaction, pause and reflect taking in any additional information our senses are registering and contrasting the event with any past experience with similar events we may have, attempting to see from the point of view of the PHD, and then responding. Your pre-frontal lobe plays a key
role in mediating all these actions (inhibiting auto-reaction, pausing, reflecting, responding). Depending on the situation, all these actions may take only a few seconds or several minutes or even longer.

Premature Cognitive Commitments (PCC)

Our brains are programmed to come to quick conclusions based on minimal information (because such function provided survival advantage). Such conclusions are key examples of premature cognitive commitments (PCCs) (a term coined by Dr. Ellen Langer, psychologist in her book *Mindfulness*). Judgmental thinking is typically based on PCCs. This tendency of our brain is valuable in emergency situations but can be counter-productive and at times even destructive in day-to-day situations involving caring. As you become more aware of your thoughts and feelings during caring, you will be able to catch yourself in such acts of prematurely committing to an understanding of what is going on (especially challenging behaviors). For example, the PHD is having an anger outburst and you quickly decide that this is “manipulative, attention seeking behavior” or that the PHD is “just being mean”. Many PCCs involve harsh judgment about ourselves (e.g., “I am a terrible caregiver”) or catastrophic predictions (e.g., “this resident with dementia is going to tear down the place”). PCC will push you to react as if this conclusion was absolutely correct and there was no alternative explanation for the behavior or alternative future scenarios (especially positive ones). PCC will further increase intensity of negative emotions that triggered them in the first place. Auto-reaction and PCCs by care-partners often make the behavior of the PHD worse. Mindful care-partnering will slowly but surely help you become aware of how often our PCCs are incorrect. As you learn to pause, take in the context, get to know the narrative of the PHD (how the PHD got to this point), your understanding of challenging behaviors will become more nuanced, what needs to be done to address the challenging situation will become clearer, and your success in addressing the challenging behavior will greatly improve.

Benefits of Mindful Care-Partnering

As you delve into this practice of mindful care-partnering, you will become aware of your fears, expectations, and judgments and how those color how you relate to the PHD. Mindful care-partnering will also help you see the situation from PHD’s point of view. Mindful care-partnering is all about honoring relationships, connectedness, and a sense of belonging. Your relationship with PHD may feel chaotic or different from the ideal you want. You may find yourself drifting through life on “autopilot,” constantly worrying about the future or feeling upset (guilty, resentful) about the past instead of living in the here and now. This method of caring teaches you to stay “in the present” and stay connected with the PHD as well as yourself in kinder and softer ways.

Regular meditation practice (e.g., 30-minute Breath Awareness Meditation early in the morning before starting one’s day; at least a few minutes of Breath Awareness Meditation when experiencing strong negative emotions [e.g., anger]) and mindfulness practices throughout the day (e.g., mindful eating, mindful walking) has been shown to dramatically improve one’s sense of wellbeing and inner strength in just eight weeks of daily practice. Yoga, Tai Chi and Qi Gong are other mindfulness-based practices (often called mind-body practices) that promote

69
psychological and spiritual wellness and make us more effective care-partners. Meditation and mindfulness practices increase our emotional and spiritual resilience (emotional and spiritual bank-balance) and lower our risk for burnout, depression and persistent anxiety. Every time we engage in meditation and mindfulness practices, we are depositing money in our emotional and spiritual bank! As we build this bank balance, hassles of daily life will seem mere annoyances rather than “terrible” events, and we will bounce back from traumatic events faster and without becoming demoralized. We will be less likely to dwell in self-pity or resentment and more likely to be aware of many things we can still be grateful for. We will find improved capacity to be patient, to see silver lining in clouds and make lemonade from lemons that life gives us (again and again). We will be able to make ourselves do things we don’t want to do (e.g., regularly exercising, eating healthy foods, having a good sleep routine, taking time to enjoy simple things in life [HYGGE {Danish art of contentment, comfort and connection}]) and we will also be able to more easily restrain ourselves from doing things we should not (e.g., putting self or others down under the influence of strong negative emotions). Meditation and mindfulness practices will help us laugh at our foibles with light-hearted spontaneity and avoid taking life too seriously. We will be able to let go of materialistic attachments and move towards a simpler and easier life where awareness of being alive and surrounded by love will be enough reason to be happy.

**From Knowledge to Skills to Practice**

Mindful care-partnering challenges you to examine your expectations. Your fears may motivate you to try and direct the PHD in certain ways and often will blind you to what is truly in their best interests. PHDs have their own trajectories, beliefs, ideas, and opinions, often different from ours, and what we think theirs should be. When you challenge yourself to pay attention to the present moment and see caring as collaboration, an artistic endeavor, you gain opportunities to truly see PHDs, to understand and appreciate them, and solve problems with them rather than for them.

Everyone has times where inner musings (inner chatter, conversation with oneself in our minds) take over, but this means we lose opportunities to connect with the PHD by being blind to the present moment. The key is in paying attention to the small things, like a child, fully embodying every moment, as opposed to life being a movie in our minds or trying to match it to our expectations.

You can think of mindful care-partnering as an opportunity to get to know yourself better and to accelerate your own psychological and spiritual growth. A moment of mindful care-partnering may allow you to catch yourself and realize an opportunity to change direction. When you pay attention to the present you may recognize that some thoughts or fears are not based in reality and liberate yourself to a more productive course of action.

As with most skills, mindful care-partnering takes time and practice. Even small changes are profoundly healing and transformative. Some examples of small steps are being aware of your breath, the feeling of your hands, or your own body language, the feel of PHD’s hand in yours. The simplest method of becoming mindful is to get connected with our senses. Bring your
awareness to your breathing, for example, in the belly or the nostrils, to focus on one thing at one time. Practicing this at a moment of tranquility, and drawing on this experience when a conflict happens, will help you respond mindfully to the conflict instead of reacting emotionally, and find better ways of dealing with it.

Instead of preparing for situations in the future, or using a cookbook approach, mindful care-partnering asks that you trust yourself to be aware of the next moment and find a solution when that moment comes. The goal is not to eliminate chaos and negative feelings; these are a natural part of life filled with caring responsibilities. The goal in to be kind and curious when those moments arise.

It is crucial to give yourself credit for the little moments of success and accept yourself and others with all the imperfections. Choosing to engage in mindful care-partnering is empowering yourself to be present for the challenges and successes.

Source:
Sheer Terror

I was mumbling in indecipherable shrinkspeak as I tried to crawl out of a state of stunned disbelief. My thoughts a mush of rampant flashes. My inner debates raging over strategies that changed by the minute. Tears started to roll out. Not tears of sorrow, but of sheer terror. My stomach flipped I wanted to throw up. My body shook to toes as my heart raced away. Where are you mom? Where have you wandered off to?

🙏
The plight of a psychiatrist caregiver whose mother (my patient) has dementia and had wandered away from home. After 2 longest hours of daughter’s sheer terror, my patient was found, safe and sound, several blocks away.

Wandering is a common AAD and Alzheimer’s Association website as excellent strategies for prevention and management. https://www.alz.org/help-support/caregiving/stages-behaviors/deambulacion
Unthinkable Future

“Sorry. I am being rude, talking about you in front of you.”
My patient with dementia was pacing.
She plunked herself into the chair and tucked her hair behind her ears.
Husband’s health was declining rapidly.
He had no words.
His soul felt too heavy.
His wife in an ALF?
That future was unthinkable at this moment, and untenable when it arrived.

His body revealed nothing of his psychic wounds.
Wounds that ran deep and felt permanent.
“Ah Assisted Living Facility, a place of boundless tranquility” he quipped.
“Humor is a good start. But we need more than a strong heart. No one can predict the future.
Let’s live in the possibility that there is an alternate destiny.
Let’s focus on getting you back to healthy.”

🙏
ALF = assisted living facility / community
“You back to healthy” = getting the caregiver husband back to healthy – less stressed, less worried, positive and hopeful as much as is possible in the given context.

His wife was having lot of agitation at home, not sleeping, asking the same questions again and again and at times, not recognizing him and demanding he leave as otherwise she will call 911. And she had called 911. By the time the police came, she had forgotten her concerns.
Anything but dementia

Her clothes were brighter than a field of wildflowers with rich reds and sunny yellows, countless shades of green and blue, deep blacks and grey and purple. I had never seen such colorful elegance before.

We sat on a bench outside, all bundled up. The air had begun to get cold. She liked sitting here.

The sun dipped below the tops of the trees. We watched a dragonfly move lazily amongst the reeds. “Why would they name it dragonfly? It looks nothing like a dragon” she said.

We chatted about this and that and more this and more that. Anything but her husband’s Dementia, her struggles keeping him home, his hallucinations, his anger, her guilt preventing her from asking her only son for help, her fears about future, and her grief.

🙏

It is hard for many family caregivers to talk about their emotional pain. That is okay. It is important just to be with them and talk about this and that. This also promotes healing.
My World

I am a dementia caregiver.  
My world is  
Desolate.  
Not a single living soul.  
Not a bird.  
Not a fly.  
In my world, waves roar  
for no one in particular.  
Where all events  
are beyond comprehension.  
Not a single logical thought.  
Just reflections and reverie,  
sinister at times  
and yet  
also filled with desire  
to look forever  
at the monotonous movements  
of the ocean waves.  
I have become a prisoner  
of my own meditations,  
egulped  
by an overwhelming sense of powerlessness,  
facing  
an inescapably bitter fate

🙏

Caregiver sadness, grief, despair, demoralization and depression are real. We should do everything we can to help. Caregiver journey typically is a lonely journey much of the time. Caregiver depression is one of the key factors increasing the risk of repeated visits of the person with dementia to ED for AAD emergencies.
Long buried sorrow

What is her yelling?
A sign, a signal,
a symptom of moral anguish
calling our souls
to understand her plight?
My patient had never learned
to experience her feelings
as there was no one
in her childhood or later
who accepted her fully,
who understood
and supported her.
And now with advanced dementia,
long buried sorrow
that could never be expressed
is set free and
only touch and soothing voice
can do the miracle
of bringing her some solace.
No meds please,
no meds.

🙏

My 82-year-old patient with advanced dementia living in a nursing home. The staff (well-meaning) asked me if an antidepressant would help. Trauma-informed care plan was created and implemented and it did help her.

Past trauma (includes emotional neglect) is under-appreciated and under-recognized in persons with dementia experiencing AAD. Let’s make inquiry about past trauma and detailed understanding of its context a routine in assessment of AAD.
Restraints

Another patient with advanced dementia
in restraints.
Physical
Chemical
Again
for trying to leave.
I still cannot believe
that conditions so deplorable
exist in our country’s hospitals.
Perhaps I am naïve.
There is no humanity
in our current system of health care.
And no one cares.
No one is watching.
At least no one
with any real power

🙏

Many of us who work in the hospital experience secondary trauma of seeing our patients with dementia and AAD in restraints. Dementia Friendly Hospital Systems and Practices within Age-Friendly Health Systems is the answer. We have the power. All we need is courageous leaders to provide emotional support to the clinical team and adequate funding to make this a reality.
Thanx

My patient smiled, a smile that stopped me in my tracks. I found the smile unexpectedly pleasant. A vigor, a curious sweetness in it. There was also youth in it, a wild hope. Did all my patients with dementia have all this in them? Have I been blind for 23 years? I sat next to her, closed my eyes and breathed quietly. I opened my eyes, turned towards her and said “Thanks”

🙏

This is my 93-year-old patient with advanced dementia in a nursing home. I thanked her for opening up my eyes to the rich inner life persons with dementia have that I have not been fully aware of. Not all persons with dementia develop AAD.

23 years = my being a geriatric psychiatrist
Delirium on Dementia

What exactly happened, Doc?
He doesn’t remember anything.

He was drifting
in dream filled delirium.
Confusion interspersed with
bouts of panic.
Terror blazing in his eyes,
courtesy hallucinations.
Only after intravenous medicine
did his inner chaos settle
and rambling settled into
coherent thinking.
His hip was on fire
and in just one hour
blossomed into a raging inferno.
A shot of opioid
settled that.
The unspoken horrors
of delirium
after hip fracture.
It’s a good thing
he doesn’t remember anything.
Some have PTSD
from such a horrific experience.

🙏

My patient with moderate stage Alzheimer’s dementia fractured his hip after a fall. Emotionally he is doing well now. Dementia has accelerated from all this stress.

His wife asked me the question and I was explaining what happened.

His wife upon my guidance spent several hours every day with him. This was key to his recovery. Unfortunately, she remembers everything and watching his agony remains a stressful memory. With prayers, mindfulness, family support and support groups, she is doing better each day.
An Angel

The nurse aide washed the sweat from his face with a warm, damp cloth. Dressed him with deft, gentle hands. Wheeled him down to the dining hall. He was as light as a bundle of rags. The aide was determined to put some muscle on him. “Don’t you worry. I will make sure,” the aide assured his wife of 67 years. “Then, you both can plan your new adventure.” The wife smiled, turned to me, and said, “She is an angel.”

🙏
My patient (90 years old with advanced dementia, lost a lot of weight) and his wife loved traveling to remote places, their “adventures.”
A Blessing for Caregivers

May the love in your soul guide you
May the courage within you overcome fears about caregiving
May the heart within you conquer the pain involved in caregiving
May you be given the best education and training to overcome challenges in this journey
May you have the commitment to care with compassion, to learn from failures, to be patient, and be the best caregiver you can be
May you have great respect for yourself
May you show growing compassion and patience towards yourself as you come to realize how much the wellbeing of (name of the person needing care) depends on you.
May you have the wisdom to hear
The unspoken gratitude (name of the person needing care) has for you
And above all, may you be given the wonderful gift of meeting the beauty that is within you
May you be blessed, and may you find life Enriched by your efforts of caregiving.

Adapted from John O’Donohue’s poems

Namaste
When Is Enough, Enough? Personal Reflections on the Threats and Ills Facing Persons living with Dementia
Abhilash Desai MD

Why do the terrible deprivations that befall the millions of persons living dementia not routinely keep the rest of us awake at night? The reasons cover a wide range of sociocultural forces, from apathy by the medical community to the fragmented health care system to lack of adequate funding by the government. Running deep are some dominant currents: of identity, in particular, the concern that despite the fact that each of us carries multiple identities, there are forces at play which strive to straitjacket us into just one, typically a narrative identity (an identity that relies solely on neurocognitive functions, especially memory); of autonomy, especially why it is so important to the idea of freedom and justice (autonomy routinely deprived to persons living with dementia); and of inequality, notably in the context of “jaw-dropping” advances in biomedical research which have brought enormous longevity dividends for the few while leaving millions to cope on their own with poor cognitive health. At the heart of the tragedy is the injustice meted out to the individuals living with dementia in long-term care facilities—plagued by their lack of effort to meet the most basic mental health need, the fundamental right to a life lived with dignity and companionship.

Being a physician, I focus my lens on medical education. Let me remind you that, while the United States is home to more than 6 million persons living with dementia, of which more than 1 million are taking antipsychotic medication (typically used as chemical restraint and carrying dangerous risks of swallowing difficulty and stroke), it also boasts the largest number of world-class medical schools and universities. In the case of medical education, despite producing many of the world's most eminent physicians, most of our country's population living with dementia struggle to receive basic good-quality physical and mental health care.

Much health care to persons living with dementia is now provided by physicians and advanced practice providers (APP [e.g., nurse practitioners, physician assistants]) who have minimal education and skills training in management of agitation in the context of dementia. Furthermore, physician and APP clinical practice is seriously compromised (in terms of time spent with patient and family) by poor reimbursement for preventive and palliative services. Persons living with dementia who cannot afford to live at home have to rely on a long-term care system that is short of virtually all resources (e.g., poor quantity and inadequate training and support of direct care staff, very low salaries with minimal health benefits of certified nursing assistants and nurses). Much of the payments from our government to the nursing homes disappears somewhere (no one seems to know where – or a few people do know but would like to keep it a secret) but definitely doesn’t trickle down to improve salaries of direct care staff. There is no transparency in long-term care financing and without transparency, radical changes that are needed in long-term care systems will not take place.

Of course, these are sweeping generalizations that mask islands of excellence in both sectors, but the exceptions do not make the rule. The fundamental question for those who are bewildered by the coexistence of world-class hospitals, complete with helipads and suites that
could make the Ritz Carlton blush, and decrepit long-term care facilities without qualified geriatric health professionals, is why does this not keep us awake at night? Perhaps the most distressing indicator of how immoral the health care system has become is the observation that, whereas our country invests heavily in costly high-tech biomedical devices, medications (the latest being Lecanemab [Leqembi] and Donanemab [Kisunla] for early Alzheimer Disease), and surgical interventions as a means to help people live 1-2 years longer, caring for persons living with dementia has become one of the leading causes of poverty for the person and their families.

What explanation could there be for this appalling situation? It seems as if our society has no social imperative to care for cognitively diminished people, and the physician and APP community (researchers, clinicians, and educators) have – for all practical purposes - divorced themselves from this cause. It seems that we physicians and APPs have chosen to deal with this injustice by simply looking away. Perhaps the indifference of the physician and APP community is because we are besieged with helplessness and burnout; more repugnant is the possibility that some simply don't care.

Most disheartening of all, for physicians and APPs with geriatric expertise, is the recognition that the medical community not only has distanced itself from this injustice but also actually has contributed to it. Whether this takes the shape of the physician / APP who treats their patient living with dementia with a sense that their situation is hopeless, or the physician / APP who ignores their patient with dementia completely and directs interaction to the caregivers accompanying the patient, or the physician / APP who prescribes unnecessary and inappropriate medication, or the surgeon who subjects the person with dementia to dangerous surgeries and procedures, the fact remains that physicians / APPs have lost their moorings to their original purpose: to address human suffering in an ethical, evidence-based, and dignified way. Even the professional societies that govern medicine have not achieved their rightful role in generating social realization.

This overarching division between the physician community and the persons living with dementia and the silence with which it is tolerated, not to mention the smugness with which it is sometimes dismissed, should keep us awake at night. Most tragically, these deprivations are not hard to overcome, as it requires what every physician / APP has (or should have): compassion. After all, medical schools are the world's factory of brilliant, creative, skillful, and (at least when they were medical students) kindest of all professionals. Of course, the physician / APP community does occasionally become sufficiently seized by injustice to protest, alongside fellow patients (with dementia) who live without dignity or support. But this action has been only when the injustice has come too close for comfort, rather than out of solidarity with cognitively diminished individuals. It is no accident that the recent outrage around the use of antipsychotic medication for persons with dementia (and the associated increased risk of stroke and mortality) was precipitated by the findings that two-thirds of persons with dementia are receiving antipsychotics unnecessarily or inappropriately. Of course, persons living with dementia (especially those who live in long-term care facilities) commonly experience excess disability and suffering due to unnecessary and/or inappropriate prescription of many
dangerous medications (not just antipsychotics), but there is no comparable outrage for that cause. The same could be said about aggressive and burdensome medical care, a horror that haunts the last months of life of persons with dementia but emerged as a mass movement only when its high cost became apparent and an impediment to the financial security of the cognitively privileged and, in particular, the government.

We need, first and foremost, a clear-headed understanding of what most ails persons living with dementia. We must identify the obstacles to overcome and acknowledge that there is something astonishing and perplexing about quiet acceptance, with relatively little political murmur, of the continuation of the misery of the least-advantaged people in our country. The complete exclusion of cognitively diminished individuals from daily social interactions with cognitively privileged individuals is one major reason why there is no national compact to address the injustice of iniquitous health care. Dementia-friendly communities, a compassionate and innovative approach for all persons living with dementia, offer a potent solution to this challenge. If we leave aside the fundamental principle of justice as the driver for dementia-friendly communities, an often-cited concern is that that we cannot afford them. Yet, such an attitude also ignores the basic economics that public investment in innovative and humane social solutions is the primary driver for the economic success of caring for persons with dementia in many communities (and, indeed, in the United States, in communities like Fox Valley in Wisconsin and communities in Seattle, Washington). The idea that financing dementia-friendly communities should be left to the market is perhaps the most perverse of all arguments, because buying innovative social solutions is not the same as buying a car or a television; the enormous asymmetry of information and power renders persons with dementia and their family members highly vulnerable to exploitation and vitiates the efficiency of market competition.

Perhaps one way to force action on this injustice might be to require all cognitively privileged individuals to befriend at least one person living with dementia. Although this may sound churlish, such a move would remind those who contribute to the injustice, even if only by their inaction, of how a person with dementia lives. And dies. It is time for the physician / APP community to make the injustice of health care for persons with dementia the focus of its mission for the development of a more humane future for all.

## Holistic Dementia Care Medicine: 11 Domains

<table>
<thead>
<tr>
<th>#</th>
<th>Type</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>Psychoeducation</td>
<td>Improve brain health literacy and address incorrect understanding and unrealistic expectations from psychiatric medications</td>
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<tr>
<td>2</td>
<td>Medical evaluation and workup</td>
<td>Identify and correct reversible medical (e.g., pain, dehydration, hearing and vision deficits, dental problems) causes of neuropsychiatric symptoms</td>
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<tr>
<td>3</td>
<td>De-prescribing</td>
<td>Identify and de-prescribe medications causing neuropsychiatric problems and medications that are unnecessary</td>
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<tr>
<td>4</td>
<td>STEPS</td>
<td>Support system (family and professional care partners) Training Empowerment Praise Support to improve healing and therapeutic approaches and prevent or discontinue counter therapeutic approaches and interventions</td>
</tr>
<tr>
<td>5</td>
<td>Nidotherapy</td>
<td>Improve physical environment and engagement with outdoors and nature (includes occupational therapy assessment)</td>
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<tr>
<td>6</td>
<td>Lifestyle medicine</td>
<td>Improve brain healthy lifestyle (diet, exercise, socialization, sleep, meditation, mind-body exercises, cognitive stimulation)</td>
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<tr>
<td>7</td>
<td>PERMA</td>
<td>Positive emotions, Engagement, Relationships, Meaning, Accomplishment (positive psychology science-based interventions)</td>
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<tr>
<td>8</td>
<td>Emotional regulation skills training</td>
<td>To prevent and reduce dysregulated emotional states in the person with dementia and their care partners</td>
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<tr>
<td>9</td>
<td>ACT</td>
<td>Acceptance Commitment Therapy – for the person with dementia and their care partners to accept the need for holistic approach and commit to it.</td>
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<tr>
<td>10</td>
<td>DEIBA</td>
<td>Diversity, Equity, Inclusion, Belonging, Anti-Ableism Includes Anti-Ageism and Anti-Dementiaism Includes addressing social determinants of health and bolstering community resources</td>
</tr>
<tr>
<td>11</td>
<td>Peer-led interventions</td>
<td>Peer-led and peer-supported interventions including support groups and advocacy groups</td>
</tr>
</tbody>
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### Key Resources:

1. Saint Alphonsus Memory Center, Boise, Idaho.  
   [https://www.saintalphonsus.org/specialty/memory-center/](https://www.saintalphonsus.org/specialty/memory-center/)
2. The DICE Approach.  
   [https://diceapproach.com](https://diceapproach.com)
3. ABC Dementia - Community Dementia Education.  
   [https://community.abcdementia.org](https://community.abcdementia.org)
4. Laura LaForte Therapy.  
   [https://www.lafortetherapy.com](https://www.lafortetherapy.com)
   [https://ppc.sas.upenn.edu/learn-more/perma-theory-well-being-and-perma-workshops](https://ppc.sas.upenn.edu/learn-more/perma-theory-well-being-and-perma-workshops)
New Research

1. Severe aggression and mortality risk: Research from Belgium indicate that severe aggression in individuals with dementia is associated with greater mortality risk (more likely to die within one year) as compared to those with mild or no aggressive behaviors. Van den Bulcke L, et al. November 13, 2023. JAMDA. Clinical implications: Very important to discuss goals of care and What Matters – quality of life over quantity of life discussions.