DEMENTIA MEASURES GROUP OVERVIEW

2015 PQRS OPTIONS FOR MEASURES GROUPS:

2015 PQRS MEASURES IN DEMENTIA MEASURES GROUP:
#47  Care Plan
#280  Dementia: Staging of Dementia
#281  Dementia: Cognitive Assessment
#282  Dementia: Functional Status Assessment
#283  Dementia: Neuropsychiatric Symptom Assessment
#284  Dementia: Management of Neuropsychiatric Symptoms
#285  Dementia: Screening for Depressive Symptoms
#286  Dementia: Counseling Regarding Safety Concerns
#287  Dementia: Counseling Regarding Risks of Driving
#288  Dementia: Caregiver Education and Support

INSTRUCTIONS FOR REPORTING:

- It is not necessary to submit the measures group-specific intent G-code for registry-based submissions. However, the measures group-specific intent G-code has been created for registry only measures groups for use by registries that utilize claims data.

G8902: I intend to report the Dementia Measures Group

- Report the patient sample method:
  20 Patient Sample Method via registries: 20 unique patients (a majority of which must be Medicare Part B FFS patients) meeting patient sample criteria for the measures group during the reporting period (January 1 through December 31, 2015).

- Patient sample criteria for the Dementia Measures Group are all patients with two denominator eligible visits regardless of age, with a specific diagnosis of dementia accompanied by a specific patient encounter:

  One of the following diagnosis codes indicating Dementia:
  ICD-9-CM [for use 1/1/2015 – 9/30/2015]: 094.1, 290.0, 290.10, 290.11, 290.12, 290.13, 290.20, 290.21, 290.3, 290.40, 290.41, 290.42, 290.43, 290.8, 290.9, 294.10, 294.11, 294.20, 294.21, 294.8, 331.0, 331.11, 331.19, 331.82
  ICD-10-CM [for use 10/1/2015 – 12/31/2015]: A52.17, F01.50, F01.51, F02.80, F02.81, F03.90, F03.91, F05, F06.8, G30.0, G30.1, G30.8, G30.9, G31.01, G31.09, G31.83

  Accompanied by:
  One of the following patient encounter codes: 90791, 90792, 90832, 90834, 90837, 96116, 96118, 96119, 96120, 96150, 96151, 96152, 96154, 97003, 97004, 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99304, 99305, 99306, 99307, 99308, 99309, 99310, 99324, 99325, 99326, 99327, 99328, 99334, 99335, 99336, 99337, 99341, 99342, 99343, 99344, 99345, 99347, 99348, 99349, 99350

- Report a numerator option on all applicable measures within the Dementia Measures Group for each patient within the eligible professional’s patient sample.

- Measure #47 need only be reported on patients 65 years and older.
• Instructions for qualifying numerator option reporting for each of the measures within the Dementia Measures Group are displayed on the next several pages. The following composite Quality Data Code (QDC) has been created for registries that utilize claims data. This QDC may be reported in lieu of individual QDCs when all quality clinical actions for all applicable measures within the group have been performed.

**Composite QDC G8761:** All quality actions for the applicable measures in the Dementia Measures Group have been performed for this patient

• To report satisfactorily the Dementia Measures Group it requires **all applicable** measures for each patient within the eligible professional’s patient sample to be reported a minimum of once during the reporting period.

• Measures groups containing a measure with a 0% performance rate will not be counted as satisfactorily reporting the measures group. The recommended clinical quality action must be performed on at least one patient for each measure within the measures group reported by the eligible professional. Performance exclusion quality-data codes are not counted in the performance denominator. If the eligible professional submits all performance exclusion quality-data codes, the performance rate would be 0/0 and would be considered satisfactorily reporting. If a measure within a measures group is not applicable to a patient, the patient would not be counted in the performance denominator for that measure (e.g., Preventive Care Measures Group - Measure #39: Screening or Therapy for Osteoporosis for Women would not be applicable to male patients according to the patient sample criteria). If the measure is not applicable for all patients within the sample, the performance rate would be 0/0 and would be considered satisfactorily reporting.

• **NOTE:** The detailed instructions in this specification apply exclusively to the reporting and analysis of the included measures under the measures group option.
**Measure #47 (NQF 0326): Care Plan – National Quality Strategy Domain: Communication and Care Coordination**

**DESCRIPTION:**
Percentage of patients aged 65 years and older who have an advance care plan or surrogate decision maker documented in the medical record or documentation in the medical record that an advance care plan was discussed but the patient did not wish or was not able to name a surrogate decision maker or provide an advance care plan

**NUMERATOR:**
Patients who have an advance care plan or surrogate decision maker documented in the medical record or documentation in the medical record that an advance care plan was discussed but patient did not wish or was not able to name a surrogate decision maker or provide an advance care plan

**Numerator Instructions:** If patient’s cultural and/or spiritual beliefs preclude a discussion of advance care planning, report 1124F.

**Definition:**
Documentation that Patient did not Wish or was not able to Name a Surrogate Decision Maker or Provide an Advance Care Plan – May also include, as appropriate, the following:
- That the patient’s cultural and/or spiritual beliefs preclude a discussion of advance care planning, as it would be viewed as harmful to the patient’s beliefs and thus harmful to the physician-patient relationship.

**Numerator Options:**
**Performance Met:** Advance Care Planning discussed and documented; advance care plan or surrogate decision maker documented in the medical record (1123F)
OR
**Performance Met:** Advance Care Planning discussed and documented in the medical record; patient did not wish or was not able to name a surrogate decision maker or provide an advance care plan (1124F)
OR
**Performance Not Met:** Advance care planning not documented, reason not otherwise specified (1123F with 8P)
Measure #280: Dementia: Staging of Dementia -- National Quality Strategy Domain: Communication and Care Coordination

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia whose severity of dementia was classified as mild, moderate or severe at least once within a 12 month period

NUMERATOR:
Patients whose severity of dementia was classified as mild, moderate or severe at least once within a 12 month period

Numerator Instructions: Dementia severity can be assessed using one of a number of available valid and reliable instruments available from the medical literature. Examples include, but are not limited to:
- Global Deterioration Scale (GDS)
- Functional Assessment Staging Tool (FAST)
- Clinical Dementia Rating (CDR)
- Dementia Severity Rating Scale
- Mini-Mental State Examination (MMSE) [Note: While simple and quick to administer, the MMSE is a blunt instrument for staging Alzheimer’s disease. The MMSE has not been well validated for non-Alzheimer’s dementias.]
- Formal Neuropsychological Evaluation

Definitions:
Mild dementia - Can be classified quantitatively as MMSE score of > 18, GDS or FAST stage 4, CDR of 1; qualitatively as being likely to have difficulty with balancing a checkbook, preparing a complex meal, or managing a complicated medication schedule. (APA, 2007)
Moderate dementia - Can be classified quantitatively as MMSE score of 10–18, GDS or FAST stages 5 and 6, CDR of 2; qualitatively as experiencing difficulties with simpler food preparation, household cleanup, and yard work and requiring assistance with some aspects of self-care (eg, picking out the proper clothing to wear). (APA, 2007)
Severe dementia - Can be classified quantitatively as MMSE score of < 10, GDS or FAST stages 6 and 7, CDR of 3; qualitatively as requiring considerable or total assistance with personal care, such as dressing, bathing, and toileting. (APA, 2007)

Numerator Note: The proposed scoring cut-offs listed above are offered only as a guide and are quoted verbatim from the referenced clinical guideline. The scoring and appropriate severity cut-offs for any of these instruments must be interpreted in the context of the patient’s age, education, and ethnicity.

Numerator Options:
Performance Met: Dementia severity classified, mild (1490F)
OR
Performance Met: Dementia severity classified, moderate (1491F)
OR
Performance Met: Dementia severity classified, severe (1493F)
OR
Performance Not Met: Dementia severity not classified, reason not otherwise specified (1490F with 8P)
▲ Measure #281: Dementia: Cognitive Assessment -- National Quality Strategy Domain: Effective Clinical Care

**DESCRIPTION:**
Percentage of patients, regardless of age, with a diagnosis of dementia for whom an assessment of cognition is performed and the results reviewed at least once within a 12 month period

**NUMERATOR:**
Patients for whom an assessment of cognition is performed and the results reviewed at least once within a 12 month period

**Numerator Instructions:**
Cognition can be assessed by the clinician during the patient's clinical history. Cognition can also be assessed by direct examination of the patient using one of a number of instruments, including several originally developed and validated for screening purposes. This can also include, where appropriate, administration to a knowledgeable informant. Examples include, but are not limited to:

- Blessed Orientation-Memory-Concentration Test (BOMC)
- Montreal Cognitive Assessment (MoCA)
- St. Louis University Mental Status Examination (SLUMS)
- Mini-Mental State Examination (MMSE) [*Note: The MMSE has not been well validated for non-Alzheimer’s dementias.*]
- Short Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE)
- Ascertain Dementia 8 (AD8) Questionnaire
- Minimum Data Set (MDS) Brief Interview for Mental Status (BIMS) [*Note: Validated for use with nursing home patients only*]
- Formal neuropsychological evaluation

**Numerator Options:**
- **Performance Met:** Cognition assessed and reviewed (1494F)
- **Medical Performance Exclusion:** Documentation of medical reason(s) for not assessing cognition (eg, patient with very advanced stage dementia, other medical reason) (1494F with 1P)
- **Patient Performance Exclusion:** Documentation of patient reason(s) for not assessing cognition (1494F with 2P)
- **Performance Not Met:** Cognition not assessed and reviewed, reason not otherwise specified (1494F with 8P)
Measure #282: Dementia: Functional Status Assessment -- National Quality Strategy
Domain: Effective Clinical Care

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia for whom an assessment of functional status is performed and the results reviewed at least once within a 12 month period.

NUMERATOR:
Patients for whom an assessment of functional status is performed and the results reviewed at least once within a 12 month period.

Numerator Instructions:
Functional status can be assessed by direct examination of the patient or knowledgeable informant. An assessment of functional status should include, at a minimum, an evaluation of the patient's ability to perform instrumental activities of daily living (IADL) and basic activities of daily living (ADL). Functional status can also be assessed using one of a number of available valid and reliable instruments available from the medical literature. Examples include, but are not limited to:
- Lawton IADL Scale
- Barthel ADL Index
- Katz Index of Independence in ADL

Numerator Options:
Performance Met: Functional status for dementia assessed and results reviewed (1175F)

OR

Medical Performance Exclusion: Documentation of medical reason(s) for not assessing and reviewing functional status for dementia (eg, patient is severely impaired and caregiver knowledge is limited, other medical reason) (1175F with 1P)

OR

Performance Not Met: Functional status for dementia not assessed and results not reviewed, reason not otherwise specified (1175F with 8P)
Measure #283: Dementia: Neuropsychiatric Symptom Assessment – National Quality Strategy Domain: Effective Clinical Care

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia and for whom an assessment of neuropsychiatric symptoms is performed and results reviewed at least once in a 12 month period

NUMERATOR:
Patients for whom an assessment of neuropsychiatric symptoms is performed and results reviewed at least once in a 12 month period

Numerator Instructions: Neuropsychiatric symptoms can be assessed by direct examination of the patient or knowledgeable informant.

Examples of reliable and valid instruments that are commonly used in research settings and that can be used to assess behavior include, but are not limited to:

- Dementia Signs and Symptoms (DSS) Scale
- Neuropsychiatric Inventory (NPI)

The assessment of behavioral status may include the assessment of Behavioral and Psychological Symptoms of Dementia (BPSD). For patients residing in nursing homes, it may include an assessment of the behavioral symptom items from the Minimum Data Set (MDS).

The following is a non-exhaustive list of dimensions (based on items included in available validated instruments) that may be evaluated during an assessment of neuropsychiatric symptoms:

Activity disturbances:
- agitation
- wandering
- purposeless hyperactivity
- verbal or physical aggressiveness
- resistiveness with care
- apathy
- impulsiveness
- socially inappropriate behaviors
- appetite
- eating disturbances
- sleep problems
- diurnal/sleep-wake cycle disturbances
- repetitive behavior

Mood disturbances:
- anxiety
- dysphoria
- euphoria
- irritability
- mood lability/fluctuations
Thought and perceptual disturbances:
- having fixed false beliefs (delusions)
- hearing or seeing non-present entities (hallucinations)
- paranoia

Numerator Options:
Performance Met: Neuropsychiatric symptoms assessed and results reviewed (1181F)

OR

Performance Not Met: Neuropsychiatric symptoms not assessed and results not reviewed, reason not otherwise specified (1181F with 8P)
Measure #284: Dementia: Management of Neuropsychiatric Symptoms -- National Quality Strategy Domain: Effective Clinical Care

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia who have one or more neuropsychiatric symptoms who received or were recommended to receive an intervention for neuropsychiatric symptoms within a 12 month period

NUMERATOR:
Patients who received or were recommended to receive an intervention for neuropsychiatric symptoms within a 12 month period

Numerator Options:
Performance Met:
One or more neuropsychiatric symptoms (G8947)
AND
Neuropsychiatric intervention ordered (4525F)
OR
Performance Met: Neuropsychiatric intervention received (4526F)
OR
Other Performance Exclusion: No neuropsychiatric symptoms (G8948)
OR
Performance Not Met:
One or more neuropsychiatric symptoms (G8947)
AND
Neuropsychiatric intervention not ordered, reason not otherwise specified (4525F with 8P)
Measure #285: Dementia: Screening for Depressive Symptoms -- National Quality Strategy Domain: Effective Clinical Care

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia who were screened for depressive symptoms within a 12 month period

NUMERATOR:
Patients who were screened for depressive symptoms within a 12 month period

Numerator Instructions:
In addition to clinical qualitative approaches, dementia patients can be screened for depressive symptoms using one of a number of valid, reliable instruments available from the medical literature. Examples include, but are not limited to:

- Cornell Scale for Depression in Dementia
- Geriatric Depression Scale
- PHQ-9

Definition:
Depressive Symptoms - Depressive symptoms in a patient with dementia can include: anxiety, sadness, lack of reactivity to pleasant events, irritability, agitation, retardation, multiple physical complaints, acute loss of interest, appetite loss, lack of energy, diurnal variation of mood, difficulty falling asleep, multiple awakenings, during sleep, early morning awakenings, suicide, self-depreciation, pessimism, and mood congruent delusions. Since patients may be unable to describe their symptoms, caregiver report of depressive symptoms should be reviewed and included in the screen for depressive symptoms.

Numerator Options:
Performance Met: Screening for depression performed (3725F)

OR

Performance Not Met: Screening for depression not performed, reason not otherwise specified (3725F with 8P)

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia or their caregiver(s) who were counseled or referred for counseling regarding safety concerns within a 12 month period.

NUMERATOR:
Patients or their caregiver(s) who were counseled or referred for counseling regarding safety concerns within a 12 month period.

Numerator Instructions: Counseling should include a discussion with the patient and their caregiver(s) regarding one or more of the following common safety concerns and potential risks to the patient. When appropriate, it should also include a recommendation or referral for a home safety evaluation.

Note: For nursing home patients, different safety concerns might apply.

A number of organizations have developed educational materials that are recommended to aid implementation of the measure. These materials/tools include:


Definition:
Caregiver(s) - Person(s) who provide care to those who need supervision or assistance in illness or disability. They may provide the care in the home, in a hospital, or in an institution. Although caregiver(s) include trained medical, nursing, and other health personnel, the concept also refers to parents, spouses, or other family members, friends, members of the clergy, teachers, social workers, fellow patients.

Safety Concerns - Safety concerns include, but are not limited to:
- Fall risk
- Gait/balance
- Medication management
- Financial management
- Home safety risks that could arise from cooking or smoking
- Physical aggression posing threat to self, family caregiver, or others
- Wandering
- Access to firearms or other weapons
- Access to potentially dangerous materials
- Being left alone in home or locked in room
- Inability to respond rapidly to crisis/household emergencies
- Driving
- Operation of hazardous equipment
- Suicidality
- Abuse or neglect

Numerator Options:
Performance Met: Safety counseling for dementia provided (6101F)
OR

**Performance Met:** Safety counseling for dementia ordered (6102F)

OR

**Medical Performance Exclusion:** Documentation of medical reason(s) for not providing counseling regarding safety concerns (eg, patient in palliative care, other medical reason) (6101F with 1P)

OR

**Medical Performance Exclusion:** Documentation of medical reason(s) for not ordering safety counseling (eg, patient in palliative care, other medical reason) (6102F with 1P)

OR

**Performance Not Met:** Safety counseling for dementia not provided, reason not otherwise specified (6101F with 8P)

OR

**Performance Not Met:** Safety counseling for dementia not ordered, reason not otherwise specified (6102F with 8P)
Measure #287: Dementia: Counseling Regarding Risks of Driving -- National Quality Strategy Domain: Effective Clinical Care

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia or their caregiver(s) who were counseled regarding the risks of driving and the alternatives to driving at least once within a 12 month period.

NUMERATOR:
Patients or their caregiver(s) who were counseled regarding the risks of driving and the alternatives to driving at least once within a 12 month period.

Numerator Instructions:
One resource that includes patient and caregiver educational materials that can be used to aid implementation of the measure is the *Physician's Guide to Assessing and Counseling Older Drivers*, developed by the American Medical Association in cooperation with the National Highway Traffic Safety Administration. This document is available on the AMA website.

Definition:
Caregiver(s) - Person(s) who provide care to those who need supervision or assistance in illness or disability. They may provide the care in the home, in a hospital, or in an institution. Although caregiver(s) include trained medical, nursing, and other health personnel, the concept also refers to parents, spouses, or other family members, friends, members of the clergy, teachers, social workers, fellow patients.

Numerator Options:
*Performance Met:* Counseling provided regarding risks of driving and the alternatives to driving (6110F)

OR

*Medical Performance Exclusion:* Documentation of medical reason(s) for not counseling regarding the risks of driving (eg, patient is no longer driving, other medical reason) (6110F with 1P)

OR

*Performance Not Met:* Counseling regarding risks of driving and alternatives to driving not performed, reason not otherwise specified (6110F with 8P)
Measure #288: Dementia: Caregiver Education and Support -- National Quality Strategy
Domain: Communication and Care Coordination

DESCRIPTION:
Percentage of patients, regardless of age, with a diagnosis of dementia whose caregiver(s) were provided with education on dementia disease management and health behavior changes AND referred to additional resources for support within a 12 month period

NUMERATOR:
Patients whose caregiver(s) were provided with education on dementia disease management and health behavior changes AND referred to additional resources for support within a 12 month period

Numerator Instructions:
There are a number of assessment tools available for the caregiver. These should be considered as an integral component of comprehensive caregiver education and support. The American Medical Association has developed a Caregiver Health Self-assessment Questionnaire to help caregivers analyze their own behavior and health risks and, with their physician’s help, make decisions that will benefit both the caregiver and the patient. This questionnaire is available on the AMA website.

Definitions:
Caregiver(s) - Person(s) who provide care to those who need supervision or assistance in illness or disability. They may provide the care in the home, in a hospital, or in an institution. Although caregiver(s) include trained medical, nursing, and other health personnel, the concept also refers to parents, spouses, or other family members, friends, members of the clergy, teachers, social workers, fellow patients.

Education – Education should also include advising the caregiver that he or she is at “increased risk of serious illness (including circulatory and heart conditions and respiratory disease and hypertension), increased physician visits and use of prescription medications, emotional strain, anxiety, and depression.”

Numerator Options:
Performance Met: Caregiver provided with education and referred to additional resources for support (4322F)

OR

Medical Performance Exclusion: Documentation of medical reason(s) for not providing the caregiver with education on disease management and health behavior changes or referring to additional sources for support (eg, patient does not have a caregiver, other medical reason) (4322F with 1P)

OR

Performance Not Met: Caregiver not provided with education and not referred to additional resources for support, reason not otherwise specified (4322F with 8P)
DEMENTIA MEASURES GROUP RATIONALE AND CLINICAL RECOMMENDATION STATEMENTS

Measure #47 - Care Plan

RATIONALE:
It is essential that the patient’s wishes regarding medical treatment be established as much as possible prior to incapacity. The Work Group has determined that the measure should remain as specified with no required timeframe based on a review of the literature. Studies have shown that people do change their preferences often with regard to advanced care planning, but it primarily occurs after a major medical event or other health status change. In the stable patient, it would be very difficult to define the correct interval. It was felt by the Work Group that the error rate in simply not having addressed the issue at all is so much more substantial (Teno, 1997) than the risk that an established plan has become outdated that we should not define a specific timeframe at this time. As this measure is tested and reviewed, we will continue to evaluate if and when a specific timeframe should be included.

CLINICAL RECOMMENDATION STATEMENTS:
Advance directives are designed to respect patient’s autonomy and determine his/her wishes about future life-sustaining medical treatment if unable to indicate wishes. Key interventions and treatment decisions to include in advance directives are: resuscitation procedures, mechanical respiration, chemotherapy, radiation therapy, dialysis, simple diagnostic tests, pain control, blood products, transfusions, and intentional deep sedation.

Oral statements
- Conversations with relatives, friends, and clinicians are most common form; should be thoroughly documented in medical record for later reference.
- Properly verified oral statements carry same ethical and legal weight as those recorded in writing.

Instructional advance directives (DNR orders, living wills)
- Written instructions regarding the initiation, continuation, withholding, or withdrawal of particular forms of life-sustaining medical treatment.
- May be revoked or altered at any time by the patient.
- Clinicians who comply with such directives are provided legal immunity for such actions.

Durable power of attorney for health care or health care proxy
- A written document that enables a capable person to appoint someone else to make future medical treatment choices for him or her in the event of decisional incapacity. (AGS)

The National Hospice and Palliative Care Organization provides the Caring Connection web site, which provides resources and information on end-of-life care, including a national repository of state-by-state advance directives.

Measure #280 Dementia: Staging of Dementia

RATIONALE:
Dementia is characterized by continued and progressive impairment in cognition and function including the evolution of symptoms over time. (APA, 2007)

The treatment varies throughout the disease course. (APA, 2007)

Patients with dementia, therefore, require assessment of disease severity and subsequent treatment specific and appropriate to their current stage of disease. (APA, 2007)
Early stage patients, for example, have special needs and can and should be involved in care planning and referred to community resources. (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008)

Care for late stage patients may focus on improving the quality of life for patients and caregivers, maintaining optimal function and providing maximum comfort. (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

**CLINICAL RECOMMENDATION STATEMENTS:**
Progressive dementias are generally staged globally according to the level of cognitive and functional impairment, and the same categories may be used to describe the degree of severity of any dementia. However, the staging criteria have not been well validated for non-Alzheimer’s dementias. Specific functional staging (FAST staging) has also been developed, is widely used, and can be very useful in tracking the course of Alzheimer’s disease and other dementias. The CDR is a commonly used scale to stage dementia severity. The Global Deterioration Scale (GDS) distinguishes three stages in this range. (APA, 2007)

Individuals with “mild” dementia (MMSE score of >18, GDS or FAST stage 4, CDR of 1) are likely to have difficulties with balancing a checkbook, preparing a complex meal, or managing a difficult medication schedule. Those with “moderate” impairment (MMSE score of 10–18, GDS or FAST stages 5 and 6, CDR of 2) also have difficulties with simpler food preparation, household cleanup, and yard work and may require assistance with some aspects of self-care (eg, picking out the proper clothing to wear). Those whose dementia is “severe” (MMSE score of <10, GDS or FAST stages 6 and 7, CDR of 3) require considerable or total assistance with personal care, such as dressing, bathing, and toileting. Research has shown that measurable cognitive abilities remain throughout the course of severe dementia. In the terminal phase, patients become bed bound, develop contractures, require constant care, and may be susceptible to accidents and infectious diseases, which ultimately prove fatal. (APA, 2007)

**Measure #281 – Dementia:Cognitive Assessment**

**RATIONALE:**
Dementia is often characterized by the gradual onset and continuing cognitive decline in one or more domains including memory, executive function, language, judgment, and spatial abilities. (APA, 2007) Cognitive deterioration represents a major source of morbidity and mortality and poses a significant burden on affected individuals and their caregivers. (NIH, 2010) Although cognitive deterioration follows a different course depending on the type of dementia, significant rates of decline have been reported. For example, one study found that the annual rate of decline for Alzheimer’s disease patients was more than four times that of older adults with no cognitive impairment. (Wilson et al., 2010) Nevertheless, measurable cognitive abilities remain throughout the course of dementia. (APA, 2007) Initial and ongoing assessments of cognition are fundamental to the proper management of patients with dementia. These assessments serve as the basis for identifying treatment goals, developing a treatment plan, monitoring the effects of treatment, and modifying treatment as appropriate.

**CLINICAL RECOMMENDATION STATEMENTS:**
Ongoing assessment includes periodic monitoring of the development and evolution of cognitive and noncognitive psychiatric symptoms and their response to intervention (Category I). Both cognitive and noncognitive neuropsychiatric and behavioral symptoms of dementia tend to evolve over time, so regular monitoring allows detection of new symptoms and adaptation of treatment strategies to current needs…Cognitive symptoms that almost always require assessment include impairments in memory, executive function, language, judgment, and spatial abilities. It is often helpful to track cognitive status with a structured simple examination. (APA, 2007)
Conduct and document an assessment and monitor changes in cognitive status using a reliable and valid instrument. Cognitive status should be reassessed periodically to identify sudden changes, as well as to monitor the potential beneficial or harmful effects of environmental changes, specific medications, or other interventions. Proper assessment requires the use of a standardized, objective instrument that is relatively easy to use, reliable (with less variability between different assessors), and valid (results that would be similar to gold-standard evaluations). (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008)

Measure #282 – Dementia: Functional Status Assessment

RATIONALE:

CLINICAL RECOMMENDATION STATEMENTS:
A detailed assessment of functional status may also aid the clinician in documenting and tracking changes over time as well as providing guidance to the patient and caregivers. Functional status is typically described in terms of the patient’s ability to perform instrumental activities of daily living such as shopping, writing checks, basic housework, and activities of daily living such as dressing, bathing, feeding, transferring, and maintaining continence. These regular assessments of recent cognitive and functional status provide a baseline for assessing the effect of any intervention, and they improve the recognition and treatment of acute problems, such as delirium. (APA, 2007)

Conduct and document an assessment and monitor changes in daily functioning, including feeding, bathing, dressing, mobility, toileting, continence, and ability to manage finances and medications… Functional assessment includes evaluation of physical, psychological, and socioeconomic domains. Physical functioning may focus on basic activities of daily living (ADLs) that include feeding, bathing, dressing, mobility, and toileting. Assessment of instrumental (or intermediate) activities of daily living (IADLs) addresses more advanced self-care activities, such as shopping, cooking, and managing finances and medications. Standardized assessment instruments such as the Barthel or Katz indices can provide information on the patient’s capacity for self-care and independent living. Proxies or patient surrogates can complete a number of these instruments when necessary. The initial assessment of functional abilities is important to determine a baseline to which future functional deficits may be compared. (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008)

Measure #283 – Dementia: Neuropsychiatric Symptom Assessment

RATIONALE:
Neuropsychiatric symptoms appear to be common for patients with dementia. In community samples of dementia patients, the prevalence of neuropsychiatric symptoms range from 40-88%. (Lyketsos CG et al. JAMA. 2002; 288:1475-1483., Ikeda M et al. J Neurol Neurosurg Psychiatry.

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**CLINICAL RECOMMENDATION STATEMENTS:**

It is important for the [clinician] treating a patient with dementia to regularly assess cognitive deficits or behavioral difficulties that potentially pose a danger to the patient or others. (APA, 2007)

Conduct and document an assessment and monitor changes in behavioral symptoms, psychotic symptoms, or depression. (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008)

**For mild to moderate Alzheimer’s disease**

Assessment of patients with mild to moderate AD [Alzheimer’s Disease] should include measures of behavior and other neuropsychiatric symptoms. (Grade B, Level 3) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

**For severe Alzheimer’s disease**

Assessment should include cognition (eg, MMSE), function, behaviour, medical status, nutrition, safety and caregiver health. (Grade B, Level 3) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

**Measure #284 – Dementia: Management of Neuropsychiatric Symptoms**

**RATIONALE:**

Nonpharmacologic interventions should be considered in all cases and in some will be the mainstay of management. Examples of approaches that may be useful include behavioural management for depression, education programs for caregivers and staff to teach them how to recognize, manage, and sometimes prevent behavioral problems, stress reduction for caregivers, and, for patients living at home, enrollment in adult day programs offering structured activities and social stimulation. The evidence evaluating non-pharmacological interventions varies considerably in quality and amount, but broadly supports an individualized approach that includes one or more such interventions. A management plan that assesses the severity and intrusiveness of problematic behaviors can assist clinicians in determining what pharmacologic or non-pharmacologic interventions might be appropriate. 

CLINICAL RECOMMENDATION STATEMENTS:

For mild to moderate Alzheimer's disease
The management of BPSD [Behavioral and Psychological Symptoms of Dementia] should include a careful documentation of behaviours and identification of target symptoms, a search for potential triggers or precipitants, recording of the consequences of the behaviour, an evaluation to rule out treatable or contributory causes, and consideration of the safety of the patient, their caregiver, and others in their environment. (Grade B, Level 3) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

For severe Alzheimer's disease
The management of BPSD should begin with appropriate assessments, diagnosis, and identification of target symptoms and consideration of safety of the patient, their caregiver and others in their environment. (Grade B, Level 3) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

There are no fully comprehensive consensus guidelines for use of specific non-pharmacological approaches to neuropsychiatric symptoms. Patient heterogeneity, variations in care settings, and the broad range of non-pharmacological interventions having some empirical support impede uniform generalization. However, the following evidence statements serve as the evidence to support the measure and are quoted verbatim from the referenced clinical guidelines.

Nonpharmacologic interventions should be initiated first. Approaches that may be useful for severe Alzheimer disease include behaviourial management for depression, and education programs for caregivers and staff to teach them how to recognize behavioural problems and to teach them behaviour-modification techniques. Music therapy and controlled multisensory stimulation (Snoezelen) are useful during treatment sessions, but longer-term benefits have not been demonstrated. (Grade B, Level 1) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

Except for emergency situations, non-pharmacological strategies are the preferred first-line treatment approach for behavioral problems. Medications should be used only as a last resort, if non-pharmacological approaches prove unsuccessful and they are clinically indicated. (California Workgroup on Guidelines for Alzheimer's Disease Management, 2008)

Pharmacologic therapies should be initiated concurrently with nonpharmacologic interventions in the presence of severe depression, psychosis or aggression that puts the patient or others at risk.
of harm. (Grade B, Level 3) (Third Canadian Consensus Conference on the Diagnosis and Treatment of Dementia, 2008)

Measure #285 – Dementia: Screening for Depressive Symptoms

RATIONALE:
Depression is one of the most common co-occurring psychiatric conditions in dementia patients, affecting over 50% of patients with Alzheimer’s disease. (Starkstein SE et al. Am J Psychiatry. 2005;162:2086-2093.) Depression can be reliably detected and quantified, and can be differentiated from the other neuropsychiatric symptoms of dementia. (Lyketsos CG et al. Dement Geriatr Cogn Disord. 2004;17:55-64.) The impact of depression is significant with even mild levels of depression in dementia patients associated with higher rates of disability, impaired quality of life, and greater mortality. (APA, 2007) In particular, Alzheimer’s disease patients with depression have demonstrated “significantly more severe apathy, delusions, anxiety, pathological affective crying, irritability, deficits in activities of daily living, impairments in social functioning, and parkinsonism than Alzheimer’s disease patients without depression.” (Starkstein SE et al. Am J Psychiatry. 2005;162:2086-2093.) Furthermore, with increasing severity of depression, the severity of psychopathological and neurological impairments in dementia patients increases. (Starkstein SE et al. Am J Psychiatry. 2005;162:2086-2093.) Identifying depression in patients with dementia is therefore essential for early intervention and proper management.

CLINICAL RECOMMENDATION STATEMENTS:
Depression is a common, treatable comorbidity in patients with dementia and should be screened for (Guideline). (AAN, 2001)

Ongoing assessment includes periodic monitoring of the development and evolution of cognitive and noncognitive psychiatric symptoms and their response to intervention (Category I)…Among the neuropsychiatric symptoms that require ongoing assessment are depression (including major depression and other depressive syndromes), suicidal ideation or behavior, hallucinations, delusions, agitation, aggressive behavior, disinhibition, sexually inappropriate behavior, anxiety, apathy, and disturbances of appetite and sleep. (APA, 2007)

Conduct and document an assessment and monitor changes in behavioral symptoms, psychotic symptoms, or depression…It is important for health care professionals to be sensitive to symptoms of affective disorders associated with Alzheimer’s Disease and to facilitate early intervention. Since administering assessment tests for depression to Alzheimer’s Disease patients is often challenging and patients may be unable to describe their symptoms to the [primary care practitioner], gathering data from family members becomes especially important. The Cornell Scale for Depression in Dementia is a useful tool for providers because it captures both patient and caregiver input. (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008)

In patients with serious illness at the end of life, clinicians should regularly assess patients for pain, dyspnea, and depression. (Grade: strong recommendation, moderate quality of evidence.) (ACP, 2008)

Measure #286 – Dementia: Counseling Regarding Safety Concerns

RATIONALE:
The vast majority (87%) of individuals with Alzheimer’s disease are cared for at home by family members. (Alz Assoc, 2009) "As the disease progresses however, physical features of the home environment may present as a safety hazard or barrier to performing activities of daily living, particularly at the moderate stage of the disease process." (Gitlin LN et al. Disabil Rehabil. 2002, Vol. 24 , No. 1-3, Pages 59-71.) Safety concerns should be addressed with patients and their caregivers throughout the course of the disease.
CLINICAL RECOMMENDATION STATEMENTS:
Recommended assessments include evaluation of suicidality, dangerousness to self and others, and the potential for aggression, as well as evaluation of living conditions, safety of the environment, adequacy of supervision, and evidence of neglect or abuse (Category I). Important safety issues in the management of patients with dementia include interventions to decrease the hazards of wandering and recommendations concerning activities such as cooking, driving, hunting, and the operation of hazardous equipment. Caregivers should be referred to available books [and other materials] that provide advice and guidance about maximizing the safety of the environment for patients with dementia...As patients become more impaired, they are likely to require more supervision to remain safe, and safety issues should be addressed as part of every evaluation. Families should be advised about the possibility of accidents due to forgetfulness (eg, fires while cooking), of difficulties coping with household emergencies, and of the possibility of wandering. Family members should also be advised to determine whether the patient is handling finances appropriately and to consider taking over the paying of bills and other responsibilities. At this stage of the disease [ie, moderately impaired patients], nearly all patients should not drive. (APA, 2007)

Safety issues such as driving, fall risk, medication management, environmental hazards, wandering, and access to firearms need to be discussed periodically with the patient and caregiver. Safety concerns typically focus on three risks in particular: falling, wandering, and driving. (California Workgroup on Guidelines for Alzheimer's Disease Management, 2008)

For mild to moderate Alzheimer's disease
Assess for safety risks (eg, driving, financial management, medication management, home safety risks that could arise from cooking or smoking, potentially dangerous behaviours such as wandering). (Canadian Consensus Conference on Diagnosis and Treatment of Dementia, 2008)

Measure #287 – Dementia: Counseling Regarding Risks of Driving
RATIONALE:
Motor vehicle-related injuries are a leading cause of injury deaths in adults over 65. (AMA Physician's Guide to Assessing and Counseling Older Drivers, 2010) Per mile driven, drivers age 75 and older are involved in significantly more motor vehicle crashes than middle-aged drivers. (AMA Physician's Guide to Assessing and Counseling Older Drivers, 2010) Dementia has a negative impact on driving skills which deteriorate with increasing dementia severity. (AAN, 2010)

Compared with cognitively intact older adults drivers, studies suggest that drivers with dementia have at least a 2-fold greater risk of crashes. (Carr DB et al. JAMA. 2010;303(16):1632-1641.) “Physicians can influence their patients’ decisions to modify or stop driving. They can also help their patients maintain safe driving skills.” (AMA Physician’s Guide to Assessing and Counseling Older Drivers, 2010) Clinicians should address the risks of driving in patients with dementia for the safety of the patient and everyone on the road.

CLINICAL RECOMMENDATION STATEMENTS:
A diagnosis of Alzheimer's disease is not, on its own, a sufficient reason to withdraw driving privileges. The determining factor in withdrawing driving privileges should be an individual’s driving ability. (Alzheimer’s Association, 2001)

All patients and families should be informed that even mild dementia increases the risk of vehicular accidents (Category I). Mildly impaired patients should be advised to limit their driving to safer situations or to stop driving (Category I), and moderately impaired patients should be instructed not to drive (Category I). Advice about driving cessation should also be communicated to family members, as the implementation of the recommendation often falls on them (Category I). Relevant state laws regarding notification should be followed (Category I). (APA, 2007)
For patients with dementia, consider the following characteristics useful for identifying patients at increased risk for unsafe driving: the Clinical Dementia Rating scale (Level A), a caregiver’s rating of a patient’s driving ability as marginal or unsafe (Level B), a history of crashes or traffic citations (Level C), reduced driving mileage or self-reported situational avoidance (Level C), Mini-Mental State Examination scores of (California Workgroup on Guidelines for Alzheimer’s Disease Management, 2008) or less (Level C), and aggressive or impulsive personality characteristics (Level C). Consider the following characteristics not useful for identifying patients at increased risk for unsafe driving: a patient’s self-rating of safe driving ability (Level A) and lack of situational avoidance (Level C). There is insufficient evidence to support or refute the benefit of neuropsychological testing, after controlling for the presence and severity of dementia, or interventional strategies for drivers with dementia (Level U). Clinicians may present patients and their caregivers with the data showing that, as a group, patients with mild dementia (CDR of 1) are at a substantially higher risk for unsafe driving and thus should strongly consider discontinuing driving. At the very least, patients and their caregivers should prepare for the eventuality of driving cessation as dementia severity increases. (AAN, 2010)

Measure #288 – Dementia: Caregiver Education and Support

RATIONALE:
The vast majority (87%) of individuals with Alzheimer’s disease are cared for at home by family members. (Alz Assoc, 2009) Chodosh et al. found that greater caregiver knowledge of dementia management was associated with higher care quality. (Chodosh J et al. J Am Geriatr Soc. 2007 Aug;55(8):1260-8.) Other studies have indicated that intensive caregiver support in the form of individual and family counseling and on-going telephone counseling results in improved patient health outcomes. (Gaugler JE et al. J Am Geriatr Soc. 2005;53:2098–2105., Mittelman MS et al. Neurology. 2006;67:1592–1599.) Providing education to caregivers and referring them to additional sources for support is a critically important piece of comprehensive care for patients with dementia.
CLINICAL RECOMMENDATION STATEMENTS:
Important aspects of psychiatric management include educating patients and families about the illness, its treatment, and sources of additional care and support (eg, support groups, respite care, nursing homes, and other long-term-care facilities) and advising patients and their families of the need for financial and legal planning due to the patient's eventual incapacity (eg, power of attorney for medical and financial decisions, an up-to-date will, and the cost of long-term care) (Category I)... The family should be educated regarding basic principles of care, including 1) recognizing declines in capacity and adjusting expectations appropriately, 2) bringing sudden declines in function and the emergence of new symptoms to professional attention, 3) keeping requests and demands relatively simple, 4) deferring requests if the patient becomes overly upset or angered, 5) avoiding overly complex tasks that may lead to frustration, 6) not confronting patients about their deficits, 7) remaining calm, firm, and supportive and providing redirection if the patient becomes upset, 8) being consistent and avoiding unnecessary change, and 9) providing frequent reminders, explanations, and orientation cues... In addition to providing families with information on support groups, there are a number of benefits of referral to the local chapter or national office of the Alzheimer's Association (1-800-272-3900; http://www.alz.org), the Alzheimer's Disease Education and Referral Center (ADEAR) (1-800-438-4380; http://www.nia.nih.gov/Alzheimers/), and other support organizations. (APA, 2007).

Studies have shown that education and support for caregivers increases the chances of adherence to treatment recommendations for patients. The PCP should provide information and education about the current stage of the disease process and talk with the patient and family to establish treatment goals. Based on the agreed-upon goals, a discussion regarding the expected effects (positive and negative) of interventions on cognition, mood, and behavior will ensure that the prescribed treatment strategy is appropriate to family values and culture. (California Workgroup on Guidelines for Alzheimer's Disease Management, 2008)

Seamless resource referral and access to critical services for both patients and caregivers are considered essential. The PCP should encourage the caregiver to participate in educational programs, support groups, respite services, and adult day service programs. The local Alzheimer's Association chapter or other local agency support groups and community resources such as the Caregiver Resources Centers should be recommended. (California Workgroup on Guidelines for Alzheimer's Disease Management, 2008).