Electronic Clinical Quality Measures

(ECQM)

Measure Guidance Manual

Version 3.0
December 2020
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1.0 Introduction

Electronic Clinical Quality Measures (eCQM) are standardized metrics that measure and track the quality of health care services that eligible professionals (EP), eligible hospitals (EH), and critical access hospitals (CAH) provide. The results of the measures are used to calculate a quality score. This process helps to ensure that our health care system is delivering effective, safe, efficient, patient-centered, equitable, and timely care. While the eCQMs are not practice guidelines, they are indicative through measuring positive or negative outcomes of good clinical practices.

eCQM performance rates are used by various governing bodies to evaluate programs and in the case of Centers for Medicare & Medicaid Services (CMS), payments for Medicare services may be affected. eCQMs measure many aspects of patient care, including:

- Patient and Family Engagement
- Patient Safety
- Care Coordination
- Population/Public Health
- Efficient Use of Healthcare Resources
- Clinical Process/Effectiveness

Indian Health Service Resource Patient Management System Certified Electronic Health Record (IHS RPMS-CEHR) generates patient-based files containing the data needed to create CQM reports in standardized format. These are called Quality Reporting Data Architecture (QRDA) Category I (CAT-I) files. The ECQM Engine is a browser-enabled graphical user interface for the Indian Health Service (IHS) which extracts the data from multiple CAT-I files and generates QRDA Category III (CAT-III) aggregated report files which may be submitted to CMS. The ECQM Engine also outputs human-readable reports that can be used in quality improvement activities at individual sites.

eCQMs are tools that help measure and track the quality of health care services that EP, EH, and CAH provide, as generated by a provider's electronic health record (EHR). Health care providers are required to electronically report eCQMs, which use data from EHRs and/or health information technology systems to measure health care quality.

eCQMs are a component of the 2015 Edition Health Information Technology (Health IT) certification criteria necessary for participating in various CMS Programs. RPMS EHR is being updated to meet these expanded criteria.
Each year, CMS makes updates to the eCQMs approved for CMS programs to reflect changes in:

- Evidence-based Medicine
- Code Sets
- Measure Logic

Measures will need to be updated for the 2021 and subsequent reporting periods.

The 2019 and 2020 reporting period for EHs and CAHs who report electronically is one self-selected quarter of calendar year (CY) data. If they submit electronically for 2019, they must submit four eCQMs from the list of 16 available eCQMs. If they submit electronically for 2020, they must submit four eCQMs from the list of eight available eCQMs (indicated with # in the list that follows).

The 2019 and 2020 eCQM reporting period for EP is the full CY, except for first time meaningful users, for whom the reporting period is any continuous 90-day period within the CY. All participating EPs are required to report on any six eCQMs relevant to their scope of practice from the set of those available (12 of the 24 eCQM are currently available within the 2015 Certified EHR).

For the 2015 Certified RPMS EHR, a total of 16 EH/CAH eCQMs and 12 EP eCQMs were selected for inclusion based on user input. The measures included are as follows:

**Selected EH/CAH eCQM included in the 2015 Certified RPMS EHR (For the 2020 Reporting Period)**

<table>
<thead>
<tr>
<th>Notes: The presence of an asterisk (*) in any of the eCQM entries that follow indicates a new measure added in 2018.</th>
</tr>
</thead>
<tbody>
<tr>
<td>The presence of a pound sign (#) in any of the eCQM entries that follow indicates a 2020 measure.</td>
</tr>
</tbody>
</table>

- CMS9v8 # Newborn Exclusive Breast Milk Feeding
- CMS26v6 Home Management Plan of Care (HMPC) Document Given to Patient/Caregiver
- CMS31v7 Newborn Hearing Screening Prior to Hospital Discharge
- CMS32v8 Median Time from ED Arrival to ED Departure for Discharged ED Patients
- CMS53v7 Primary PCI Received within 90 Minutes of Hospital Arrival
- CMS55v7 Median Time from ED Arrival to ED departure for Admitted ED Patients
- CMS71v9 # Anticoagulation Therapy for Atrial Fibrillation/Flutter
• CMS72v8 # Antithrombotic Therapy By End of Hospital Day 2
• CMS102v7 * Assessed for Rehabilitation
• CMS104v8 # Ischemic stroke-Discharged on Anti-thrombotic Therapy
• CMS105v8 **# Discharged on Statin Medication
• CMS107v7 Ischemic or hemorrhagic stroke-Stroke education
• CMS108v8 **# Venous Thromboembolism Prophylaxis
• CMS111v8 # Median Admit Decision Time to ED Departure Time for Admitted Patients
• CMS113v7 * Elective Delivery
• CMS190v8 **# Intensive Care Unit Venous Thromboembolism Prophylaxis

**Selected EP eCQMs (For the 2020 Reporting Period)**

<table>
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<tr>
<th>Notes: The presence of an asterisk (*) in any of the eCQM entries that follow indicates a new measure added in 2018.</th>
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• CMS2v9 Preventive Care & Screening: Screening for Clinical Depression and Follow-up
• CMS69v8 Preventive Care & Screening: Body Mass Index (BMI) Screening and Follow-up
• CMS117v8 Childhood Immunization Status
• CMS122v8 Hemoglobin A1c (HbA1c) Poor Control (>9%)
• CMS127v8 * Pneumococcal Vaccination Status for Older Adults
• CMS131v8 Eye Exam
• CMS134v8 Diabetes: Medical Attention for Nephropathy
• CMS138v8 Preventive Care & Screening: Tobacco Use: Screening and Cessation Intervention
• CMS139v8 * Falls: Screening for Future Fall Risk
• CMS155v8 Weight Assessment/Counseling for Nutrition/Physical Activity Children/Adolescents
• CMS156v8 Use of High-Risk Medications in the Elderly
• CMS165v8 Controlling High Blood Pressure
While RPMS EHR offers multiple data entry options, eCQM included in this manual present a limited subset of these options in order to efficiently capture the required data elements and achieve the highest possible score. Each eCQM is presented in several views:

- The “data entry view.” This outlines one data entry process to achieve the best possible outcome. These data entry processes have been tested during development and certification.

- A detailed description of each measure from the CMS web site, giving the rationale and logic for each measure.

- A list of the Value Sets for each measure is included in Appendix E. Value sets are groups of codes (Systematized Nomenclature of Medicine-Clinical Terms (SNOMED-CT), International Classification of Diseases (ICD), RxNORM, CPT (Current Procedural Terminology), HCPCS (Healthcare Common Procedure Coding System), etc., that are approved by CMS for each measure. A value set can contain from one to several hundred codes. Only these codes are counted for the measure. CMS updates these value sets periodically, removing some, adding others. IHS OIT updates the terminology used for eCQM based on these changes and distributes terminology updates. It is important to install these updates to be sure that only active and approved codes are being used. OIT “maps” to these new codes to simplify the workload for individual sites.

Since the contents of any given Value Set are fluid and change over time, it is not practical to simply list the currently available codes. Instead, it is better if the user logs into the Value Set Authority Center (VSAC), creates an account (free), and searches for the current values within the set. Past value sets are also viewable for trouble shooting purposes. See Appendix A to learn how to view these Value Sets.

There are six appendices attached:

- Appendix A Using the Value Set Authority Center
- Appendix B Using the United States Health Information Knowledgebase
- Appendix C Using The Electronic Clinical Quality Improvement Center
- Appendix D Using Treatment/Regimen/Follow Up in RPMS EHR
- Appendix E Value Sets
- Appendix F Rules of Behavior
2.0 Hospital Measures for 2020 Reporting Period

2.1 CMS 9v8 Exclusive Breast Milk Feeding in Hospital

2.1.1 Detail

| Description: | Exclusive breast milk feeding during the newborn's entire hospitalization. The measure is reported as an overall rate which includes all newborns that were exclusively fed breast milk during the entire hospitalization. |
| Rationale: | Exclusive breast milk feeding for the first 6 months of neonatal life has long been the expressed goal of World Health Organization (WHO), Department of Health and Human Services (DHHS), American Academy of Pediatrics (AAP) and American College of Obstetricians and Gynecologists (ACOG). ACOG has recently reiterated its position (ACOG, 2007). A recent Cochrane review substantiates the benefits (Kramer & Kakuma, 2002). Much evidence has now focused on the prenatal and intrapartum period as critical for the success of exclusive (or any) BF (Centers for Disease Control and Prevention [CDC], 2007; Petrova, Hegyi, & Mehta, 2007; Shealy et al., 2005; Taveras et al., 2004). Exclusive breast milk feeding rate during birth hospital stay has been calculated by the California Department of Public Health for the last several years using newborn genetic disease testing data. Healthy People 2010 and the CDC have also been active in promoting this goal. |
| Clinical Recommendation Statement: | Exclusive breast milk feeding for the first 6 months of neonatal life can result in numerous long-term health benefits for both mother and newborn and is recommended by a number of national and international organizations. Evidence suggests that the prenatal and intrapartum period is critical for the success of exclusive (or any) breast feeding. Therefore, it is recommended that newborns are fed breast milk only from birth to discharge. |
| Improvement Notation: | Improvement noted as an increase in the rate |
| Guidance | A discharge to a designated cancer center or children's hospital should be captured as a discharge to an acute care facility. It is acceptable to calculate Gestational Age using the American College of Obstetricians and Gynecologists ReVITALize guidelines, which define Gestational Age as calculated using the best obstetrical Estimated Due Date (EDD) based on the formula: Gestational Age= (280-(EDD-Reference Date))/7 where Reference Date is the date on which you are trying to determine gestational age. For PC-05, Reference Date is the Birth Date. Note however that the calculation may yield a non-whole number and gestational age should be rounded off to the nearest completed week. For example, an infant born on the 5th day of the 36th week (35 weeks and 5/7 days) is at a gestational age of 35 weeks, not 36 weeks. |
Initial Population: Inpatient hospitalizations for single newborns with an estimated gestational age at birth of >=37 weeks who are born in the hospital and who did not have a diagnosis of galactosemia, were not subject to parenteral nutrition, and had a length of stay less than or equal to 120 days that ends during the measurement period.

Denominator Initial Population

Denominator Exclusions Inpatient hospitalizations for newborns who were admitted to the Neonatal Intensive Care Unit (NICU), who were transferred to an acute care facility, or other health care facility, or who expired during the hospitalization.

Numerator Inpatient hospitalizations for newborns who were fed breast milk only since birth.

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.1.2 Data Entry
1. Admit patient to Nursery.
2. Diagnosis “Single Live Birth Born in Hospital.”
3. Gestational Age = or > than 37 weeks (document triage tab, personal health, birth measurements), OR Birth Weight >= 3000 gm.

![Image of Update Birth Measurements]

Figure 2-1: Update Birth Measurements
4. Excluded if diagnosis “Galactosemia,” parenteral nutrition, or prolonged length of stay (>120 days).

5. Excluded if admitted to NICU, expires during hospitalization, or transfer to acute care facility.

6. Enter **Infant Feeding** on **Triage** tab, **Exclusive Breast Feeding** to include in numerator.

![Figure 2-2: Triage tab](image)

2.2 **CMS 26v6 Home Management Plan of Care Document Given to Patient/Caregiver**

2.2.1 **Detail**

| Description: | An assessment that there is documentation in the medical record that a Home Management Plan of Care (HMPC) document was given to the pediatric asthma patient/caregiver |
**Rationale:**

Asthma is the most common chronic disease in children and a major cause of morbidity and health care costs nationally (Adams, et al, 2001). In 2005, 5.2% of children with asthma had at least one asthma attack in the previous year (3.8 million children). Nearly two of every three children who currently have asthma had at least one attack in the past 12 months. Chronic asthma in children can account for an annual loss of more than 14 million school days per year, according to the Asthma and Allergy Foundation (Asthma Facts and Figures).

It is clear from multiple sources of evidence including the National Heart Lung and Blood Institute (NHLBI) Guidelines that actual self-management of asthma by the patient or caregiver leads to more positive outcomes. Appropriate self-management is completely reliant upon patient education. Patient education is more effective when it aims at training self-management skills that will alter behavior (Norris, et al, 2001).

**Rationale (cont.)**

NHLBI notes that review of asthma management by expert clinicians is necessary but not sufficient to improve outcomes. Active learning, participating and verbalization of understanding are all strategies that a healthcare organization must incorporate with parents or caregivers of asthmatic children in order for them to understand and make the appropriate changes that can impact the disease in the child in question. Education programs have been effective in improving lung function, feelings of self-esteem, and consequently decreased missed days of school in children and adolescents (Phipatanakul, 2004). Acute hospitalization follow up is imperative to a successful discharge from the hospital, providing the caretaker with the resource information needed to contact the follow up facility, medical office or clinic setting (Schatz, et al, 2009).

Environmental control consists of removal of asthma triggers from the environment. Multiple studies support the positive correlation of household maintenance factors such as control of cockroach dust, and the number of acute asthma attacks in asthmatic children (McConnell, et al, 2005 and Eggleston, et al, 2005). Evidence from Carter et al, (2001) supported by the National Institute of Health (NIH) grant found specifically that reduction in triggers such as household conditions i.e., dust mites, cockroach, cats and presence of molds and fungus, resulted in a decrease in acute care visits and an overall positive outcome of children.

Rescue action education related to early recognition of symptoms and proper action to control incidence of asthma attacks is noted to have positive outcomes for asthmatic children (Ducharme and Bhogal, 2008).

**Clinical Recommendation Statement:**

Under-treatment and/or inappropriate treatment of asthma are recognized as major contributors to asthma morbidity and mortality. National guidelines for the diagnosis and management of asthma in children, recommend establishing a plan for maintaining control of asthma and for establishing plans for managing exacerbation.
### Guidance:

The home management plan of care document should be a separate and patient-specific written instruction. The document must be present in the form of an explicit and separate document specific to the patient rather than components or segments of the plan spread across discharge instruction sheets, discharge orders, education sheets, or other instruction sheets.

The home management plan of care is represented in the eMeasure logic by a Logical Observation Identifiers Names and Codes (LOINC) for an asthma action plan document. This form, or equivalent, contains most of the components required for the home management plan of care, including information on:

- **Methods and timing of rescue actions:** the home management plan of care addresses what to do if asthma symptoms worsen after discharge, including all of the following:
  - When to take action, i.e., assessment of severity (e.g., peak flow meter reading, signs and symptoms to watch for).
  - What specific steps to take, i.e., initial treatment instructions (e.g., inhaled relievers up to three treatments of 2-4 puffs by MDI at 20-minute intervals or single nebulizer treatment).
  - Contact information to be used, when an asthma attack occurs or is about to occur.
  - Appropriate use of long-term asthma medications (controllers), including the medication name, dose, frequency, and method of administration.
  - Appropriate use of rescue, quick-relief, or short acting medications of choice to quickly relieve asthma exacerbations (relievers), including the medication name, dose, frequency, and method of administration.
  - Environmental control and control of other triggers: information on avoidance or mitigation of environmental and other triggers.

The home management plan of care can only be considered to comply with the criteria outlined in the measure logic if it meets the requirements outlined above and is appropriately filled-out with information specific to the patient.

Patient refusal includes refusal by a caregiver. The caregiver is defined as the patient's family or any other person (e.g., home health, VNA provider, prison official or other law enforcement personnel) who will be responsible for care of the patient after discharge.

The "Discharge To Home Or Police Custody" value set also intends to capture the following discharge disposition values:

- Assisted Living Facilities
- Court/Law Enforcement - includes detention facilities, jails, and prison
- Home - includes board and care, foster or residential care, group or personal care homes, and homeless shelters
- Home with Home Health Services
- Outpatient Services including outpatient procedures at another hospital, Outpatient Chemical Dependency Programs and Partial Hospitalization.
Initial Population: Pediatric asthma inpatients with an age of 2 through 17 years, and length of stay less than or equal to 120 days that ends during the measurement period.

Denominator: Patients discharged to home or police custody.

Denominator Exclusions: None

Numerator: Pediatric asthma inpatients with either of the following:
- Documentation that they or their caregivers were given a written Home Management Plan of Care (HMPC) document that addresses all of the following:
  - Arrangements for follow-up care
  - Environmental control and control of other triggers
  - Method and timing of rescue actions
  - Use of controllers
  - Use of relievers
- Pediatric asthma inpatients with documentation that they or their caregivers refused a written Home Management Plan of Care (HMPC) document. Patient refusal includes refusal by a caregiver. The caregiver is defined as the patient's family or any other person (e.g., home health, VNA provider, prison official or other law enforcement personnel) who will be responsible for care of the patient after discharge.

Numerator Exclusions: Not Applicable

Denominator Exceptions: None

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.2.2 Data Entry

Description: An assessment that there is documentation in the medical record that a Home Management Plan of Care (HMPC) document was given to the pediatric asthma patient/caregiver.

1. Admit Patient 2-17 y/o to hospital, diagnosis = Asthma (Inpatient coder codes).

![Figure 2-3: Integrated Problem List](image)

Figure 2-4: Self-Management Plan

Figure 2-5: Edit Patient Education Event dialog
3. When Discharged, discharge to Home or Police Custody (ADT Package), hospital stay less than 120 days.

### 2.3 CMS 31v7 Hearing Screening Prior To Hospital Discharge

#### 2.3.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>This measure assesses the proportion of births that have been screened for hearing loss before hospital discharge.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Birthing facility staff should review the effectiveness and timeliness of screening relative to nursery discharge. Benchmarks set within the EHCP may trigger hospital or jurisdictional compliance activities, such as re-writing of procedural guidelines or re-training of screening staff.</td>
</tr>
<tr>
<td>Improvement Notation:</td>
<td>Improvement noted as an increase in rate.</td>
</tr>
<tr>
<td>Guidance:</td>
<td>Complete hearing screening is defined as having both ears screened during the inpatient encounter. The Medical Reasons value set (OID 2.16.840.1.114222.4.1.214079.1.1.7) includes SNOMED codes to capture medical contraindications, surgical contraindications, and not indicated because there is a reason to withhold a certain medical procedure/treatment because it could harm a patient. A comprehensive list of codes for these conditions is not available because the judgement is often subjective. This value set is not intended to be used for other reasons a screening was not performed, such as parent refusal, technical failure, or a missed procedure.</td>
</tr>
</tbody>
</table>
Transmission Format: TBD

Initial Population: Live birth encounters at a hospital or birthing facility where the newborn was discharged with hospital stays \(\leq 120\) days that ends during the measurement period.

Denominator: Denominator is equal to the Initial Population.

Denominator Exclusions: A live birth encounter where the newborn expires prior to discharge and has not received a complete hearing screening.

Numerator: A live birth encounter where a complete newborn hearing screening is performed prior to discharge or the newborn is not screened due to medical reasons.

Numerator Exclusions: Not applicable

Denominator Exceptions: None

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.3.2 Data Entry

Description: This measure assesses the proportion of births that have been screened for hearing loss before hospital discharge.

1. Admit patient, Diagnosis = Live Newborn Born in Hospital

2. During hospitalization, do Hearing Left and Right (Wellness tab, Exams, Newborn Hearing Screen, enter Pass or Referral Needed for both ears).
3. If **not** done for Medical reasons:

   a. On Triage tab, **Personal Health**, select **Refusal**.

   b. Select **Refusal Type = Exam**.

   c. Select **Newborn Hearing Screen (left and right)**.
d. Choose reason for refusal = Contraindicated.

4. Discharge patient (must be less than 120 days and end during measurement period).

2.4 CMS 32v8 Median Time from ED Arrival to ED Departure for Discharged ED Patients

2.4.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Median elapsed time from ED arrival to Emergency Room (ER) departure for patients discharged from the ED.</th>
</tr>
</thead>
</table>
| Stratification: | Report total score and the following strata:
 | Stratification 1 - all patients with principal diagnosis consistent with mental disorders.
 | Stratification 2 - all patients transferred to another acute care hospital.
 | Stratification 3 - all patients who do not have a principal diagnosis consistent with mental disorders and who are not transferred. |
### Rate Aggregation:

Calculate the ED encounter duration at the facility in minutes for each ED encounter in the measure population; report the median time for all calculations performed. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the length of time the patient was in the ED, also stated as: the Datetime difference between the ED facility location departure time and the ED facility location arrival time. The calculation requires the median across all ED encounter durations.

### Rationale:

In recent times, EDs have experienced significant overcrowding. Although once only a problem in large, urban, teaching hospitals, the phenomenon has spread to other suburban and rural healthcare organizations. According to a 2002 national U.S. survey, more than 90 percent of large hospitals report EDs operating "at" or "over" capacity. Overcrowding and heavy emergency resource demand have led to a number of problems, including ambulance refusals, prolonged patient waiting times, increased suffering for those who wait, rushed and unpleasant treatment environments, and potentially poor patient outcomes. Approximately one third of hospitals in the U.S. report increases in ambulance diversion in a given year, whereas up to half report crowded conditions in the ED. In a recent national survey, 40 percent of hospital leaders viewed ED crowding as a symptom of workforce shortages. ED crowding may result in delays in the administration of medication such as antibiotics for pneumonia and has been associated with perceptions of compromised emergency care. For patients with non-ST-segment-elevation myocardial infarction, long ED stays were associated with decreased use of guideline-recommended therapies and a higher risk of recurrent myocardial infarction. When EDs are overwhelmed, their ability to respond to community emergencies and disasters may be compromised.

### Clinical Recommendation Statement:

Reducing the time patients remain in the ED can improve access to treatment and increase quality of care. Reducing this time potentially improves access to care specific to the patient condition and increases the capability to provide additional treatment.
Guidance: This measure uses a continuous variable. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the length of time the patient was in the ED, also stated as: the ED departure time minus the ED arrival time. The calculation requires the median of all ED encounter durations. This measure specification defines how to determine an individual ED stay. Reporting requires the median of all patient stays ([Encounter: encounter ED].facility location department date and time minus [Encounter: encounter ED].facility location arrival date and time).

For each population, results should be reported without stratification and then with each stratum applied. For this measure, the number of encounters that fall into the Initial Population are reported without stratification, then reported according to the defined stratification. The number of encounters that fall into the Measure Population are reported without stratification, then reported according to the defined stratification. The computed continuous variable defined by the Measure Observation is reported for the Measure Population also, then reported according to the defined stratification.

Initial Population: ED encounters during the measurement period.

Measure Population Exclusions: ED encounters where the patient expired during the encounter or where the ED visit is followed within an hour by an inpatient encounter at the same physical facility.

Measure Observations: Median elapsed time (in minutes) from ED arrival to ER departure or for patients placed in observation services, use the time of the order for observation for ED departure for patients discharged from the ED.

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.4.2 Data Entry

Description: Median elapsed time from ED arrival to ER departure for patients discharged from the ED.

Admit to ER (AMER or ED Dashboard):
- Exclude expired patients (Discharge Status = patient expired)
- Exclude admitted patients (ADT package)

Discharge from ER – AMER or ED Dashboard - timestamps discharge time:
- Enter Primary Purpose of Visit (POV) through ER IPL; enter disposition
**Stratification 1:** Patients with Mental Health Diagnosis (Set as POV) in IPL (see value sets)

**Stratification 2:** Discharge status = Discharge (transfer) to another Acute Care Facility

**Stratification 3:** Patients discharged alive and neither of above applies.

### 2.5 CMS 53v7 Primary PCI Received Within 90 Minutes of Hospital Arrival

#### 2.5.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Acute myocardial infarction (AMI) patients with a diagnostic Electrocardiogram (ECG) and ST-segment elevation, who receive a primary percutaneous coronary intervention (PCI) during their hospital stay, and the time from hospital arrival to PCI is 90 minutes or less</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>The early use of primary angioplasty in patients with ST-segment myocardial infarction (STEMI) results in a significant reduction in mortality and morbidity. The earlier primary coronary intervention is provided, the more effective it is (Brodie, 1998 and DeLuca, 2004). National guidelines recommend the prompt initiation of PCI in patients presenting with ST-elevation myocardial infarction (O’Gara, 2013; and Levine 2011).</td>
</tr>
<tr>
<td>Guidance:</td>
<td>To identify the ECG closest to hospital arrival, the denominator will identify the most recent ECG performed that starts less than 1 hour prior to the ED or inpatient encounter OR the first ECG performed starts after the ED or inpatient encounter started. PCI procedures analyzed in this measure must be primary procedures. Primary includes emergent or urgent PCI procedures and not those described by the physician/APN/PA anywhere in the record as elective, not emergent, not immediate, not primary, not urgent, or secondary. From a clinical standpoint, Primary PCI is loosely defined as percutaneous coronary intervention performed in the acute setting in patients with ST-segment elevation MI which is intended to restore perfusion in the infarct-related artery.</td>
</tr>
<tr>
<td>Initial Population:</td>
<td>Patients age 18 and older at the time of hospital admission with a principal diagnosis of an Acute or Evolving Myocardial Infarction (ST-segment elevation MI) with hospital stays &lt;= 120 days during the measurement period</td>
</tr>
<tr>
<td>Denominator:</td>
<td>Initial population with an ECG performed and a primary PCI procedure closest to the inpatient admission that does not start after fibrinolytic therapy</td>
</tr>
</tbody>
</table>
Denominator Exclusions: Patients who transferred from another hospital's inpatient, outpatient, or ED and ambulatory surgery center facilities are excluded

Numerator: Initial population whose time from hospital arrival to primary PCI is 90 minutes or less

Numerator Exclusions: Not applicable

Denominator Exceptions: Patients who did not receive PCI within 90 minutes and have a documented reason. Documented reasons include:
"Diagnosis: Cardiopulmonary arrest"
"Procedure, Performed: Endotracheal Intubation"
"Procedure, Performed: Aortic balloon pump insertion"
"Procedure, Performed: Ventricular Assist Device placement"

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex

2.5.2 Data Entry

Description: AMI patients with a diagnostic ECG and ST-segment elevation, who receive a primary PCI during their hospital stay, and the time from hospital arrival to PCI is 90 minutes less.

Admitted patients, 18 or older, with principal diagnosis of acute or evolving myocardial infarction (ST-elevated MI, “STEMI”), LOS <120 days ending during measurement period. Primary POV entered by Inpatient Coder after discharge.

Denominator also requires ECG performed AND PCI begun before fibrinolytic therapy given (timestamp when finished by pharmacy, in ER or inpatient).

Denominator exclusions: Patients transferred from another facility (ER, IP, OPD, or ambulatory surgery center).

Numerator: Patients with time from admission to primary PCI 90 minutes or less (AMI Tool allows entry of times for EKG, PCI in ER or inpatient).
Figure 2-13: Problem Management AMI tool

- Check **ECG Done.** Time and add finding.
- Check “fibrinolytic therapy not initiated.”
- Check **PCI Done.** This captures time. Procedure must still be entered in IPL for outpatient or by Inpatient Coder.

Denominator exceptions (documented reasons for not providing PCI):

- Cardiopulmonary arrest
- Endotracheal intubation
- Aortic balloon pump insertion
- Ventricular assist device placement

### 2.6 CMS 55v7 Median Time from ED Arrival to ED Departure for Admitted ED Patients

#### 2.6.1 Detail

| Description: | Median time from ED arrival to time of departure from the ER for patients admitted to the facility from the ED |
**Stratification:**
Report total score and the following strata:
Stratification 1 - all patients seen in the ED and admitted as an inpatient who do not have an inpatient encounter principal diagnosis consistent with psychiatric/mental health disorders
Stratification 2 - all patients seen in the ED and admitted as an inpatient who have an inpatient encounter principal diagnosis consistent with psychiatric/mental health disorders

**Rate Aggregation:**
Calculate the ED encounter duration in minutes for each ED encounter in the measure population; report the median time for all calculations performed. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the length of time the patient was in the ED, also stated as: the Datetime difference between the ED facility location departure date/time and the ED facility location arrival date/time. The calculation requires the median across all ED encounter durations.

**Rationale:**
In recent times, EDs have experienced significant overcrowding. Although once only a problem in large, urban, teaching hospitals, the phenomenon has spread to other suburban and rural healthcare organizations. According to a 2002 national U.S. survey, more than 90% of large hospitals report EDs operating "at" or "over" capacity. Approximately one third of hospitals in the US report increases in ambulance diversion in a given year, whereas up to half report crowded conditions in the ED. In a recent national survey, 40% of hospital leaders viewed ED crowding as a symptom of workforce shortages. ED crowding may result in delays in the administration of medication such as antibiotics for pneumonia and has been associated with perceptions of compromised emergency care. For patients with non-ST-segment-elevation myocardial infarction, long ED stays were associated with decreased use of guideline-recommended therapies and a higher risk of recurrent myocardial infarction. Overcrowding and heavy emergency resource demand have led to a number of problems, including ambulance refusals, prolonged patient waiting times, increased suffering for those who wait, rushed and unpleasant treatment environments, and potentially poor patient outcomes. When EDs are overwhelmed, their ability to respond to community emergencies and disasters may be compromised.

**Clinical Recommendation Statement:**
Reducing the time patients remain in the ED can improve access to treatment and increase quality of care. Reducing this time potentially improves access to care specific to the patient condition and increases the capability to provide additional treatment.
This measure specification defines how to determine the duration of an individual ED encounter. Reporting requires the median of all ED encounter durations defined as [Encounter: encounter ED] facility location departure date and time minus [Encounter: encounter ED] ED facility location arrival date and time.

Calculate the ED time in minutes for each patient in the measure population; report the median time for all calculations performed. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the length of time the patient was in the ED, also stated as: the Datetime difference between the ED facility location departure datetime and the ED facility location arrival datetime. The calculation requires the median across all ED encounter durations.

For each population, results should be reported without stratification and then with each stratum applied. For this measure, the number of encounters that fall into the Initial Population are reported without stratification, then reported according to the defined stratification. The number of encounters that fall into the Measure Population are reported without stratification, then reported according to the defined stratification. The computed continuous variable defined by the Measure Observation is reported for the Measure Population also, then reported according to the defined stratification.

**Initial Population:**
Inpatient Encounters ending during the measurement period with Length of Stay (Discharge Date minus Admission Date) less than or equal to 120 days, and preceded within an hour by an ED visit at the same physical facility

**Measure Population:**
Initial Population

**Measure Population Exclusions:**
ED encounters with an admission source from another "Hospital Setting" (any different facility, even if part of the same hospital system) resulting in an inpatient stay

**Measure Observations:**
Time (in minutes) from ED facility location arrival to ED facility location departure for patients admitted to the facility from the ED

**Supplemental Data Elements:**
For every patient evaluated by this measure, also identify payer, race, ethnicity, and sex.

### 2.6.2 Data Entry

**Description:**
Median time from ED arrival to time of departure from the ER for patients admitted to the facility from the ED.

Admit patient to ER, use AMER or ED Dashboard to timestamp arrival time (if patient transferred to ER from hospital settings within 6 hours, they are not included in Denominator for measure)
When patient discharged in AMER or ED Dashboard (timestamps discharge time) and admitted from ER, the time between ER discharge and Inpatient must be less than 1 hour to be included. Enter “Admit” for disposition

Stratification 1: POV in ER is not mental health related (see value sets)

Stratification 2: POV in ER is mental health related

### 2.7 CMS 71v9 Anticoagulation Therapy for Atrial Fibrillation/Flutter

#### 2.7.1 Detail

<table>
<thead>
<tr>
<th>Description.</th>
<th>Ischemic stroke patients with atrial fibrillation/flutter who are prescribed or continuing to take anticoagulation therapy at hospital discharge.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale.</td>
<td>Nonvalvular atrial fibrillation (NVAF) is a common arrhythmia and an important risk factor for stroke. It is one of several conditions and lifestyle factors that have been identified as risk factors for stroke. It has been estimated that over 2 million adults in the United States have NVAF. While the median age of patients with atrial fibrillation is 75 years, the incidence increases with advancing age. For example, The Framingham Heart Study noted a dramatic increase in stroke risk associated with atrial fibrillation with advancing age, from 1.5% for those 50 to 59 years of age to 23.5% for those 80 to 89 years of age. Furthermore, a prior stroke or transient ischemic attack (TIA) are among a limited number of predictors of high stroke risk within the population of patients with atrial fibrillation. Therefore, much emphasis has been placed on identifying methods for preventing recurrent ischemic stroke as well as preventing first stroke. Prevention strategies focus on the modifiable risk factors such as hypertension, smoking, and atrial fibrillation. Analysis of five placebo-controlled clinical trials investigating the efficacy of warfarin in the primary prevention of thromboembolic stroke, found the relative risk of thromboembolic stroke was reduced by 68% for atrial fibrillation patients treated with warfarin. The administration of anticoagulation therapy, unless there are contraindications, is an established effective strategy in preventing recurrent stroke in high stroke risk-atrial fibrillation patients with TIA or prior stroke.</td>
</tr>
<tr>
<td>Clinical Recommendation Statement.</td>
<td>The administration of anticoagulation therapy, unless there are contraindications, is an established effective strategy in preventing recurrent stroke in high stroke risk atrial fibrillation patients with TIA or prior stroke.</td>
</tr>
</tbody>
</table>
**Guidance.**

The "Non-elective Inpatient Encounter" value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the "Inpatient Encounter" value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective Inpatient Encounters include emergency, urgent, and unplanned admissions. The "Medication, Discharge" datatype refers to the discharge medication list and is intended to express medications ordered for post-discharge use.

**Initial Population.**

Inpatient hospitalizations for patients age 18 and older, discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during measurement period.

**Denominator.**

Inpatient hospitalizations for patients with a principal diagnosis of ischemic stroke, and a history of atrial ablation, or current or history of atrial fibrillation/flutter.

**Denominator Exclusions.**

Inpatient hospitalizations for patients admitted for elective carotid intervention. This exclusion is implicitly modeled by only including non-elective hospitalizations. Inpatient hospitalizations for patients discharged to another hospital. Inpatient hospitalizations for patients who left against medical advice. Inpatient hospitalizations for patients who expired. Inpatient hospitalizations for patients discharged to home for hospice care. Inpatient hospitalizations for patients discharged to a health care facility for hospice care. Inpatient hospitalizations for patients with Comfort measures documented.

**Numerator.**

Inpatient hospitalizations for patients prescribed or continuing to take anticoagulation therapy at hospital discharge.

**Denominator Exceptions.**

Inpatient hospitalizations for patients with a documented reason for not prescribing anticoagulation therapy at discharge.

**Supplemental Data Elements.**

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 2.7.2 Data Entry

**Description:**

Ischemic stroke patients with atrial fibrillation/flutter who are prescribed or continuing to take anticoagulation therapy at hospital discharge.

Admit patient (non-elective, patient registration), age 18+. Includes observation.

Diagnosis: Principle diagnosis = Ischemic Stroke plus EITHER atrial ablation, or current atrial flutter/fibrillation (can enter PCC post discharge).

Prescribe discharge anticoagulant medication

Value set: 2.16.840.1.113883.3.117.1.7.1.200

Discharge with diagnosis = Ischemic Stroke.
Denominator exclusions:

- Discharge to acute care facility (ADT package, d/c status).
- Left against medical advice (AMA) (ADT package, d/c status).
- Expired (ADT package, type of discharge).
- Discharged to home or healthcare facility for hospice care (ADT package, type of discharge).
- Comfort measures –document in “Palliative Procedures” in IPL/TREG (Edit POV, Add Visit Instructions/Care Plans/Goal Activities, Treatment/Regimen/Follow-up, Palliative Care).

Figure 2-14: Palliative Care list

Denominator Exception:

- Patients who refuse or cannot take anticoagulant medication.
- Document on Wellness tab, Personal Health, Patient refusals, type = medication, reason contraindicated or patient refusal.

Numerator:

- Prescribe anticoagulant, mark as “discharge medication”, finish in pharmacy.

2.8 CMS 72v8 Antithrombotic Therapy End of Hospital Day 2

2.8.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Ischemic stroke patients administered antithrombotic therapy by the end of hospital day 2.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>The effectiveness of antithrombotic agents in reducing stroke mortality, stroke-related morbidity and recurrence rates has been studied in several large clinical trials. While the use of these agents for patients with acute ischemic stroke and transient ischemic attacks continues to be the subject of study, substantial evidence is available from completed studies. Data at this time suggest that antithrombotic therapy should be administered within 2 days of symptom onset in acute ischemic stroke patients to reduce stroke mortality and morbidity as long as no contraindications exist. Anticoagulants at doses to prevent venous thromboembolism are insufficient antithrombotic therapy to prevent recurrent ischemic stroke or TIA.</td>
</tr>
</tbody>
</table>
Clinical Recommendation Statement: Antithrombotic therapy should be administered within 2 days of symptom onset in acute ischemic stroke patients to reduce stroke mortality and morbidity as long as no contraindications exist.

Guidance: The "Non-elective Inpatient Encounter" value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the "Inpatient Encounter" value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective Inpatient Encounters include emergency, urgent, and unplanned admissions. NPO is not a valid reason for not administering antithrombotic therapy by end of hospital day 2 as another route of administration can be used (i.e., rectal or intravenous). In the denominator exclusions, the intent is to only exclude patients with a total length of stay of <2 days, including ED visit (if there is one). For the eCQM we model both of the scenarios of admission via the ED as well as direct admits. This statement addresses direct admits.

Initial Population: Inpatient hospitalization for patients age 18 and older discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during measurement period.

Denominator: Inpatient hospitalizations for patients with a principal diagnosis of ischemic stroke.

Denominator Exclusions: inpatient hospitalizations for patients who have a duration of stay less than 2 days...Inpatient hospitalization for patients with an INR greater than 3.5.

Numerator: Inpatient hospitalization for patients who had antithrombotic therapy administered the day of or day after hospital arrival.

Denominator Exceptions: Inpatient hospitalization for patients with a documented reason for not administering antithrombotic therapy the day of or day after hospital arrival.

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.8.2 Data Entry

Description: Ischemic stroke patients administered antithrombotic therapy by the end of hospital day 2.

Admit patient, age over 18, diagnosis “Ischemic Stroke.” Includes observation.
Duration of hospitalization (including ER) over 2 days and less than 120 days (admission from ER within one hour of ER Discharge).

Excludes patients with documented “Comfort measures” on admission or day 1 (document hospice care on IPL/TREG).

Excludes patients who received thrombolytic therapy during hospital stay or within 24 hours before admission. Documented by med order or POV “IV infusion of thrombolytic.”

Exclude patients with INR>3.5 (INR Value set: 2.16.840.1.113883.3.117.1.7.1.213) Documented administration of antithrombotic medication (Value Set 2.16.840.1.113883.3.117.1.7.1.201) during first 2 days of admission (order med, finish in pharmacy, administer with BCMA); use “Stroke Tool.”

Discharge dx = Ischemic stroke.

2.9 CMS 102v7 Assessed for Rehabilitation

2.9.1 Detail

| Description: | Ischemic or hemorrhagic stroke patients who were assessed for rehabilitation services. |
### Rationale:
Each year about 700,000 people experience a new or recurrent stroke, which is the nation’s third leading cause of death. Approximately two thirds of these individuals survive and require rehabilitation. Stroke is a leading cause of serious, long-term disability in the United States, with about 4.4 million stroke survivors alive today. Forty percent of stroke patients are left with moderate functional impairment and 15 to 30 percent with severe disability. More than 60% of those who have experienced stroke, serious injury, or a disabling disease have never received rehabilitation. Stroke rehabilitation should begin as soon as the diagnosis of stroke is established, and life-threatening problems are under control. Among the high priorities for stroke are to mobilize the patient and encourage resumption of self-care activities as soon as possible. A considerable body of evidence indicates better clinical outcomes when patients with stroke are treated in a setting that provides coordinated, multidisciplinary stroke-related evaluation and services. Effective rehabilitation interventions initiated early following stroke can enhance the recovery process and minimize functional disability. The primary goal of rehabilitation is to prevent complications, minimize impairments, and maximize function.

### Guidance
The "Non-elective Inpatient Encounter" value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the "Inpatient encounter" value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective admissions include emergency, urgent and unplanned admissions.

### Clinical Recommendation Statement:
A considerable body of evidence indicates better clinical outcomes when patients with stroke are treated in a setting that provides coordinated, multidisciplinary stroke-related evaluation and services. Effective rehabilitation interventions initiated early following stroke can enhance the recovery process and minimize functional disability.

### Initial Population:
Patients age 18 and older discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during the measurement period.

### Denominator:
Initial Population.

### Denominator Exclusions:
- Patients with Comfort measures documented.
- Patients discharged to another hospital.
- Patients who left against medical advice.
- Patients who expired.
- Patients discharged to home for hospice care.
- Patients discharged to a health care facility for hospice care.
- Patients admitted for elective carotid intervention. This exclusion is implicitly modeled by only including non-elective hospitalizations.

### Numerator:
Ischemic or hemorrhagic stroke patients assessed for or who received rehabilitation services.
### 2.9.2 Data Entry

**Description:** Ischemic or hemorrhagic stroke patients who were assessed for rehabilitation services.

Admit, patient over age 18, diagnosis = Ischemic Stroke or Hemorrhagic Stroke.

- Excludes patients discharged to acute care facility, leave AMA, patient expires, discharged to hospice care (home or facility) (Document in ADT package discharge status).
- Excludes patients with documented “Comfort measures” (TREG).

![Figure 2-17: Palliative Care list](image)

- Includes patients discharged to rehab facility.

Numerator:

Document Rehabilitation Therapy or Rehabilitation Assessment, IPL, TREG, Rehab Services:

![Figure 2-18: Rehab Services list](image)

Numerator includes refusals (patient or medical reason); document in Personal Health component, select **SNOMED** and select one of several options:

- **Assessment of mobility**
- **Diagnostic physical therapy**
- **Occupational therapy assessment**
- **Physical therapy assessment**
Figure 2-19: Service Refusal

Figure 2-20: SNOMED CT Lookup

Figure 2-21: Service Refusal

Figure 2-22: Personal Health tab
### 2.10 CMS 104v8 Discharged on Antithrombotic Therapy

#### 2.10.1 Detail

<table>
<thead>
<tr>
<th><strong>Description:</strong></th>
<th>Ischemic stroke patients prescribed or continuing to take antithrombotic therapy at hospital discharge.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rationale:</strong></td>
<td>The effectiveness of antithrombotic agents in reducing stroke mortality, stroke-related morbidity and recurrence rates has been studied in several large clinical trials. While the use of these agents for patients with acute ischemic stroke and transient ischemic attacks continues to be the subject of study, substantial evidence is available from completed studies. Data at this time suggest that antithrombotic therapy should be prescribed at discharge following acute ischemic stroke to reduce stroke mortality and morbidity as long as no contraindications exist. For patients with a stroke due to a cardioembolic source (e.g., atrial fibrillation, mechanical heart valve), warfarin is recommended unless contraindicated. In recent years, novel oral anticoagulant agents (NOACs) have been developed and approved by the U.S. Food and Drug Administration (FDA) for stroke prevention and may be considered as an alternative to warfarin for select patients. Anticoagulation therapy is not generally recommended for secondary stroke prevention in patients presumed to have a non-cardioembolic stroke. Anticoagulants at doses to prevent venous thromboembolism are insufficient antithrombotic therapy to prevent recurrent ischemic stroke or TIA.</td>
</tr>
<tr>
<td><strong>Clinical Recommendation Statement:</strong></td>
<td>Clinical trial results suggest that antithrombotic therapy should be prescribed at discharge following acute ischemic stroke to reduce stroke mortality and morbidity as long as no contraindications exist.</td>
</tr>
<tr>
<td><strong>Guidance</strong></td>
<td>The &quot;Non-elective Inpatient Encounter&quot; value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the &quot;Inpatient encounter&quot; value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective admissions include emergency, urgent and unplanned admissions. The &quot;Medication, Discharge&quot; datatype refers to the discharge medication list and is intended to express medications ordered for post-discharge use.</td>
</tr>
<tr>
<td><strong>Initial Population:</strong></td>
<td>Inpatient hospitalizations for patients age 18 and older, discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during the measurement period</td>
</tr>
<tr>
<td><strong>Denominator:</strong></td>
<td>Inpatient hospitalizations for patients with a principal diagnosis of Ischemic stroke.</td>
</tr>
</tbody>
</table>
Denominator Exclusions:

Inpatient hospitalizations for patients admitted for elective carotid intervention. This exclusion is implicitly modeled by only including non-elective hospitalizations. Inpatient hospitalizations for patients discharged to another hospital. Inpatient hospitalizations for patients who left against medical advice. Inpatient hospitalizations for patients who expired. Inpatient hospitalizations for patients discharged to home for hospice care. Inpatient hospitalizations for patients discharged to a health care facility for hospice care. Inpatient hospitalizations for patients with Comfort measures documented.

Numerator:

Inpatient hospitalizations for patients prescribed or continuing to take antithrombotic therapy at hospital discharge.

Numerator Exclusions:

Not Applicable.

Denominator Exceptions:

Inpatient hospitalizations for patients with a documented reason for not prescribing antithrombotic therapy at discharge. Inpatient hospitalizations for patients who receive Ticagrelor as an antithrombotic therapy during the hospitalization.

Supplemental Data Elements:

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.10.2 Data Entry

Description:

Ischemic stroke patients prescribed or continuing to take antithrombotic therapy at hospital discharge.

- Admit (non-elective), age over 18, diagnosis = Ischemic Stroke. Includes observation services (Value set: 2.16.840.1.113762.1.4.1111.143)
- Excludes patients with documented “Comfort Measures” (IPL/TREG).

Figure 2-23: Palliative Care list

- Excludes patients discharged to acute care facility (ADT package, d/c), patient expires, left AMA, or discharged for hospice care (home or facility).
- Exceptions for patients who refuse antithrombotic (patient refusal or medical contraindication), enter in Personal Health Component, medication.
- Numerator requires prescription for antithrombotic medication (list found in Value Set: 2.16.840.1.113883.3.117.1.7.1.201, see Appendix A) during hospitalization, marked as Discharge Medication.)
2.11 CMS 105v8 Discharged on Statin Medication

2.11.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Ischemic stroke patients who are prescribed or continuing to take statin medication at hospital discharge.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>There is an extensive and consistent body of evidence supporting the use of statins for secondary prevention in patients with clinically evident atherosclerotic cardiovascular disease (ASCVD), which includes individuals with ischemic stroke due to large artery atherosclerosis, individuals with ischemic stroke due to intrinsic small vessel disease, and individuals with ischemic stroke not directly due to atherosclerosis but with clinically evident atherosclerotic disease in an uninvolved cerebral or noncerebral bed. Both women and men with clinical ASCVD are at increased risk for recurrent ASCVD and ASCVD death. High-intensity statin therapy should be initiated or continued as first-line therapy in women and men less than or equal to 75 years of age who have clinical ASCVD, unless contraindicated. In patients with clinical ASCVD and a contraindication to high-intensity statin therapy, moderate-intensity therapy should be considered as an alternative if it can be tolerated. In individuals greater than 75 years of age, the potential for ASCVD risk reduction benefits, adverse effects, drug-drug interactions, and patient preferences should be considered, and statin therapy individualized based on these considerations (Stone, 2013).</td>
</tr>
<tr>
<td>Guidance</td>
<td>The &quot;Non-elective Inpatient Encounter&quot; value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the &quot;Inpatient Encounter&quot; value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective Inpatient Encounters include emergency, urgent, and unplanned admissions. The &quot;Medication, Discharge&quot; datatype refers to the discharge medication list and is intended to express medications ordered for post-discharge use.</td>
</tr>
<tr>
<td>Clinical Recommendation Statement:</td>
<td>For patients with stroke of atherosclerotic origin, intensive lipid lowering therapy with statins should be initiated.</td>
</tr>
</tbody>
</table>
**Initial Population:** Inpatient hospitalizations for patients age 18 and older, discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during measurement period

**Denominator:** Inpatient hospitalizations for patients with a principal diagnosis of ischemic stroke

**Denominator Exclusions:** Inpatient hospitalizations for patients admitted for elective carotid intervention. This exclusion is implicitly modeled by only including non-elective hospitalizations. Inpatient hospitalizations for patients discharged to another hospital. Inpatient hospitalizations for patients who left against medical advice. Inpatient hospitalizations for patients who expired. Inpatient hospitalizations for patients discharged to a health care facility for hospice care. Inpatient hospitalizations for patients with Comfort measures documented.

**Numerator:** Inpatient hospitalizations for patients prescribed or continuing to take statin medication at hospital discharge

**Numerator Exclusions:** Not Applicable.

**Denominator Exceptions:** Inpatient hospitalizations for patients with a reason for not prescribing statin medication at discharge. Inpatient hospitalizations for patients with a maximum LDL-c result of less than 70 mg/dL <= 30 days prior to arrival or any time during the hospital stay. Inpatient hospitalizations for patients with a statin allergy.

**Supplemental Data Elements:** For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 2.11.2 Data Entry

**Description:** Ischemic stroke patients who are prescribed or continuing to take statin medication at hospital discharge.

- Admit (non-elective), patient over 18, diagnosis = Ischemic Stroke, discharged within measurement period.
- Includes observation services (Value set: 2.16.840.1.113762.1.4.1111.143)
- Denominator exception: Allergy to statin (enter allergy in EHR); Refusal (patient or contraindicated), (enter personal health, refusals, medication).
- Denominator exclusion: LDL-C < 70 within 30 days of hospitalization; Patients with “Comfort Measures” documented (document in IPL/TREG):
• Excludes patients who expire during hospitalization, discharged to another hospital, left AMA, discharged for hospice care (home or facility).

• Numerator: Order statin (2.16.840.1.113762.1.4.1110.19), mark as **Discharge Medication**.

2.12 **CMS 107v7 Stroke Education**

2.12.1 **Detail**

| Description: | Ischemic or hemorrhagic stroke patients or their caregivers who were given educational materials during the hospital stay addressing all of the following: activation of emergency medical system, need for follow-up after discharge, medications prescribed at discharge, risk factors for stroke, and warning signs and symptoms of stroke. |
### Rationale:

There are many examples of how patient education programs for specific chronic conditions have increased healthful behaviors, improved health status, and/or decreased health care costs of their participants. Clinical practice guidelines include recommendations for patient and family education during hospitalization as well as information about resources for social support services. Some clinical trials have shown measurable benefits in patient and caregiver outcomes with the application of education and support strategies. The type of stroke experienced, and the resulting outcomes will play a large role in determining not only the course of treatment but also what education will be required. Patient education should include information about the event (e.g., cause, treatment, and risk factors), the role of various medications or strategies, as well as desirable lifestyle modifications to reduce risk or improve outcomes. Family/caregivers will also need guidance in planning effective and realistic care strategies appropriate to the patient's prognosis and potential for rehabilitation.

### Clinical Recommendation Statement:

Some clinical trials have shown measurable benefits in patient and caregiver outcomes with the application of education and support strategies. Patient education should include information about the event (e.g., cause, treatment, and risk factors), the role of various medications or strategies, as well as desirable lifestyle modifications to reduce risk or improve outcomes. Family/caregivers will also need guidance in planning effective and realistic care strategies appropriate to the patient's prognosis and potential for rehabilitation.

### Guidance

Written information given to the patient is required to address each and every one of the educational components. These components are modeled in the population criteria and data criteria as communication from provider to patient: activation of emergency medical system, follow-up after discharge, medications prescribed at discharge, risk factors and signs and symptoms, and are intended to be specific to stroke. The "Non-elective admissions" value set intends to capture all non-scheduled hospitalizations. This value set is a subset of the "Inpatient encounter" value set, excluding concepts that specifically refer to elective hospital admissions. Non-elective admissions include emergency, urgent and unplanned admissions.

### Initial Population:

Patients age 18 and older discharged from inpatient care (non-elective admissions) with a principal diagnosis of ischemic or hemorrhagic stroke and a length of stay less than or equal to 120 days that ends during the measurement period.

### Denominator:

Ischemic stroke or hemorrhagic stroke patients discharged to home, home care, or court/law enforcement.

### Denominator Exclusions:

Patients with Comfort measures documented.
### Numerator:

Ischemic or hemorrhagic stroke patients with written documentation that they or their caregivers were given educational material addressing all of the following:
- Activation of emergency medical system.
- Follow-up after discharge.
- Medications prescribed at discharge.
- Risk factors for stroke.
- Warning signs and symptoms of stroke.

### Numerator Exclusions:

Not Applicable.

### Denominator:

None.

### Denominator Exceptions:

None.

### Supplemental Data Elements:

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 2.12.2 Data Entry

**Description:**

Ischemic or hemorrhagic stroke patients or their caregivers who were given educational materials during the hospital stay addressing all of the following: activation of emergency medical system, need for follow-up after discharge, medications prescribed at discharge, risk factors for stroke, and warning signs and symptoms of stroke.

Denominator:

- Patients admitted (non-elective), age 18+ (ADT package).
- Principle diagnosis = stroke (hemorrhagic or ischemic) (post discharge PCC entry).
- Discharged during measurement period (LOS <120 days) to home, home care, or court/law enforcement.

Denominator exclusions: Comfort care documented (TREG, Palliative Care).

![Figure 2-27: Palliative Care list](image)

Numerator:

- Written documentation that patient or caregivers given education material including:
  - Warning signs and symptoms of stroke (POV, raising awareness of risk factors).
- FU after discharge (IPL/TREG - follow up arranged).

- Activation of EMS.
  Can enter IPL and search for “Activation of emergency…”

  or:

  Can enter Education code (from Chest Pain or Cardiovascular Disease):

- Meds at discharge (Patient Education – Cerebrovascular Disease – Medications).
- Risk factors for stroke (POV, raising awareness of risk factors).

  Education topic, Cerebrovascular Disease – Disease Process:
CEREBROVASCULAR DISEASE—DISEASE PROCESS

OUTCOME:
The patient/family will understand the cerebrovascular disease.

STANDARD:
1. Discuss the cerebrovascular disease process and types strokes.
2. Explain the risk factors related to the development of cerebrovascular disease.

Figure 2-32: Disease Process

2.13 CMS 108v8 Venous Thromboembolism Prophylaxis

2.13.1 Detail

| Description: | This measure assesses the number of patients who received Venous Thromboembolism (VTE) prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after hospital admission or surgery end date for surgeries that start the day of or the day after hospital admission. |
| **Rationale:** | Hospitalized patients at high-risk for VTE may develop an asymptomatic deep vein thrombosis (DVT) and die from pulmonary embolism (PE) even before the diagnosis is suspected. The majority of fatal events occur as sudden or abrupt death, underscoring the importance of prevention as the most critical action step for reducing death from PE (Geerts et al., 2008). The estimated annual incidence of deep-vein thrombosis (DVT) and pulmonary embolism (PE), known collectively as venous thromboembolism (VTE), is approximately 900,000 (Geerts et al., 2008). Approximately two-thirds of cases of DVT or PE are associated with recent hospitalization. This is consistent with the 2001 report by The Agency for Healthcare Research and Quality (AHRQ). AHRQ indicates that "the appropriate application of effective preventive measures in hospitals has major potential for improving patient safety by reducing the incidence of venous thromboembolism" (Shojania et al., 2001). Despite its proven effectiveness, rates of appropriate thromboprophylaxis remain low in both medical and surgical patients. A recent analysis from the ENDORSE survey, which evaluated prophylaxis rates in 17,084 major surgery patients, found that more than one third of patients at risk for VTE (38%) did not receive prophylaxis and that rates varied by surgery type (Cohen et al., 2008). In a review of evidence-based patient safety practices, the Agency for Healthcare Research and Quality defined thromboprophylaxis against VTE as the "number one patient safety practice" for hospitalized patients (Shojania et al., 2001). Updated "safe practices" published by the National Quality Forum (NQF) recommend routine evaluation of hospitalized patients for risk of VTE and use of appropriate prophylaxis (National Quality Forum, 2006). As noted by the ACCP, a vast number of randomized clinical trials provide irrefutable evidence that thromboprophylaxis reduces VTE events, and there are studies that have also shown that fatal PE is prevented by thromboprophylaxis (Geerts et al., 2008). Some select surgeries have previously been monitored in the Surgical Care Improvement Project; since performance on these surgeries has achieved very high levels, they are not included in this measure. |
| **Clinical Recommendation Statement:** | Failure to recognize and protect patients at risk for VTE increases the chances for acutely ill hospitalized patients at high risk for developing a deep vein thrombosis or dying from a pulmonary emboli. Screening all patients is the only evidence-based practice in reducing incidence of disease. All hospitalized patients should be evaluated for primary VTE prophylaxis and given appropriate prophylaxis when indicated. |
| **Guidance** | When low dose unfractionated heparin is administered for VTE Prophylaxis, the intended administration route for low dose unfractionated heparin is subcutaneous. Reasons for no pharmacological and no mechanical VTE prophylaxis must be explicitly documented by the MD/APN/PA or pharmacist and linked with VTE prophylaxis. Ambulation alone is not a sufficient reason for not administering VTE prophylaxis. In order for ambulation/patient ambulating to be considered as an acceptable reason, there needs to be explicit documentation, e.g., "patient out of bed and ambulating in halls - no VTE prophylaxis needed." |
| **Initial Population:** | Inpatient hospitalizations for patients age 18 and older, discharged from hospital inpatient acute care without a diagnosis of venous thromboembolism (VTE) or obstetrics with a length of stay less than or equal to 120 days that ends during the measurement period. |
| **Denominator:** | Inpatient hospitalizations for all patients in the initial population. |
| **Denominator Exclusions:** | * Inpatient hospitalizations for patients who have a length of stay less than 2 days * Inpatient hospitalizations for patients with Comfort measures documented anytime between the day of arrival and the day after hospital admission * Inpatient hospitalizations for patients with Comfort measures documented by the day after surgery end date for surgeries that start the day of or the day after hospital admission * Inpatient hospitalizations for patients who are direct admits to intensive care unit (ICU), or transferred to ICU the day of or the day after hospital admission with ICU length of stay greater than or equal to one day * Inpatient hospitalizations for patients with a principal diagnosis of mental disorders or stroke * Inpatient hospitalizations for patients with a principal procedure of Surgical Care Improvement Project (SCIP) VTE selected surgeries |
| **Numerator:** | Inpatient hospitalizations for patients who received VTE prophylaxis: - the day of or the day after hospital admission - the day of or the day after surgery end date for surgeries that end the day of or the day after hospital admission Inpatient hospitalizations for patients who have documentation of a reason why no VTE prophylaxis was given: - between the day of arrival and the day after hospital admission - the day of or the day after surgery end date (for surgeries that end the day of or the day after hospital admission) |
| **Numerator Exclusions:** | Not Applicable. |
| **Denominator Exceptions:** | None. |
| **Supplemental Data Elements:** | For every patient evaluated by this measure also identify payer, race, ethnicity, and sex. |
2.13.2 Data Entry

| Description: | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after hospital admission or surgery end date for surgeries that start the day of or the day after hospital admission. |

Denominator:
- Hospital (not ICU) admission, non-elective, patient 18+
- Include observation.
- Discharged within measurement period, LOS <120 days
- No diagnosis on admission of VTE OR Obstetrics

Denominator exclusions:
- LOS <2 days
- “Comfort Measures” documented between arrival and day after admission or surgery
- Direct admit to ICU or transfer to ICU day of admission or next day, LOS ≥1 day
- Principle diagnosis of stroke or mental disorder. See Appendix A for information on using Value Sets (Value Set 2.16.840.1.113883.3.117.1.7.1.212 – Hemorrhagic Stroke) or
- (Value Set 2.16.840.1.113883.3.117.1.7.1.247 – Ischemic Stroke) or
- (Value Set 2.16.840.1.113883.3.464.1003.105.12.1004 – Mental Health Diagnosis)
- Principle diagnosis of selected surgeries:
  - General Surgery
  - Gyn Surgery
  - Hip Fracture Surgery
  - Hip Replacement Surgery
  - Intracranial Neurosurgery
  - Knee Replacement Surgery

![Palliative Care](image)
- Urologic Surgery

Numerator:
- Patients who received VTE prophylaxis within 1 day of admission:
  - Low dose unfractionated heparin
  - Low molecular weight heparin
  - Injectable Factor Xa inhibitor
  - Warfarin
  - Intermittent pneumatic compression devices (IPL/TREG/Anticoag DVT Prevention)
  - Venous foot pumps (TREG/Anticoag DVT Prevention)
  - Graduated compression stockings (TREG/Anticoag DVT Prevention)

Figure 2-34: Treatment/Regimen list

or:
- Oral Factor Xa Inhibitor given
  - AND
  - Prior or present diagnosis of AF or VTE
  - OR Present or Prior Knee or Hip Replacement
- Patients with documentation of a reason no VTE prophylaxis given:
  - Low risk for VTE or anticoagulant administered
    - Low risk assessment score
    - INR >3.0
  - On medication:
    - Direct thrombin inhibitor
    - Unfractionated heparin
    - Glycoprotein IIB/IIA inhibitors
– Patient refusal (Personal history, refusal of treatment by patient)
– Medical reason (Personal history, refusal, Complication of medical treatment)

2.14 CMS 111v8 Median Admit Decision Time to ED Departure Time for Admitted Patients

2.14.1 Detail

| Description: | Median time (in minutes) from admit decision time to time of departure from the ED for ED patients admitted to inpatient status. |
| Stratification: | Report total score and the following strata: Stratification 1 - all patients seen in the ED and admitted as an inpatient who do not have an inpatient encounter principal diagnosis consistent with psychiatric/mental health disorders. Stratification 2 - all patients seen in the ED and admitted as an inpatient who have an inpatient encounter principal diagnosis consistent with psychiatric/mental health disorders. |
| Rate Aggregation: | Calculate the duration in minutes between the Decision to Admit time and the departure time for each ED encounter in the measure population; report the median time for all calculations performed. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the duration the patient was in the ED after the decision to admit, also stated as: the Datetime difference between the ED facility location departure date/time and the Decision to Admit date/time. The calculation requires the median across all ED encounter durations. |
## Rationale:

Reducing the time patients remain in the ED can improve access to treatment and increase quality of care. Reducing this time potentially improves access to care specific to the patient condition and increases the capability to provide additional treatment. In recent times, EDs have experienced significant overcrowding. Although once only a problem in large, urban, teaching hospitals, the phenomenon has spread to other suburban and rural healthcare organizations. According to a 2002 national U.S. survey, more than 90 percent of large hospitals report EDs operating "at" or "over" capacity. Approximately one third of hospitals in the U.S. report increases in ambulance diversion in a given year, whereas up to half report crowded conditions in the ED. In a recent national survey, 40 percent of hospital leaders viewed ED crowding as a symptom of workforce shortages. ED crowding may result in delays in the administration of medication such as antibiotics for pneumonia and has been associated with perceptions of compromised emergency care. For patients with non-ST-segment-elevation myocardial infarction, long ED stays were associated with decreased use of guideline-recommended therapies and a higher risk of recurrent myocardial infarction. Overcrowding and heavy emergency resource demand have led to a number of problems, including ambulance refusals, prolonged patient waiting times, increased suffering for those who wait, rushed and unpleasant treatment environments, and potentially poor patient outcomes. When EDs are overwhelmed, their ability to respond to community emergencies and disasters may be compromised.

## Clinical Recommendation Statement:

The most common cause of ED crowding is the boarding of admitted patients in the ED. Numerous studies have demonstrated the potential for errors, life threatening delays in treatment, and diminished overall quality is enormous.
### Guidance:

This measure specification defines how to determine the duration from a Decision to Admit and the departure from an Emergency Department stay. Reporting requires the median of all admit decision to ED departure durations defined as [Encounter: encounter ED] facility location departure date and time minus [Encounter: encounter ED] ED admit decision date and time. Decision to Admit: Documentation of the decision to admit the patient from the ED that is closest to the inpatient admission. Specification: as admission processes vary at different hospitals, this can use any of the following: 1) admission order (this may be an operational order rather than the hospital admission to inpatient status order), 2) disposition order (must explicitly state to admit), 3) documented bed request, or 4) documented acceptance from admitting physician. This is not the "bed assignment time" or "report called time". Calculate the ED time in minutes for each patient in the measure population; report the median time for all calculations performed. The specification provides elements from the clinical electronic record required to calculate for each ED encounter, i.e., the length of time the patient was in the Emergency Department from the time of decision to admit, also stated as: the Datetime difference for the Emergency Department facility location departure date/time minus the Decision to Admit date/time. The calculation requires the median across all ED encounter durations. For each population, results should be reported without stratification and then with each stratum applied. For this measure, the number of encounters that fall into the Initial Population are reported without stratification, then reported according to the defined stratification. The number of encounters that fall into the Measure Population are reported without stratification, then reported according to the defined stratification. The computed continuous variable defined by the Measure Observation is reported for the Measure Population also, then reported according to the defined stratification.

### Initial Population:

Inpatient Encounters ending during the measurement period with Length of Stay (Discharge Date minus Admission Date) less than or equal to 120 days, and where the decision to admit was made during the preceding emergency department visit at the same physical facility unless the ED and admitting hospital share the same CCN.

### Measure Population:

Initial Population.

### Measure Population Exclusions:

ED encounters with an admission source in "Hospital Setting" (any different facility, even if part of the same hospital system) resulting in an inpatient stay

### Supplemental Data Elements:

For every patient evaluated by this measure, also identify payer, race, ethnicity, and sex.
2.14.2 Data Entry

**Description:** Median time (in minutes) from admit decision time to time of departure from the ED for ED patients admitted to inpatient status.

- Denominator = ER visits resulting in Inpatient admission (w/in one hour of ER departure); admission ends within measurement period (LOS<120 days).
- Exclusions: Patients sent to ER from Hospital.
- Decision to Admit time entered in AMER or ED Dashboard.
- Discharge from ER time entered in AMER or ED Dashboard; Disposition “Admit”.
- System calculates time difference (Admission must occur within 60 minutes of ED Discharge).
- Report all patients, plus:
  - Stratification 1: Patients without a principle diagnosis of psychiatric/mental health disorder. See Appendix A on using Value Sets. (Value Set: 2.16.840.1.113883.3.117.1.7.1.299).
  - Stratification 2: Patients with a principle diagnosis of psychiatric/mental health disorder (Value Set: 2.16.840.1.113883.3.117.1.7.1.299).

2.15 CMS 113v7 Elective Delivery

2.15.1 Detail

**Description:** Patients with elective vaginal deliveries or elective cesarean births at >= 37 and < 39 weeks of gestation completed.
**Rationale:**
For almost 3 decades, the ACOG and the AAP have had in place a standard requiring 39 completed weeks gestation prior to ELECTIVE delivery, either vaginal or operative (ACOG, 1996). A survey conducted in 2007 of almost 20,000 births in HCA hospitals throughout the U.S. carried out in conjunction with the March of Dimes at the request of ACOG revealed that almost 1/3 of all babies delivered in the United States are electively delivered with 5% of all deliveries in the U.S. delivered in a manner violating ACOG/AAP guidelines. Most of these are for convenience, and result in significant short-term neonatal morbidity (neonatal intensive care unit admission rates of 13-21%) (Clark et al., 2009). According to Glantz (2005), compared to spontaneous labor, elective inductions result in more cesarean births and longer maternal length of stay. The American Academy of Family Physicians (2000) also notes that elective induction doubles the cesarean delivery rate. Repeat elective cesarean births before 39 weeks gestation also result in higher rates of adverse respiratory outcomes, mechanical ventilation, sepsis and hypoglycemia for the newborns (Tita et al., 2009).

**Clinical Recommendation Statement:**
Guidelines from the American College of Obstetricians and Gynecologists and the AAP do not support non-medically indicated elective deliveries at <39 weeks gestation. Evidence suggests that early term deliveries result in significant short-term neonatal morbidity. Therefore, it is recommended that elective deliveries should not be performed at <39 weeks gestation unless medically indicated.

**Guidance**
Stillbirth: v2017A of chart-abstracted measure PC-01: Elective Delivery contains a denominator exclusion data element for Stillbirth. The value set for eCQM Denominator Exclusion data element "Assessment, Performed: Conditions Possibly Justifying Elective Delivery" includes SNOMED CT and ICD-10-CM concepts representing Stillbirth and History of Stillbirth. Wherever the gestational age is mentioned with relative timing to delivery, the intent is to capture the estimated gestational age on the day of delivery. It is acceptable to calculate Gestational Age using the American College of Obstetricians and Gynecologists ReVITALize guidelines, which define Gestational Age as calculated using the best obstetrical Estimated Due Date (EDD) based on the formula: Gestational Age = (280-(EDD-Reference Date))/7 where Reference Date is the date on which you are trying to determine gestational age. For PC-01, Reference Date is the Date of Delivery. Note however that the calculation may yield a non-whole number and gestational age should be rounded off to the nearest completed week. For example, an infant born on the 5th day of the 36th week (35 weeks and 5/7 days) is at a gestational age of 35 weeks, not 36 weeks.

**Initial Population:**
Patients age >= 8 years and < 65 admitted to the hospital for inpatient acute care to undergo a delivery procedure and had a length of stay less than or equal to 120 days that ends during the measurement period.
Denominator: Patients delivering newborns with >= 37 and < 39 weeks of gestation completed

Denominator Exclusions: Patients with conditions possibly justifying elective delivery prior to 39 weeks gestation

Numerator: Patients with elective deliveries by either: Medical induction of labor while not in labor prior to the procedure Cesarean birth while not in labor and with no history of a prior uterine surgery

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

2.15.2 Data Entry

Description: Patients with elective vaginal deliveries or elective cesarean births at >= 37 and < 39 weeks of gestation completed.

Denominator= Patients, age >8-<65 admitted for elective vaginal or C/S delivery.

- Baby’s estimated gestational age near term (= or > 37 weeks and < 39 weeks).
  - Personal Health – Birth Measurements.

Figure 2-35: Birth Measurements
Denominator exclusions: Conditions justifying elective delivery before 39 weeks. See Appendix A on using Value Sets. (Value Set: 2.16.840.1.113883.3.117.1.7.1.286).

Numerator:

- Medical induction when not in labor (before start of induction):
  - Medical induction (e.g., oxytocin, dinoprostone) w/in 24 hours of onset of labor.
  - Artificial Rupture of Membranes within 24 hours of onset of labor.
- C/S when not in labor before start of procedure, AND:
  - No history of prior uterine surgery:
    - Classical C/S.
    - Myomectomy.
    - Transabdominal cerclage.
    - Metroplasty.
    - Uterine horn.
  - No history of uterine surgery:
    - Perforation of uterus.
    - Uterine window.
    - Uterine rupture.
• Cornual ectopic pregnancy:

This measure is being retired. Documentation done by Coder and requires following information:

• Date/time labor established.
• Characterize labor.

**MNEMONIC: V DELIVERY VDEL VDELIVERY ALLOWED VISIT RELATED ONLY**

<table>
<thead>
<tr>
<th>Labor Established (Date/Time):</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) 6893006  First stage of labor</td>
</tr>
<tr>
<td>2) 29236002  Labor established</td>
</tr>
<tr>
<td>3) 6383007  Premature labor</td>
</tr>
<tr>
<td>4) 84457005  Spontaneous onset of labor</td>
</tr>
<tr>
<td>5) 366325002  Progress of labor - first stage - finding</td>
</tr>
<tr>
<td>6) 289211007  First stage of labor established</td>
</tr>
</tbody>
</table>

Please Choose a SNOMED CONCEPT ID from the list above: (1-6)

**Figure 2-37: Roll-and-scroll sequence**

• Induced or not.
• If induced, include time and date.
• Method of induction.

**LABOR INDUCED?: Y YES**

<table>
<thead>
<tr>
<th>DATE/TIME LABOR INDUCTION INITIATED:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) 408816000  Artificial rupture of membranes</td>
</tr>
<tr>
<td>2) 408818004  Induction of labor by artificial rupture of membranes</td>
</tr>
<tr>
<td>3) 177129005  Surgical induction of labor</td>
</tr>
<tr>
<td>4) 177139005  Oxytocin induction of labor</td>
</tr>
<tr>
<td>5) 31208007  Medical induction of labor</td>
</tr>
<tr>
<td>6) 308037008  Syntocinon induction of labor</td>
</tr>
<tr>
<td>7) 237002008  Stimulation of labor</td>
</tr>
<tr>
<td>8) 288190009  Induce labor - IV drip</td>
</tr>
<tr>
<td>9) 288189000  Induce labor - IV injection</td>
</tr>
<tr>
<td>10) 177136006  Prostaglandin induction of labor</td>
</tr>
</tbody>
</table>

Please Choose a SNOMED CONCEPT ID from the list above: (1-10)

**Figure 2-38: Roll-and-scroll sequence (cont.)**

• EGA of newborn.
• Sex of newborn.
• Live or stillborn.
Newborn Data: None recorded
Select one of the following:
    A        Add a Newborn Entry
    N        No Change
Which action: N//A Add a Newborn Entry
Enter NEWBORN Delivery Date/Time
Note: There are no EGA measurements on file between a day before and the
delivery date. You will need to manually enter the EGA.
EGA AT DELIVERY:
SEX AT BIRTH:
LIVE/STILLBORN:

Figure 2-39: Roll-and-scroll sequence (cont.)

2.16 CMS 190v8 Intensive Care Unit Venous
Thromboembolism Prophylaxis

2.16.1 Detail

| Description: | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after the initial admission (or transfer) to the ICU or surgery end date for surgeries that start the day of or the day after ICU admission (or transfer). |
| **Rationale:** | Approximately two-thirds of cases of Deep Vein Thrombosis (DVT) or Pulmonary Emboli (PE) are associated with recent hospitalization. This is consistent with the 2001 report by Agency for Healthcare Research and Quality (Shojania et al., 2001). AHRQ reports that "the appropriate application of effective preventive measures in hospitals has major potential for improving patient safety, by reducing the incidence of VTE." Almost all hospitalized patients have at least one risk factor for Venous Thromboembolism (VTE), and approximately 40% have three or more risk factors. Without thromboprophylaxis, the incidence of objectively confirmed, hospital-acquired DVT is approximately 10% to 40% among medical or general surgical patients and 40% to 60% following major orthopedic surgery (Geerts et al., 2008). Commonly, criteria for admission to the Intensive Care Unit (ICU) itself, puts patients at an increased risk for developing VTE, and subsequent increased risk of morbidity from PE. Some risk factors are related to the acute illness present that allowed for the admission to the ICU unit, and some risk factors may be acquired during the ICU admission due to subsequent medical treatments, for example limitations of mobility, presence of central venous lines or mechanical ventilation and subsequent pharmacological paralysis. Reports of DVT in the population of ICU patients vary in relation to the acuity of the illness in this population. DVT in ICU patients diagnosed with routine venography or Doppler ultrasound found ranges between 10% to 100%. Five studies prospectively screened patients who were not receiving thromboprophylaxis during their ICU stays. The rates of DVT using Fibrinogen Uptake Test, Doppler Ultrasound or venography ranged from 13 to 31% (Geerts et al., 2008). It is essential for all ICUs to assess each patient upon admission to the ICU unit, a change in level of status, for the need for VTE prophylaxis due to the above increased development of risk factors (Geerts et al., 2004). Some select surgeries have previously been monitored in the Surgical Care Improvement Project; since performance on these surgeries has achieved very high levels, they are not included in this measure. |
| **Clinical Recommendation Statement:** | Failure to recognize and protect patients at risk for VTE increases the chances for critically ill hospitalized patients for developing a deep vein thrombosis or dying from a pulmonary emboli. Screening all patients is the only evidence based practice in reducing incidence of disease. All intensive care unit (ICU) patients should be evaluated for primary VTE prophylaxis, and given appropriate prophylaxis when indicated. |
**Guidance:**
The definition of an ICU for the purpose of the measures noted above is that used by the CDC in the NHSN Patient Safety Project. An intensive care unit can be defined as a nursing care area that provides intensive observation, diagnosis, and therapeutic procedures for adults and/or children who are critically ill. An ICU excludes nursing areas that provide step-down, intermediate care or telemetry only and specialty care areas.

Reasons for no pharmacological and no mechanical VTE prophylaxis must be explicitly documented by the MD/APN/PA or pharmacist and linked with VTE prophylaxis. Ambulation alone is not a sufficient reason for not administering VTE prophylaxis. In order for ambulation/patient ambulating to be considered as an acceptable reason, there needs to be explicit documentation, e.g., "patient out of bed and ambulating in halls - no VTE prophylaxis needed."

**Initial Population:**
Inpatient hospitalizations for patients age 18 and older, discharged from hospital inpatient acute care without a diagnosis of venous thromboembolism (VTE) or obstetrics with a length of stay less than or equal to 120 days that ends during the measurement period

**Denominator:**
Inpatient hospitalizations for patients directly admitted or transferred to ICU during the hospitalization

**Denominator Exclusions:**
* Inpatient hospitalizations for patients who have a hospital length of stay (LOS) less than 2 days
* Inpatient hospitalizations for patients with Comfort measures documented anytime between the day of arrival and the day after ICU admission or transfer
* Inpatient hospitalizations for patients with Comfort measures documented by the day after surgery end date for surgeries that end the day of or the day after hospital admission
* Inpatient hospitalizations for patients with a principal procedure of surgical care improvement Project (SCIP) VTE selected surgeries that end the day of or the day after ICU admission or transfer

**Numerator:**
Inpatient hospitalizations for patients who received VTE prophylaxis: - the day of or the day after ICU admission (or transfer) - the day of or the day after surgery end date for surgeries that end the day of or the day after ICU admission (or transfer)

**Numerator Exclusions:**
Not Applicable

**Denominator Exceptions:**
Inpatient hospitalizations for patients with ICU LOS less than one day

**Supplemental Data Elements:**
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.
2.16.2 Data Entry

| Description: | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after the initial admission (or transfer) to the ICU or surgery end date for surgeries that start the day of or the day after ICU admission (or transfer). |

Inpatient encounter, patient > 18, discharged without a diagnosis of VTE or Obstetrics, LOS < 120 days. Include observation.

Denominator: Patients admitted to or transferred to ICU during admission.

Denominator Exclusions:
- LOS < 2 days.
- Comfort measures documented from start of hospitalization to day after first ICU stay.
- Comfort measures documented day after general anesthesia done after ICU admission.

Figure 2-40: Palliative Care list

- Primary Diagnosis selected surgeries done by the day after ICU admission:
  - General Surgery.
  - Gyn Surgery.
  - Hip Fracture Surgery.
  - Hip Replacement Surgery.
  - Intracranial Neurosurgery.
  - Knee Replacement Surgery.
  - Urologic Surgery.

Numerator:
- Received VTE prophylaxis day of or day after ICU admission or surgical procedure.
  - Low dose unfractionated heparin.
  - Low molecular weight heparin.
  - Injectable Factor Xa inhibitor.
  - Warfarin.
- Intermittent pneumatic compression devices (IPL/TREG/Anticoag DVT Prevention).
- Venous foot pumps (IPL/TREG/Anticoag DVT Prevention).
- Graduated compression stockings (IPL/TREG/Anticoag DVT Prevention).

Figure 2-41: Treatment/Regimen list

or:

- Oral Factor Xa Inhibitor given.

and:

- Prior or present diagnosis of AF or VTE.
- OR Present or Prior Knee or Hip Replacement surgery.

- Documented reason for no VTE prophylaxis given.
- Medical reason (document refusal file, complication of medical care).
- Low risk for VTE.
  - Low risk assessment score.
  - INR >3.0.
  - On medication:
    - Direct thrombin inhibitor.
    - Unfractionated heparin.
    - Glycoprotein IIB/IIA inhibitors.

Patient refusal (document as refusal of treatment by patient in refusal file, Personal history).

Denominator Exceptions: ICU stay <1 day
3.0 Provider Measures for 2020 Reporting Period

3.1 CMS 2v9 Preventive Care and Screening for Depression

3.1.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients aged 12 years and older screened for depression on the date of the encounter or 14 days prior to the date of the encounter using an age appropriate standardized depression screening tool AND if positive, a follow-up plan is documented on the date of the eligible encounter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Depression is a serious medical illness associated with higher rates of chronic disease, increased health care utilization, and impaired functioning (Pratt &amp; Brody, 2014). 2016 U.S. survey data indicate that 12.8 percent of adolescents (2.2 million adolescents) had a major depressive episode (MDE) in the past year, with nine percent of adolescents (2.2 million adolescents) having one MDE with severe impairment; 6.7 percent of adults aged 18 or older (16.2 million adults) had at least one MDE in the past year, with 4.3 percent of adults (10.3 million adults) having one MDE with severe impairment in the past year (Substance Abuse and Mental Health Services Administration, 2017). Data indicate that severity of depressive symptoms factor into having difficulty with work, home, or social activities. For example, as the severity of depressive symptoms increased, rates of having difficulty with work, home, or social activities related to depressive symptoms increased. For those twelve and older with mild depressive symptoms, 45.7% reported difficulty with activities and those with severe depressive symptoms, 88.0% reported difficulty (Pratt &amp; Brody, 2014). Children and teens with major depressive disorder (MDD) has been found to have difficulty carrying out their daily activities, relating to others, and growing up healthy with an increased risk of suicide (Siu &amp; the U.S. Preventive Services Task Force [USPSTF], 2016). Additionally, perinatal depression (considered here as depression arising in the period from conception to the end of the first postnatal year) affects up to 15% of women. Depression and other mood disorders, such as bipolar disorder and anxiety disorders, especially during the perinatal period, can have devastating effects on women, infants, and families (Molenaar et al., 2018). Maternal suicide rates rise over hemorrhage and hypertensive disorders as a cause of maternal mortality (American College of Obstetricians and Gynecologists, 2015). Negative outcomes associated with depression make it crucial to screen in order to identify and treat depression in its early stages. While Primary Care Providers (PCPs) serve as the first line of defense in the detection of depression, studies show that PCPs fail to recognize up to 50% of depressed patients (Borner, 2010, p. 948).</td>
</tr>
</tbody>
</table>
Rationale: (cont)

Coyle et al. (2003) suggested that the picture is more grim for adolescents, and that more than 70% of children and adolescents suffering from serious mood disorders go unrecognized or inadequately treated" (Borner et al., 2010, p. 948 ). In nationally representative U.S. surveys, about eight percent of adolescents reported having major depression in the past year. Only 36% to 44% of children and adolescents with depression receive treatment, suggesting that the majority of depressed youth are undiagnosed and untreated" (Siu on behalf of USPSTF, 2016, p. 360 & 364). Evidence supports that screening for depression in pregnant and postpartum women is of moderate net benefit and treatment options for positive depression screening should be available for patients twelve and older including pregnant and postpartum women. If preventing negative patient outcomes is not enough, the substantial economic burden of depression for individuals and society alike makes a case for screening for depression on a regular basis. Depression imposes economic burden through direct and indirect costs: "In the United States, an estimated $22.8 billion was spent on depression treatment in 2009, and lost productivity cost an additional estimated $23 billion in 2011" (Siu & USPSTF, 2016, p. 383-384). This measure seeks to align with clinical guideline recommendations as well as the Healthy People 2020 recommendation for routine screening for mental health problems as a part of primary care for both children and adults (U.S. Department of Health and Human Services, 2014) and makes an important contribution to the quality domain of community and population health.

Clinical Recommendation Statement:

Adolescent Recommendation (12-18 years): "The USPSTF recommends screening for MDD in adolescents aged 12 to 18 years. Screening should be implemented with adequate systems in place to ensure accurate diagnosis, effective treatment, and appropriate follow-up (B recommendation)" (Siu on behalf of USPSTF, 2016, p. 360). "Clinicians and health care systems should try to consistently screen adolescents, ages 12-18, for major depressive disorder, but only when systems are in place to ensure accurate diagnosis, careful selection of treatment, and close follow-up" (Wilkinson et al., 2013, p. 16). Adult Recommendation (18 years and older): "The USPSTF recommends screening for depression in the general adult population, including pregnant and postpartum women. Screening should be implemented with adequate systems in place to ensure accurate diagnosis, effective treatment, and appropriate follow-up (B recommendation)" (Siu & USPSTF, 2016, p. 380). The Institute for Clinical Systems Improvement (ICSI) health care guideline, Adult Depression in Primary Care, provides the following recommendations: 1. "Clinicians should routinely screen all adults for depression using a standardized instrument." 2. "Clinicians should establish and maintain follow-up with patients." 3. "Clinicians should screen and monitor depression in pregnant and post-partum women." (Trangle et al., 2016, p. 8-10).
<table>
<thead>
<tr>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Screening:</strong> Completion of a clinical or diagnostic tool used to identify people at risk of developing or having a certain disease or condition, even in the absence of symptoms.</td>
</tr>
<tr>
<td><strong>Standardized Depression Screening Tool</strong> - A normalized and validated depression screening tool developed for the patient population in which it is being utilized. Examples of depression screening tools include but are not limited to:</td>
</tr>
<tr>
<td>Adolescent Screening Tools (12-17 years).</td>
</tr>
<tr>
<td>Patient Health Questionnaire for Adolescents (PHQ-A).</td>
</tr>
<tr>
<td>Beck Depression Inventory-Primary Care Version (BDI-P).</td>
</tr>
<tr>
<td>Mood Feeling Questionnaire (MFQ).</td>
</tr>
<tr>
<td>Center for Epidemiologic Studies Depression Scale (CES-D).</td>
</tr>
<tr>
<td>Patient Health Questionnaire (PHQ-9).</td>
</tr>
<tr>
<td>Pediatric Symptom Checklist (PSC-17)</td>
</tr>
<tr>
<td>PRIME MD-PHQ2.</td>
</tr>
<tr>
<td>Adult Screening Tools (18 years and older).</td>
</tr>
<tr>
<td>Patient Health Questionnaire (PHQ9).</td>
</tr>
<tr>
<td>Beck Depression Inventory (BDI or BDI-II).</td>
</tr>
<tr>
<td>Center for Epidemiologic Studies Depression Scale (CES-D).</td>
</tr>
<tr>
<td>Depression Scale (DEPS).</td>
</tr>
<tr>
<td>Duke Anxiety-Depression Scale (DADS).</td>
</tr>
<tr>
<td>Geriatric Depression Scale (GDS).</td>
</tr>
<tr>
<td>Cornell Scale for Depression in Dementia (CSDS).</td>
</tr>
<tr>
<td>PRIME MD-PHQ2.</td>
</tr>
<tr>
<td>Hamilton Rating Scale for Depression (HAM-D).</td>
</tr>
<tr>
<td>Quick Inventory of Depressive Symptomatology Self-Report (QID-SR).</td>
</tr>
<tr>
<td>Computerized Adaptive Testing Depression Inventory (CAT-DI).</td>
</tr>
<tr>
<td>Computerized Adaptive Diagnostic Screener (CAD-MDD).</td>
</tr>
<tr>
<td>Perinatal Screening Tools.</td>
</tr>
<tr>
<td>Edinburgh Postnatal Depression Scale.</td>
</tr>
<tr>
<td>Postpartum Depression Screening Scale.</td>
</tr>
<tr>
<td>Patient Health Questionnaire 9 (PHQ-9).</td>
</tr>
<tr>
<td>Beck Depression Inventory.</td>
</tr>
<tr>
<td>Beck Depression Inventory-II.</td>
</tr>
<tr>
<td>Center for Epidemiologic Studies Depression Scale.</td>
</tr>
<tr>
<td>Zung Self-rating Depression Scale.</td>
</tr>
</tbody>
</table>
| **Definition (cont):** | Follow-Up Plan:  
Documented follow-up for a positive depression screening must include one or more of the following:  
Additional evaluation or assessment for depression.  
Suicide Risk Assessment.  
Referral to a practitioner who is qualified to diagnose and treat depression.  
Pharmacological interventions.  
Other interventions or follow-up for the diagnosis or treatment of depression. |
**Guidance:**

A depression screen is completed on the date of the encounter or up to 14 days prior to the date of the encounter using an age appropriate standardized depression screening tool AND if positive, either additional evaluation for depression, suicide risk assessment, referral to a practitioner who is qualified to diagnose and treat depression, pharmacological interventions, or other interventions or follow-up for the diagnosis or treatment of depression is documented on the date of the eligible encounter. Depression screening is required once per measurement period, not at all encounters; this is patient based and not an encounter based measure. Screening Tools: * The name of the age appropriate standardized depression screening tool utilized must be documented in the medical record * The depression screening must be reviewed and addressed in the office of the provider, filing the code, on the date of the encounter. Positive pre-screening results indicating a patient is at high risk for self-harm should receive more urgent intervention as determined by the provider practice. * The screening should occur during a qualified encounter or up to 14 days prior to the date of the qualifying encounter. * Standardized depression screening tools should be normalized and validated for the age appropriate patient population in which they are used. Follow-Up Plan: * The follow-up plan must be related to a positive depression screening, example: "Patient referred for psychiatric evaluation due to positive depression screening." Examples of a follow-up plan include but are not limited to: * Additional evaluation or assessment for depression such as psychiatric interview, psychiatric evaluation, or assessment for bipolar disorder * Completion of any Suicide Risk Assessment such as Beck Depression Inventory or Beck Hopelessness Scale * Referral to a practitioner or program for further evaluation for depression, for example, referral to a psychiatrist, psychologist, social worker, mental health counselor, or other mental health service such as family or group therapy, support group, depression management program, or other service for treatment of depression * Other interventions designed to treat depression such as psychotherapy, pharmacological interventions, or additional treatment options * Pharmacologic treatment for depression is often indicated during pregnancy and/or lactation. Review and discussion of the risks of untreated versus treated depression is advised. Consideration of each patient's prior disease and treatment history, along with the risk profiles for individual pharmacologic agents, is important when selecting pharmacologic therapy with the greatest likelihood of treatment effect.

<table>
<thead>
<tr>
<th><strong>Initial Population:</strong></th>
<th>All patients aged 12 years and older at the beginning of the measurement period with at least one eligible encounter during the measurement period.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Denominator:</strong></td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td><strong>Denominator Exclusions:</strong></td>
<td>Patients with an active diagnosis for depression or a diagnosis of bipolar disorder.</td>
</tr>
</tbody>
</table>
Numerator: Patients screened for depression on the date of the encounter or up to 14 days prior to the date of the encounter using an age appropriate standardized tool AND if positive, a follow-up plan is documented on the date of the eligible encounter.

Numerator Exclusions: Not Applicable.

Denominator Exceptions: Patient Reason(s) Patient refuses to participate OR Medical Reason(s) Patient is in an urgent or emergent situation where time is of the essence and to delay treatment would jeopardize the patient's health status OR Situations where the patient's cognitive capacity, functional capacity or motivation to improve may impact the accuracy of results of standardized depression assessment tools. For example: certain court appointed cases or cases of delirium.

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

3.1.2 Data Entry

Description: Percentage of patients aged 12 years and older screened for depression on the date of the encounter using an age appropriate standardized depression screening tool AND if positive, a follow-up plan is documented on the date of the positive screen.

Enter encounter, patient age 12+

(Multiple OPD encounters accepted, (Value Set: 2.16.840.1.113883.3.600.1916).

See Appendix A on Using Value Sets.

Exclusion if IPL has active diagnosis of “Depression” (Value Set: 2.16.840.1.113883.3.600.145) or “Bipolar Diagnosis” (Value Set: 2.16.840.1.113883.3.600.450) entered before encounter date.

Exceptions:

- Patient refusals (document in Personal Health component, wellness tab, exams).
- Medical Reasons: Patient is in an urgent or emergent situation where time is of the essence and to delay treatment would jeopardize the patient's health status.
- Other: Situations where the patient's functional capacity or motivation to improve may impact the accuracy of results of standardized depression assessment tools. For example: certain court appointed cases or cases of delirium.

Screen patient for depression (PHQ 2), enter result in Vital Signs,
Or enter “positive” or “negative” in Exams.

- If depression screen is Negative (PHQ ≤2 score in vital signs, or “negative” screen in exams), this will be included in numerator.
- If depression screen is Positive (PHQ ≥3, or “positive” exam), then must perform intervention on same day as encounter in order to be included in numerator. Multiple intervention options available (need only one):
  - Intervention: PHQ 9 or PHQt with any results (Vital Signs).
  - Referral or consult to mental health worker.

- Same day visit with mental health worker, POV mental health care management
- Suicide risk assessment. Value set 2.16.840.1.113883.3.600.559
Order medication for depression. See Appendix A on Using Value Sets.
- (Value Set: 2.16.840.1.113883.3.600.469)-adolescent, or
- (Value Set: 2.16.840.1.113883.3.600.470) -adult

3.2 CMS 50v8 Closing the Referral Loop: Receipt of Specialist Report

3.2.1 Detail
Percentage of patients with referrals, regardless of age, for which the referring provider receives a report from the provider to whom the patient was referred.

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients with referrals, regardless of age, for which the referring provider receives a report from the provider to whom the patient was referred.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Problems in the outpatient referral and consultation process have been documented, including lack of timeliness of information and inadequate provision of information between the specialist and the requesting physician (Gandhi, 2000; Forrest, 2000; Stille, 2005). In a study of physician satisfaction with the outpatient referral process, Gandhi et al. (2000) found that 68% of specialists reported receiving no information from the primary care provider prior to referral visits, and 25% of primary care providers had still not received any information from specialists 4 weeks after referral visits. In another study of 963 referrals (Forrest, 2000), pediatricians scheduled appointments with specialists for only 39% and sent patient information to the specialists in only 51% of the time.</td>
</tr>
</tbody>
</table>
| Rationale (cont): | In a 2006 report to Congress, MedPAC found that care coordination programs improved quality of care for patients, reduced hospitalizations, and improved adherence to evidence-based care guidelines, especially among patients with diabetes and CHD. Associations with cost-savings were less clear; this was attributed to how well the intervention group was chosen and defined, as well as the intervention put in place. Additionally, cost-savings were usually calculated in the short-term, while some argue that the greatest cost-savings accrue over time (MedPAC, 2006).

Improved mechanisms for information exchange could facilitate communication between providers, whether for time-limited referrals or consultations, on-going co-management, or during care transitions. For example, a study by Branger et al. (1999) found that an electronic communication network that linked the computer-based patient records of physicians who had shared care of patients with diabetes significantly increased frequency of communications between physicians and availability of important clinical data. There was a 3-fold increase in the likelihood that the specialist provided written communication of results if the primary care physician scheduled appointments and sent patient information to the specialist (Forrest, 2000).

Care coordination is a focal point in the current health care reform and our nation’s ambulatory health information technology (HIT) framework. The National Priorities Partnership recently highlighted care coordination as one of the most critical areas for development of quality measurement and improvement (NPP, 2008). |
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Definition:</td>
<td>Referral: A request from one physician or other eligible provider to another practitioner for evaluation, treatment, or co-management of a patient's condition. This term encompasses referral and consultation as defined by Centers for Medicare and Medicaid Services.</td>
</tr>
</tbody>
</table>
**Guidance:**
The provider who refers the patient to another provider is the provider who should be held accountable for the performance of this measure. The provider to whom the patient was referred should be the same provider that sends the report. If there are multiple referrals for a patient during the measurement period, use the first referral. The consultant report that will fulfill the referral should be completed after the referral and should be related to the referral for which it is attributed. If there are multiple consultant reports received by the referring provider which pertain to a particular referral, use the first consultant report to satisfy the measure. Eligible professionals or eligible clinicians reporting on this measure should note that all data for the reporting year is to be submitted by the deadline established by CMS. Therefore, eligible professionals or eligible clinicians who see patients towards the end of the reporting period (i.e., December in particular), should communicate the consultant report as soon as possible in order for those patients to be counted in the measure numerator. Communicating the report as soon as possible will ensure the data is included in the submission to CMS.

<table>
<thead>
<tr>
<th>Initial Population:</th>
<th>Number of patients, regardless of age, who were referred by one provider to another provider, and who had a visit during the measurement period.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator:</td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td>Denominator Exclusions:</td>
<td>None.</td>
</tr>
<tr>
<td>Numerator:</td>
<td>Number of patients with a referral, for which the referring provider received a report from the provider to whom the patient was referred.</td>
</tr>
<tr>
<td>Numerator Exclusions:</td>
<td>Not Applicable.</td>
</tr>
<tr>
<td>Denominator Exceptions:</td>
<td>None.</td>
</tr>
<tr>
<td>Supplemental Data Elements:</td>
<td>For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.</td>
</tr>
</tbody>
</table>

### 3.2.2 Data Entry

| Description: | Percentage of patients with referrals, regardless of age, for which the referring provider receives a report from the provider to whom the patient was referred. |

Patient visit during measurement period (any age), face to face encounter.

Multiple OPD encounters accepted.

Ophthalmology (Value Set 2.16.840.1.113883.3.526.3.1285).
Preventive care (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1022, …1023, …1024, …1025)

Office visits (Value Set 2.16.840.1.113883.3.464.1003.101.12.1001)

See Appendix A on using Value Sets.

Referral made anytime during measurement period (RCIS).

![Figure 3-5: Add Referral pane](image)

Patient visits referred provider, consult report completed.

Report received, attached to note with provider identified as signer, signed as received by provider (RCIS).

![Figure 3-6: Clinical Consultation pane](image)

Close out referral in RPMS, Referral Management, Close Out Referral, Current Fiscal Year.

### 3.3 CMS 69v8 Preventive Care and Screening: BMI Screening and Follow-Up Plan

#### 3.3.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage of patients aged 18 years and older with a BMI documented during the current encounter or during the previous twelve months AND with a BMI outside of normal parameters, a follow-up plan is documented during the encounter or during the previous twelve months of the current encounter. Normal Parameters: Age 18 years and older BMI =&gt; 18.5 and &lt; 25 kg/m2.</td>
</tr>
</tbody>
</table>
### Rationale:

BMI Above Normal Parameters Obesity is a chronic, multifactorial disease with complex psychological, environmental (social and cultural), genetic, physiologic, metabolic and behavioral causes and consequences. The prevalence of overweight and obese people is increasing worldwide at an alarming rate in both developing and developed countries. Environmental and behavioral changes brought about by economic development, modernization and urbanization have been linked to the rise in global obesity. The health consequences are becoming apparent (Fitch, 2013). Hales et al. (2017), report that the prevalence of obesity among adults and youth in the United States was 39.8% and 18.5% respectively, from 2015-2016. They note that obesity prevalence was higher among adults in the 40-59 age bracket than those in the 20-39 age bracket, for both men and women. Hales et al. (2017) also disaggregated the data according to ethnicity and noted that obesity prevalence was higher among non-Hispanic black and Hispanic adults and youth when compared with other races ethnicities. While obesity prevalence was lower among non-Hispanic Asian men and women, obesity prevalence among men, was comparable between non-Hispanic black and non-Hispanic white men. Obesity prevalence was higher among Hispanic men compared with non-Hispanic black men. While the prevalence among non-Hispanic black and Hispanic women was comparable, the prevalence for both groups was higher than that of non-Hispanic white women. Most notably, Hales et al. (2017), report that the prevalence of obesity in the United States remains higher than the Healthy People 2020 goals of 14.5% among youth and 30.5% among adults. More than a third of U.S. adults have a body mass index \( [\text{BMI}] \geq 30 \text{ kg/m}^2 \); substantially at increased risk for diabetes and cardiovascular disease (CVD) (Flegal et al., 2012; Ogden et al., 2014). Behavioral weight management treatment has been identified as an effective first-line treatment for obesity with an average initial weight loss of eight to ten percent. This percentage weight loss is associated with a significant risk reduction for diabetes and CVD (Wadden, Butryn & Wilson, 2007). Despite the availability of effective interventions, two-thirds of obese U.S. patients were not offered or referred to weight management treatment during their primary care visit between 2005 and 2006, (Ma et al., 2009). In addition, the rate of weight management counseling in primary care significantly decreased by ten percent (40% to 30%) between 1995-1996 and 2007-2008 (Kraschnewski et al., 2013). This suggests that the availability of evidence based clinical guidelines since 2008 obesity management in primary care remains suboptimal (Fitzpatrick & Stevens, 2017). BMI continues to be a common and reasonably reliable measurement to identify overweight and obese adults who may be at an increased risk for future morbidity. Although good quality evidence supports obtaining a BMI, it is important to recognize it is not a perfect measurement. BMI is not a direct measure of adiposity and as a consequence it can over or underestimate adiposity.)
Rationale (cont)

BMI is a derived value that correlates well with total body fat and markers of secondary complications, e.g., hypertension and dyslipidemia (Barlow & the Expert Committee, 2007). In contrast with waist circumference, BMI and its associated disease and mortality risk appear to vary among ethnic subgroups. Female African American populations appear to have the lowest mortality risk at a BMI of 26.2-28.5 kg/m² and 27.1-30.2 kg/m² for women and men, respectively. In contrast, Asian populations may experience lowest mortality rates starting at a BMI of 23 to 24 kg/m². The correlation between BMI and diabetes risk also varies by ethnicity (LeBlanc et al., 2011, pp. 2-3). Screening for BMI and follow-up therefore is critical to closing this gap and contributes to quality goals of population health and cost reduction. However, due to concerns for other underlying conditions (such as bone health) or nutrition related deficiencies providers are cautioned to use clinical judgment and take these into account when considering weight management programs for overweight patients, especially the elderly (National Heart, Lung, and Blood Institute [NHLBI] Obesity Education Initiative, 1998, p. 91). It is important to enhance beneficiary access to all existing providers of Intensive Behavioral Therapy for obesity (IBT) which would result in decreased healthcare costs and lower obesity rates. Dietary counseling performed by a Registered Dietitian Nutritionist (RDN) is more effective than by a primary care clinician. IBT provided by RDNs for 6-12 months shows significant mean weight loss of up to 10% of body weight, maintained over one year's time (Raynor & Champagne, 2016).

BMI below Normal Parameters On the other end of the body weight spectrum is underweight (BMI <18.5 kg/m²), which is equally detrimental to population health. When compared to normal weight individuals (BMI 18.5-25 kg/m²), which is equally detrimental to population health. When compared to normal weight individuals (BMI 18.5-25 kg/m²), underweight individuals have significantly higher death rates with a Hazard Ratio of 2.27 and 95% confidence intervals (CI) = 1.78, 2.90 (Borrell & Lalitha, 2014). Poor nutrition or underlying health conditions can result in underweight (Fryar & Ogden, 2012). The National Health and Nutrition Examination Survey (NHANES) results from the 2007-2010 indicate that women are more likely to be underweight than men. Therefore, patients should be equally screened for underweight and followed up with nutritional counselling to reduce mortality and morbidity associated with underweight.
Clinical Recommendation Statement:

All adults should be screened annually using a BMI measurement. BMI measurements >=25kg/m2 should be used to initiate further evaluation of overweight or obesity after taking into account age, gender, ethnicity, fluid status, and muscularity; therefore, clinical evaluation and judgment must be used when BMI is employed as the anthropometric indicator of excess adiposity, particularly in athletes and those with sarcopenia (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, pp. 12-13) (Grade A). Overweight and Underweight Categories:

Underweight <18.5; Normal weight 18.5-24.9; Overweight 25-29.9; Obese class I 30-34.9; Obese class II 35-39.9; Obese class III >=40 (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, p. 15). When evaluating patients for adiposity related disease risk, waist circumference should be measured in all patients with BMI <35 kg/m2 (Garvey, et. al., 2016 AACE/ACE Guidelines, 2016, p. 13) (Grade A). BMI cutoff point value of >= 23 kg/m2 should be used in the screening and confirmation of excess adiposity in Asian adults (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, p. 13) (Grade B). In the United States the waist circumference cutoff points that can be used to indicate increased risk are >=102 cm (>40 inches) for men and >=88 cm (>35 inches) for women (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, p. 13) (Grade A). Lifestyle/Behavioral Therapy for Overweight and Obesity should include behavioral interventions that enhance adherence to prescriptions for a reduced-calorie meal plan and increased physical activity (behavioral interventions can include: self-monitoring of weight, food intake, and physical activity; clear and reasonable goal-setting; education pertaining to obesity, nutrition, and physical activity; face-to-face and group meetings; stimulus control; systematic approaches for problem solving; stress reduction; cognitive restructuring [i.e., cognitive behavioral therapy], motivational interviewing; behavioral contracting; psychological counseling; and mobilization of social support structures) (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, p. 22) (Grade A). Behavioral lifestyle intervention should be tailored to a patient's ethnic, cultural, socioeconomic, and educational background (Garvey, et al., 2016 AACE/ACE Guidelines, 2016, p. 22) (Grade B). USPSTF Clinical Guideline (Grade B Recommendation) The USPSTF recommends that clinicians offer or refer adults with a body mass index (BMI) of 30 kg/m2 or higher to intensive, multicomponent behavioral interventions.

Interventions: - Effective intensive behavioral interventions were designed to help participants achieve or maintain a >= 5% weight loss through a combination of dietary changes and increased physical activity - Most interventions lasted for 1 to 2 years, and the majority had >= 12 sessions in the first year - Most behavioral interventions focused on problem solving to identify barriers, self-monitoring of weight, peer support, and relapse prevention - Interventions also provided tools to support weight loss or weight loss maintenance (e.g., pedometers, food scales, or exercise videos) (USPSTF, 2018a)
The USPSTF recommends screening for abnormal blood glucose levels as part of cardiovascular risk assessment in adults aged 40 to 70 years who are overweight or have obesity. Patients with certain risk factors (family history of diabetes, personal history of gestational diabetes or polycystic ovarian syndrome, or being a member of certain racial/ethnic groups [African American, American Indian or Alaskan Native, Asian American, Hispanic or Latino, or Native Hawaiian or Pacific Islander]) may also be at increased risk of diabetes at a younger age or at a lower BMI and should be considered for screening (USPSTF, 2018b). Nutritional safety for the elderly should be considered when recommending weight reduction. "A clinical decision to forego obesity treatment in older adults should be guided by an evaluation of the potential benefits of weight reduction for day-to-day functioning and reduction of the risk of future cardiovascular events, as well as the patient's motivation for weight reduction. Care must be taken to ensure that any weight reduction program minimizes the likelihood of adverse effects on bone health or other aspects of nutritional status" (NHLBI Obesity Education Initiative, 1998, p. 91) (Evidence Category D). In addition, weight reduction prescriptions in older persons should be accompanied by proper nutritional counseling and regular body weight monitoring (NHLBI Obesity Education Initiative, 1998, p. 91). The possibility that a standard approach to weight loss will work differently in diverse patient populations must be considered when setting expectations about treatment outcomes (NHLBI Obesity Education Initiative, 1998, p. 97) (Evidence Category B).
**Overweight and Underweight Categories:**

Underweight <18.5; Normal weight 18.5-24.9; Overweight 25-29.9; Obese class I 30-34.9; Obese class II 35-39.9; Obese class III >=40 (AACE/ACE Guidelines, 2016. p. 15).

When evaluating patients for adiposity related disease risk, waist circumference should be measured in all patients with BMI <35 kg/m² (AACE/ACE Guidelines, 2016. p. 13) (Grade A).

In the United States the waist circumference cutoff points that can be used to indicate increased risk are >=102 cm (>40 inches) for men and >=88 cm (more than >35 inches) for women (AACE/ACE Guidelines, 2016. p. 13) (Grade A).

Lifestyle/Behavioral Therapy for Overweight and Obesity should include behavioral interventions that enhance adherence to prescriptions for a reduced-calorie meal plan and increased physical activity (behavioral interventions can include: self-monitoring of weight, food intake, and physical activity; clear and reasonable goal-setting; education pertaining to obesity, nutrition, and physical activity; face-to-face and group meetings; stimulus control; systematic approaches for problem solving; stress reduction; cognitive restructuring [i.e., cognitive behavioral therapy], motivational interviewing; behavioral contracting; psychological counseling; and mobilization of social support structures) (AACE/ACE Guidelines, 2016. p. 22) (Grade A).

Behavioral lifestyle intervention should be tailored to a patient's ethnic, cultural, socioeconomic, and educational background (AACE/ACE Guidelines, 2016. p. 22) (Grade B).

USPSTF Clinical Guideline (Grade B Recommendation)

Individuals with a BMI of 30 kg/m² or higher should be offered or referred to intensive, multicomponent behavioral interventions that include the following components:

- Behavioral management activities, such as setting weight-loss goals.
- Improving diet or nutrition and increasing physical activity.
- Addressing barriers to change.
- Self-monitoring.
- Strategizing how to maintain lifestyle changes.

Nutritional safety for the elderly should be considered when recommending weight reduction. "A clinical decision to forego obesity treatment in older adults should be guided by an evaluation of the potential benefits of weight reduction for day-to-day functioning and reduction of the risk of future cardiovascular events, as well as the patient's motivation for weight reduction. Care must be taken to ensure that any weight reduction program minimizes the likelihood of adverse effects on bone health or other aspects of nutritional status" (NHLBI Obesity Education Initiative, 1998, p. 91) (Evidence Category D). In addition, weight reduction prescriptions in older persons should be accompanied by proper nutritional counseling and regular body weight monitoring. (NHLBI Obesity Education Initiative, 1998, p. 91).

The possibility that a standard approach to weight loss will work differently in diverse patient populations must be considered when setting expectations about treatment outcomes (NHLBI Obesity Education Initiative, 1998, p. 97) (Evidence Category B).
**Definition:**

BMI- Body mass index (BMI) is a number calculated using the Quetelet index: weight divided by height squared (W/H²) and is commonly used to classify weight categories. BMI can be calculated using:

- Metric Units: BMI = Weight (kg) / (Height (m) x Height (m))
- English Units: BMI = Weight (lbs.) / (Height (in) x Height (in)) x 703

**Follow-Up Plan:** Proposed outline of treatment to be conducted as a result of a BMI out of normal parameters. A follow-up plan may include, but is not limited to:
- documentation of education,
- referral (for example a Registered Dietitian Nutritionist (RDN), occupational therapist, physical therapist, primary care provider, exercise physiologist, mental health professional, or surgeon) for lifestyle/behavioral therapy, pharmacological interventions, dietary supplements, exercise counseling and/or nutrition counseling.
Guidance:

* There is no diagnosis associated with this measure. * This measure is to be reported a minimum of once per reporting period for patients seen during the reporting period. * This measure may be reported by eligible professionals who perform the quality actions described in the measure based on the services provided at the time of the qualifying visit and the measure-specific denominator coding. BMI Measurement Guidance: * Height and Weight - An eligible professional or their staff is required to measure both height and weight. Both height and weight must be measured within twelve months of the current encounter and may be obtained from separate encounters. Self-reported values cannot be used. * The BMI may be documented in the medical record of the provider or in outside medical records obtained by the provider. * If the most recent documented BMI is outside of normal parameters, then a follow-up plan is documented during the encounter or during the previous twelve months of the current encounter. * If more than one BMI is reported during the measurement period, the most recent BMI will be used to determine if the performance has been met. * Review the exclusions and exceptions criteria to determine those patients that BMI measurement may not be appropriate or necessary. Follow-Up Plan Guidance: * The documented follow-up plan must be based on the most recent documented BMI, outside of normal parameters, example: "Patient referred to nutrition counseling for BMI above or below normal parameters." (See Definitions for examples of follow-up plan treatments).

Variation has been noted in studies exploring optimal BMI ranges for the elderly (see Donini et al., [2012]; Holme & Tonstad [2015]; Diehr et al. [2008]). Notably however, all these studies have arrived at ranges that differ from the standard range for ages 18 and older, which is \( \geq 18.5 \) and \(< 25 \text{ kg/m}^2 \). For instance, both Donini et al. (2012) and Holme and Tonstad (2015) reported findings that suggest that higher BMI (higher than the upper end of 25kg/m2) in the elderly may be beneficial. Similarly, worse outcomes have been associated with being underweight (at a threshold higher than 18.5 kg/m2) at age 65 (Diehr et al. 2008). Because of optimal BMI range variation recommendations from these studies, no specific optimal BMI range for the elderly is used. However, it may be appropriate to exempt certain patients from a follow-up plan by applying the exception criteria. Review the following to apply the Medical Reason exception criteria: The Medical Reason exception could include, but is not limited to, the following patients as deemed appropriate by the health care provider: * Elderly patients (65 or older) for whom weight reduction/weight gain would complicate other underlying health conditions such as the following examples: * Illness or physical disability * Mental illness, dementia, confusion * Nutritional deficiency such as Vitamin/mineral deficiency * Patients in an urgent or emergent medical situation where time is of the essence and to delay treatment would jeopardize the patient's health status.

Initial Population:

All patients 18 and older on the date of the encounter with at least one eligible encounter during the measurement period.
### Denominator:

Equals Initial Population

### Denominator Exclusions:

Patients who are pregnant
Patients receiving palliative or hospice care
Patients who refuse measurement of height and/or weight

### Numerator:

Patients with a documented BMI during the encounter or during the previous twelve months, AND when the BMI is outside of normal parameters, a follow-up plan is documented during the encounter or during the previous twelve months of the current encounter.

### Numerator Exclusions:

Not Applicable

### Denominator Exceptions:

Patients with a documented Medical Reason.
Patients in an urgent or emergent medical situation where time is of the essence and to delay treatment would jeopardize the patient's health status.

### Supplemental Data Elements:

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

#### 3.3.2 Data Entry

**Description:** Percentage of patients aged 18 years and older with a BMI documented during the current encounter or during the previous twelve months AND with a BMI outside of normal parameters, a follow-up plan is documented during the encounter or during the previous twelve months of the current encounter.

Normal Parameters: Age 18 years and older BMI => 18.5 and < 25 kg/m².

- Visit, patient age 18+
- Multiple encounter codes accepted, including dental, medical, counseling visits (Value Set 2.16.840.1.113883.3.600.1.1751). See Appendix A on using Value Sets.
- Exclude patients with pregnancy diagnosis, palliative care diagnosis (e.g., hospice, Comfort measures in IPL), patient refusals
- Document height and weight during visit or anytime in previous 12 months. Height and weight do not need to be measured on same date. System calculates BMI for most recent measurement pair.
- If BMI outside of normal range (18.5-25 kg/m²) then must perform intervention, either during present encounter or during the 12 months prior to the present encounter. Multiple possible interventions:
  - Refer or consult dietician, diagnosis “overweight” or “underweight”, POV for visit of over or underweight
  - Education code, prescribed diet education for obesity (MNT) or check Ed button on IPL for underweight or obesity related to POV
- Order medication for overweight (Value Set: 2.16.840.1.113883.3.600.1.1498), or underweight (Value Set: 2.16.840.1.113883.3.600.1.1499)
- If document patient refusal (wellness tab – personal health) for medical reason/contraindication, will be a denominator exception and not included

### 3.4 CMS 117 V8 Childhood Immunization Status

#### 3.4.1 Detail

| **Description:** | Percentage of children 2 years of age who had four diphtheria, tetanus and acellular pertussis (DTaP); three polio (IPV), one measles, mumps and rubella (MMR); three or four H influenza type B (HiB); three hepatitis B (Hep B); one chicken pox (VZV); four pneumococcal conjugate (PCV); one hepatitis A (Hep A); two or three rotavirus (RV); and two influenza (flu) vaccines by their second birthday |
| **Rationale:** | Infants and toddlers are particularly vulnerable to infectious diseases because their immune systems have not built up the necessary defenses to fight infection (Centers for Disease Control and Prevention 2017a). Most childhood vaccines are between 90 and 99 percent effective in preventing diseases (HealthyChildren 2015). Vaccination of each U.S. birth cohort with the current childhood immunization schedule prevents approximately 42,000 deaths and 20 million cases of disease and saves nearly $14 billion in direct costs and $69 billion in societal costs each year (Zhou 2014). Immunizing a child not only protects that child's health but also the health of the community, especially for those who are not immunized or are unable to be immunized due to other health complications (Centers for Disease Control and Prevention 2017b). When the majority of the community is immunized against a disease, other members of the community are also protected because herd immunity shields them. (National Institute of Allergy and Infectious Diseases 2014). |
| Clinical Recommendation Statement: | Advisory Committee on Immunization Practices (ACIP) Recommended Immunization Schedule for Children and Adolescents Aged 18 Years or Younger, United States, 2018 (Centers for Disease Control and Prevention 2018) Hepatitis B (HepB) "Minimum age: birth" "Birth Dose (Monovalent HepB vaccine only): -- Mother is HBsAg-Negative: 1 dose within 24 hours of birth for medically stable infants >2,000 grams. Infants <2,000 grams administer 1 dose at chronological age 1 month or hospital discharge. -- Mother is HBsAg-Positive: Give HepB vaccine and 0.5 mL of HBIG (at separate anatomic sites) within 12 hours of birth, regardless of birth weight. Test for HBsAg and anti-HBs at age 9-12 months. If HepB series is delayed, test 1-2 months after final dose. -- Mother's HBsAg status is unknown: Give HepB vaccine within 12 hours of birth, regardless of birth weight. For infants <2,000 grams, give 0.5 mL of HBIG in addition to HepB vaccine within 12 hours of birth. Determine mother's HBsAg status as soon as possible. If mother is HBsAg-positive, give 0.5 mL of HBIG to infants >2,000 grams as soon as possible, but no later than 7 days of age. Routine Series: -- A complete series is 3 doses at 0, 1-2, and 6-18 months. (Monovalent HepB vaccine should be used for doses given before age 6 weeks.) -- Infants who did not receive a birth dose should begin the series as soon as feasible. -- Administration of 4 doses is permitted when a combination vaccine containing HepB is used after the birth dose. -- Minimum age for the final (3rd or 4th) dose: 24 weeks. -- Minimum Intervals: Dose 1 to Dose 2: 4 weeks / Dose 2 to Dose 3: 8 weeks / Dose 1 to Dose 3: 16 weeks. (When 4 doses are given, substitute "Dose 4" for "Dose 3" in these calculations.) "Diptheria, tetanus, acellular pertussis vaccinations (DTap) "Minimum age: 6 weeks [4 years for Kinrix or Quadracel] " "Routine vaccination: -- 5-dose series at 2, 4, 6, and 15-18 months, and 4-6 years. Prospectively: A 4th dose may be given as early as age 12 months if at least 6 months have elapsed since the 3rd dose. Retrospectively: A 4th dose that was inadvertently given as early as 12 months may be counted if at least 4 months have elapsed since the 3rd dose. "Hib (Haemophilus influenzae type b) "Minimum age: 6 weeks" "Routine vaccination: -- ActHIB, Hiberix, or Pentacel: 4-dose series at 2, 4, 6, and 12-15 months. -- PedvaxHIB: 3-dose series at 2, 4, and 12-15 months." -- Polio (IPV) "Minimum age: 6 weeks" "Routine vaccination: -- 4-dose series at ages 2, 4, 6-18 months, and 4-6 years. Administer the final dose on or after the 4th birthday and at least 6 months after the previous dose." Measles, mumps, rubella (MMR) "Minimum age: 12 months for routine vaccination" "Routine vaccination: -- 2-dose series at 12-15 months and 4-6 years. -- The 2nd dose may be given as early as 4 weeks after the 1st dose. "Pneumococcal conjugate (PCV13) "Minimum age: 6 weeks [PCV13]" "Routine vaccination with PCV13: -- 4-dose series at 2, 4, 6, and 12-15 months." Varicella (Var) "Minimum age: 12 months" "Routine vaccination: - - 2-dose series: 12-15 months and 4-6 years. -- The 2nd dose may be given as early as 3 months after the 1st dose (a dose given after a 4-week interval may be counted)." Hepatitis A (HepA) "Minimum age: 12 months," "Routine vaccination: -- 2
doses, separated by 6-18 months, between the 1st and 2nd birthdays. (A series begun before the 2nd birthday should be completed even if the child turns 2 before the second dose is given.)" Rotavirus (RV) "Minimum age: 6 weeks" "Routine vaccination: Rotarix: 2-dose series at 2 and 4 months. RotaTeq: 3-dose series at 2, 4, and 6 months. If any dose in the series is either RotaTeq or unknown, default to 3-dose series. " Influenza (inactivated influenza vaccine (IIV) "Minimum age: 6 months" "Routine vaccination: -- Administer an age-appropriate formulation and dose of influenza vaccine annually. Children 6 months-8 years who did not receive at least 2 doses of influenza vaccine before July 1, 2017 should receive 2 doses separated by at least 4 weeks. "

**Definition:**
Recommended vaccines: Vaccines and the schedule of vaccines as recommended by the ACIP for children two years of age. The measure may differ slightly from the ACIP recommendations because the measure focuses on immunizations that are appropriate by age 2. Also, there may be small differences when there are shortages for a particular vaccine.

**Guidance:**
For the MMR, hepatitis B, VZV and hepatitis A vaccines, numerator inclusion criteria include: evidence of receipt of the recommended vaccine; documented history of the illness; or, a seropositive test result for the antigen. For the DTaP, IPV, Hib, pneumococcal conjugate, rotavirus, and influenza vaccines, numerator inclusion criteria include only evidence of receipt of the recommended vaccine.

Patients may be included in the numerator for a particular antigen if they had an anaphylactic reaction to the vaccine. Patients may be included in the numerator for the DTaP vaccine if they have encephalopathy. Patients may be included in the numerator for the IPV vaccine if they have had an anaphylactic reaction to streptomycin, polymyxin B, or neomycin. Patients may be included in the numerator for the influenza, MMR, or VZV vaccines if they have cancer of lymphoreticular or histiocytic tissue, multiple myeloma, leukemia, have had an anaphylactic reaction to neomycin, have Immunodeficiency, or have Human Immunodeficiency Virus (HIV). Patients may be included in the numerator for the hepatitis B vaccine if they have had an anaphylactic reaction to common baker's yeast.

The measure allows a grace period by measuring compliance with these recommendations between birth and age two. Children who turn 2 years of age during the measurement period and who have a visit during the measurement period

**Denominator:**
Equals Initial Population

**Denominator Exclusions:**
Exclude patients whose hospice care overlaps the measurement period

**Numerator:**
Children who have evidence showing they received recommended vaccines, had documented history of the illness, had a seropositive test result, or had an allergic reaction to the vaccine by their second birthday
### Numerator Exclusions:
Not Applicable

### Denominator Exceptions:
None

### Supplemental Data Elements:
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

#### 3.4.2 Data Entry

| Description | Percentage of children 2 years of age who had four DTaP; three polio (IPV); one MMR; three H influenza type B (HIB); three hepatitis B (Hep B); one chicken pox (VZV); four pneumococcal conjugate (PCV); one hepatitis A (Hep A); two or three rotavirus (RV); and two influenza (flu) vaccines by their second birthday. |

Create visit (inpatient or outpatient), child between age 1 and 2. Multiple encounters accepted:

- Office Visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1001),
- Home healthcare (Value Set 2.16.840.1.113883.3.464.1003.101.12.1016),
- Preventive care visits (Value sets 2.16.840.1.113883.3.464.1003.101.12.1022,…1024),
- Inpatient visits (Value Set 2.16.840.1.113883.3.666.5.307)

See Appendix A on using Value Sets.

Exclude patients discharged to hospice care or with documented hospice or Comfort measures in IPL (Value Set: 2.16.840.1.113762.1.4.1108.15).

Administer all vaccines from Immunization Forecaster unless documented reaction.

- Reactions are specific to vaccine type and assume age less than 2:
  - DPT: anaphylaxis or encephalopathy
  - Polio: reaction to vaccine, streptomycin, neomycin, polymyxin
  - MMR: disorder of immune system, malignancy of lymphatic or hematopoietic tissue, or neomycin adverse reaction
  - Diagnosis of Measles, Mumps or Rubella, OR IGG Antibody >1.1
  - HIB-anaphylaxis
  - Hepatitis B – presence of Anti-Hepatitis B virus surface ab
    - Anaphylactic reaction to Hepatitis B vaccine, common baker’s yeast
    - Diagnosis of Hepatitis B
  - Varicella/zoster – disorder of immune system, HIV, Malignant neoplasm of lymphatic or hematopoietic tissue, neomycin adverse reaction
- Diagnosis of varicella zoster or VZ IGG Aby test >1.10
  - Pneumococcal conjugate – Pneumococcal vaccine adverse reaction
  - Hepatitis A – Anaphylaxis, diagnosis of Hepatitis A, Anti Hepatitis A IgG Agn positive.
  - Rotavirus-anaphylaxis, severe combined immunodeficiency, intussusception.
  - Influenza – influenza virus vaccine adverse reaction, disorder of immune system, malignant neoplasm of lymphatic and hematopoietic tissue, neomycin adverse reaction.

3.5 CMS 122V8 Diabetes: Hemoglobin A1c Poor Control (> 9%)

3.5.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 18-75 years of age with diabetes who had hemoglobin A1c &gt; 9.0% during the measurement period.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>As the seventh leading cause of death in the U.S., diabetes kills approximately 79,500 people a year and affects more than 30 million Americans (9.4 percent of the U.S. population) (Centers for Disease Control and Prevention [CDC], 2017a, 2017b). Diabetes is a long-lasting disease marked by high blood glucose levels, resulting from the body's inability to produce or use insulin properly (CDC, 2017c). People with diabetes are at increased risk of serious health complications including vision loss, heart disease, stroke, kidney failure, amputation of toes, feet or legs, and premature death (CDC, 2016). In 2017, diabetes cost the U.S. an estimated $327 billion: $237 billion in direct medical costs and $90 billion in reduced productivity. This is a 34 percent increase from the estimated $245 billion spent on diabetes in 2012 (American Diabetes Association, 2018a). Controlling A1c blood levels help reduce the risk of microvascular complications (eye, kidney and nerve diseases) (CDC, 2014).</td>
</tr>
</tbody>
</table>
### Clinical Recommendation Statement:

American Diabetes Association (2018b): - A reasonable A1C goal for many nonpregnant adults is <7%. (Level of evidence: A) - Providers might reasonably suggest more stringent A1C goals (such as <6.5% [48 mmol/mol]) for selected individual patients if this can be achieved without significant hypoglycemia or other adverse effects of treatment (i.e., polypharmacy). Appropriate patients might include those with short duration of diabetes, type 2 diabetes treated with lifestyle or metformin only, long life expectancy, or no significant cardiovascular disease (CVD). (Level of evidence: C) - Less stringent A1C goals (such as <8% [64 mmol/mol]) may be appropriate for patients with a history of severe hypoglycemia, limited life expectancy, advanced microvascular or macrovascular complications, extensive comorbid conditions, or long-standing diabetes in whom the goal is difficult to achieve despite diabetes self-management education, appropriate glucose monitoring, and effective doses of multiple glucose-lowering agents including insulin. (Level of evidence: B)

### Guidance:

Patient is numerator compliant if most recent HbA1c level >9%, the most recent HbA1c result is missing, or if there are no HbA1c tests performed and results documented during the measurement period. If the HbA1c test result is in the medical record, the test can be used to determine numerator compliance. Only patients with a diagnosis of Type 1 or Type 2 diabetes should be included in the denominator of this measure; patients with a diagnosis of secondary diabetes due to another condition should not be included.

### Initial Population:

Patients 18-75 years of age with diabetes with a visit during the measurement period.

### Denominator:

Equals Initial Population.

### Denominator Exclusions:

Exclude patients whose hospice care overlaps the measurement period. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.

### Numerator:

Patients whose most recent HbA1c level (performed during the measurement period) is >9.0%.

### Numerator Exclusions:

Not Applicable.

### Denominator Exceptions:

None

### Supplemental Data Elements:

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.
3.5.2 Data Entry

| Description | Percentage of patients 18-75 years of age with diabetes who had hemoglobin A1c > 9% during the measurement period. |

Create visit, inpatient or outpatient, patient age 18-75 (Multiple visits accepted including):

- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Annual wellness visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Nursing facility visit: 2.16.840.1.113883.3.464.1003.101.12.1012
- Care services in long-term residential facility: 2.16.840.1.113883.3.464.1003.101.12.1014

See Appendix A on using Value Sets.

- Diagnosis diabetes on IPL (Value Set: 2.16.840.1.113883.3.464.1003.103.12.1001).
- Exclude hospital discharge to hospice or comfort care entered by coding. (Value Set: 2.16.840.1.113762.1.4.1108.15).
- Order and result HbA1c test (POC in Outpatient), included in numerator if >9.

3.6 CMS 124V8 Cervical Cancer Screening

3.6.1 Detail

| Description | Percentage of women 21-64 years of age who were screened for cervical cancer using either of the following criteria: 
Women age 21-64 who had cervical cytology performed every 3 years.
Women age 30-64 who had cervical cytology/human papillomavirus (HPV) co-testing performed every 5 years. |

<p>| Rationale | All women are at risk for cervical cancer. In 2018, an estimated 13,240 women were diagnosed with cervical cancer in the U.S., resulting in an estimated 4,170 deaths (Noone et al., 2018). If pre-cancerous lesions are detected early by Pap tests and treated, the likelihood of survival is nearly 100 percent (American Cancer Society, 2017). |</p>
<table>
<thead>
<tr>
<th>Clinical Recommendation Statement</th>
<th>USPSTF (2012)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&quot;The USPSTF recommends screening for cervical cancer in women aged 21 to 65 years with cytology (Papanicolaou smear) every 3 years or, for women aged 30 to 65 years who want to lengthen the screening interval, screening with a combination of cytology and HPV testing every 5 years.&quot; (A recommendation)</td>
</tr>
<tr>
<td></td>
<td>&quot;The USPSTF recommends against screening for cervical cancer in women younger than age 21 years.&quot; (D recommendation)</td>
</tr>
<tr>
<td></td>
<td>&quot;The USPSTF recommends against screening for cervical cancer in women older than age 65 years who have had adequate prior screening and are not otherwise at high risk for cervical cancer.&quot; (D recommendation)</td>
</tr>
<tr>
<td></td>
<td>&quot;The USPSTF recommends against screening for cervical cancer in women who have had a hysterectomy with removal of the cervix and who do not have a history of a high-grade precancerous lesion (cervical intraepithelial neoplasia grade 2 or 3) or cervical cancer.&quot; (D recommendation).</td>
</tr>
<tr>
<td></td>
<td>&quot;The USPSTF recommends against screening for cervical cancer with HPV testing, alone or in combination with cytology, in women younger than age 30 years.&quot; (D recommendation)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Guidance</th>
<th>To ensure the measure is only looking for a cervical cytology test only after a woman turns 21 years of age, the youngest age in the initial population is 23.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Patient self-report for procedures as well as diagnostic studies should be recorded in 'Procedure, Performed' template or 'Diagnostic Study, Performed' template in QRDA-1.</td>
</tr>
<tr>
<td></td>
<td>Include only cytology and HPV &quot;co-testing&quot;; in co-testing, both cytology and HPV tests are performed (i.e., the samples are collected and both tests are ordered, regardless of the cytology result) on the same date of service. Do not include reflex testing.</td>
</tr>
<tr>
<td></td>
<td>In addition, if the medical record indicates the HPV test was performed only after determining the cytology result, this is considered reflex testing and does not meet criteria for the measure.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Initial Population</th>
<th>Women 23-64 years of age with a visit during the measurement period.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator</td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td>Denominator Exclusions</td>
<td>Women who had a hysterectomy with no residual cervix or a congenital absence of cervix.</td>
</tr>
<tr>
<td></td>
<td>Exclude patients whose hospice care overlaps the measurement period.</td>
</tr>
</tbody>
</table>
### Numerator
Women with one or more screenings for cervical cancer. Appropriate screenings are defined by any one of the following criteria:
- Cervical cytology performed during the measurement period or the two years prior to the measurement period for women who are at least 21 years old at the time of the test.
- Cervical cytology/HPV co-testing performed during the measurement period or the four years prior to the measurement period for women who are at least 30 years old at the time of the test.

### Numerator Exclusions
Not Applicable.

### Denominator Exceptions
None.

### Supplemental Data Elements
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

#### 3.6.2 Data Entry
**Description:** Percentage of women 21-64 years of age who were screened for cervical cancer using either of the following criteria:
- Women age 21-64 who had cervical cytology performed every 3 years.
- Women age 30-64 who had cervical cytology/HPV co-testing performed every 5 years.

Visit, female, age 23-64 (at start of measurement year), (measure looks back to capture tests on 21-year old). Multiple visits accepted:

- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Figure 3-7: Palliative Care list](image)

- Excludes patients with IPL entry of congenital absence of cervix (SNOMED-CT Code 37687000) or hysterectomy with no residual cervix (Value Set: 2.16.840.1.113883.3.464.1003.198.12.1014).
See Appendix A on using Value Sets.

Include in numerator Pap Smear or Pap Smear with HPV test within 3 years of end of measurement period.

or:

Include in numerator patients over age 30 with Pap test within 5 years of end of measurement period, AND Pap and HPV test accessioned and resulted.

![Order a Lab Test dialog](image)

Figure 3-8: Order a Lab Test dialog

### 3.7 CMS 125V8 Breast Cancer Screening

#### 3.7.1 Detail

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of women 50-74 years of age who had a mammogram to screen for breast cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Breast cancer is one of the most common types of cancers, accounting for 15 percent of all new cancer diagnoses in the U.S. (Noone et al, 2018). In 2015, over 3 million women were estimated to be living with breast cancer in the U.S. and it is estimated that 12 percent of women will be diagnosed with breast cancer at some point during their lifetime (Noone et al, 2018). While there are other factors that affect a woman’s risk of developing breast cancer, advancing age is a primary risk factor. Breast cancer is most frequently diagnosed among women ages 55-64; the median age at diagnosis is 62 years (Noone et al, 2018). The chance of a woman being diagnosed with breast cancer in a given year increases with age. By age 40, the chances are 1 in 68; by age 50 it becomes 1 in 43; by age 60, it is 1 in 29 (American Cancer Society, 2017).</td>
</tr>
</tbody>
</table>
**Clinical Recommendation Statement:**

The USPSTF recommends biennial screening mammography for women aged 50-74 years (B recommendation). The decision to start screening mammography in women prior to age 50 years should be an individual one. Women who place a higher value on the potential benefit than the potential harms may choose to begin biennial screening between the ages of 40 and 49 years (C recommendation). (USPSTF, 2016)

The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of screening mammography in women aged 75 years or older (I statement). (USPSTF, 2016)

The USPSTF concludes that the current evidence is insufficient to assess the benefits and harms of digital breast tomosynthesis (DBT) as a primary screening method for breast cancer (I Statement). (USPSTF, 2016)

The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of adjunctive screening for breast cancer using breast ultrasonography, magnetic resonance imaging, DBT, or other methods in women identified to have dense breasts on an otherwise negative screening mammogram (I statement). (USPSTF, 2016).

**Guidance**

Patient self-report for procedures as well as diagnostic studies should be recorded in 'Procedure, Performed' template or 'Diagnostic Study, Performed' template in QRDA-1. This measure evaluates primary screening. Do not count biopsies, breast ultrasounds, or MRIs because.

**Initial Population:**

Women 51-74 years of age with a visit during the measurement period

**Denominator:**

Equals Initial Population

**Denominator Exclusions:**

Women who had a bilateral mastectomy or who have a history of a bilateral mastectomy or for whom there is evidence of a right and a left unilateral mastectomy. Exclude patients whose hospice care overlaps the measurement period. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.

**Numerator:**

Women with one or more mammograms during the 27 months prior to the end of the measurement period.

**Numerator Exclusions:**

Not Applicable.

**Denominator Exceptions:**

None.

**Supplemental Data Elements:**

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.
3.7.2 Data Entry

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of women 50-74 years of age who had a mammogram to screen for breast cancer.</th>
</tr>
</thead>
</table>

Visit during measurement period, female, age 50-74, multiple encounter codes accepted:

- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Annual wellness visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Nursing facility visit:
  2.16.840.1.113883.3.464.1003.101.12.1012

See Appendix A on using Value Sets.

- Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

  ![Figure 3-9: Palliative Care list](image)

- Exclude patients with diagnosis or history of bilateral mastectomy:
  - Two diagnoses of unilateral mastectomy
  - Either of the following:
    Right mastectomy
    (Value Set:2.16.840.1.113883.3.464.1003.198.12.1070), OR
    Left mastectomy (Value Set: 2.16.840.1.113883.3.464.1003.198.12.1069))
    Plus a second unspecified laterality mastectomy
- Numerator = Mammogram date within 27 months of end of measurement period.
3.8 CMS 127V8 Vaccination Status for Older Adults

3.8.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Pneumonia is a common cause of illness and death in the elderly and persons with certain underlying conditions. The major clinical syndromes of pneumococcal disease include pneumonia, bacteremia and meningitis, with pneumonia being the most common (Centers for Disease Control and Prevention [CDC], 2015a). Pneumonia symptoms generally include fever, chills, pleuritic chest pain, cough with sputum, dyspnea, tachypnea, hypoxia tachycardia, malaise and weakness. There is an estimated 400,000 cases of pneumonia in the U.S. each year and a 5 percent-7 percent mortality rate, although it may be higher among older adults and adults in nursing homes (CDC, 2015b; Janssens &amp; Krause, 2004). Among the 91.5 million US adults aged &gt; 50 years, 29,500 cases of IPD, 502,600 cases of nonbacteremic pneumococcal pneumonia and 25,400 pneumococcal-related deaths are estimated to occur yearly; annual direct and indirect costs are estimated to total $3.7 billion and $1.8 billion, respectively. Pneumococcal disease remains a substantial burden among older US adults, despite increased coverage with 23-valent pneumococcal polysaccharide vaccine, (PPV23) and indirect benefits afforded by PCV7 vaccination of young children (Weycker et al., 2011). Pneumococcal vaccines have been shown to be highly effective in preventing invasive pneumococcal disease. When comparing costs, outcomes and quality adjusted life years, immunization with the two recommended pneumococcal vaccines was found to be more economically efficient than no vaccination, with an incremental cost-effectiveness ratio of $25,841 per quality-adjusted life year gained (Chen et al., 2014).</td>
</tr>
</tbody>
</table>
**Clinical Recommendation Statement:**

In 2014, the Advisory Committee on Immunization Practices (ACIP) began recommending a dose of 13-valent pneumococcal conjugate vaccine (PCV13) be followed by a dose of 23-valent pneumococcal polysaccharide vaccine (PPSV23) 6-12 months later in adults aged 65 and older who have not previously received a pneumococcal vaccination, and in persons over the age of two years who are considered to be at higher risk for pneumococcal disease due to an underlying condition. The two vaccines should not be coadministered and intervals for administration of the two vaccines vary slightly depending on the age, risk group, and history of vaccination (Kobayashi et al., 2015). In 2015, ACIP updated its recommendation and changed the interval between PCV13 and PPSV23, from 6-12 months to at least one year for immunocompetent adults aged &ge;65 years who have not previously received pneumococcal vaccine. For immunocompromised vaccine-naive adults, the minimum acceptable interval between PCV13 and PPSV23 is 8 weeks. Both immunocompetent and immunocompromised adults aged &ge;65 years who have previously received a dose of PPSV23 when over the age of 65 should receive a dose of PCV13 at least one year after PPSV23 (&ge;1 year). Immunocompetent and immunocompromised adults aged &ge;65 who have previously received a dose of PPSV23 when under the age of 65, should also receive a dose of PCV13 at least one year after PPSV23 (&ge;1 year) and then another dose of PPSV23 at least one year after PCV13. It is recommended that for those that have this alternative three-dose schedule (2 PPSV23 and 1 PCV13), the three doses should be spread over a time period of five or more years (Kobayashi et al., 2015).

**Guidance:**

Patient self-report for procedures as well as immunizations should be recorded in ‘Procedure, Performed’ template or ‘Immunization, Administered’ template in QRDA-1. ACIP (Kobayashi, 2015) provides guidance about the proper interval and relative timing for the administration of two pneumococcal vaccines; this measure assesses whether patients have received at least one of either vaccine.

**Initial Population:**

Patients 65 years of age and older with a visit during the measurement period.

**Denominator:**

Equals Initial Population.

**Denominator Exclusions:**

Exclude patients whose hospice care overlaps the measurement period.

**Numerator:**

Patients who have ever received a pneumococcal vaccination before the end of the measurement period.

**Numerator Exclusions:**

Not Applicable.

**Denominator Exceptions:**

None.

**Supplemental Data Elements:**

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.
3.8.2 Data Entry

| Description: | Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine. |

Visit or Admission, age 65 or older, multiple encounter codes accepted:

- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Nursing facility visit (Value Set 2.16.840.1.113883.3.464.1003.101.11.1065).
- Annual wellness visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Long-term residential facility visit.  

See Appendix A on using Value Sets

- Excludes patients on hospice, Comfort measures, or discharged to hospice  
  (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Palliative Care list](image)

Numerator = Administer pneumococcal vaccine during visit or hospitalization if no history of receipt at any time in past (included if received in past or present encounter).
### 3.9 CMS 130V8 Colorectal Cancer Screening

#### 3.9.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale:</td>
<td>Colorectal cancer represents eight percent of all new cancer cases and is the second leading cause of cancer deaths in the United States. In 2018, an estimated 140,250 new cases of colorectal cancer and an estimated 50,630 deaths attributed to it. According to the National Cancer Institute, about 4.2 percent of men and women will be diagnosed with colorectal cancer at some point during their lifetimes. For most adults, older age is the most important risk factor for colorectal cancer, although being male and black are also associated with higher incidence and mortality. Colorectal cancer is most frequently diagnosed among people 65 to 74 years old (Noone et al., 2018). Screening can be effective for finding precancerous lesions (polyps) that could later become malignant, and for detecting early cancers that can be more easily and effectively treated. Precancerous polyps usually take about 10 to 15 years to develop into colorectal cancer, and most can be found and removed before turning into cancer. The five-year relative survival rate for people whose colorectal cancer is found in the early stage before it has spread is about 90 percent (American Cancer Society, 2017).</td>
</tr>
</tbody>
</table>

**Clinical Recommendation Statement:**
The U.S. Preventive Services Task Force (2016) recommends screening for colorectal cancer starting at age 50 years and continuing until age 75 years. This is a Grade A recommendation (U.S. Preventive Services Task Force, 2016). Appropriate screenings are defined by any one of the following: - Colonoscopy (every 10 years) - Flexible sigmoidoscopy (every 5 years) - Fecal occult blood test (annually) - FIT-DNA (every 3 years) - Computed tomographic colonography (every 5 years).

**Screening tests:**
- Colonoscopy (every 10 years)
- Flexible sigmoidoscopy (every 5 years)
- Fecal occult blood test (annually)
- FIT-DNA (every 3 years)
- Computed tomographic colonography (every 5 years)

**Guidance**
Patient self-report for procedures as well as diagnostic studies should be recorded in "Procedure, Performed" template or "Diagnostic Study, Performed" template in QRDA-1. Do not count DRE, FOBT tests performed in an office setting or performed on a sample collected via DRE.

**Initial Population:**
Patients 50-75 years of age with a visit during the measurement period

**Denominator:**
Equals Initial Population
**Description:** Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer

**Denominator Exclusions:** Exclude patients whose hospice care overlaps the measurement period. Patients with a diagnosis or past history of total colectomy or colorectal cancer. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.

**Numerator:** Patients with one or more screenings for colorectal cancer. Appropriate screenings are defined by any one of the following criteria:
- Fecal occult blood test (FOBT) during the measurement period.
- Flexible sigmoidoscopy during the measurement period or the four years prior to the measurement period.
- Colonoscopy during the measurement period or the nine years prior to the measurement period.
- FIT-DNA during the measurement period or the two years prior to the measurement period.
- CT Colonography during the measurement period or the four years prior to the measurement period.

**Numerator Exclusions:** Not Applicable.

**Denominator Exceptions:** None.

**Supplemental Data Elements:** For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.9.2 Data Entry

**Description:** Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer.

Visit or admission, patient age 50–75 at start of measurement period, multiple encounters codes accepted
- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307)
- Office visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1001)
- Preventive care (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1023, …1025)
- Home healthcare (Value Set 2.16.840.1.113883.3.464.1003.101.12.1016)
- Annual wellness care (Value Set 2.16.840.1.113883.3.526.3.1240)
- Nursing facility visit
- Services in long-term residential facility

See Appendix A for Value Set information.
• Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Figure 3-11: Palliative Care list](image)

Exclude patients with diagnosis of either of the following:

• Malignant Neoplasm of Colon

• Total Colectomy (Value Set: 2.16.840.1.113883.3.464.1003.108.12.1020).

before end of measurement period.

Include in numerator any of the following ordered and resulted:

• Colonoscopy within 10 years of end of measurement period.
• Sigmoidoscopy within 5 years of end of measurement period.
• FOBT (Stool guaiac) resulted during measurement period.
• FIT-DNA resulted within 3 years of end of measurement period.
• CT Colonography within 5 years of end of measurement period.

### 3.10 CMS 131V8 Diabetes: Eye Exam

#### 3.10.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 18-75 years of age with diabetes and an active diagnosis of retinopathy overlapping the measurement period who had a retinal or dilated eye exam by an eye care professional during the measurement period or diabetics with no diagnosis of retinopathy overlapping the measurement period who had a retinal or dilated eye exam by an eye care professional during the measurement period or in the 12 months prior to the measurement period.</th>
</tr>
</thead>
</table>
**Rationale:** As the seventh leading cause of death in the U.S., diabetes kills approximately 79,500 people a year and affects more than 30 million Americans (9.4 percent of the U.S. population) (Centers for Disease Control and Prevention [CDC], 2017a, 2017b). Diabetes is a long-lasting disease marked by high blood glucose levels, resulting from the body's inability to produce or use insulin properly (CDC, 2017c). People with diabetes are at increased risk of serious health complications including vision loss, heart disease, stroke, kidney failure, amputation of toes, feet or legs, and premature death (CDC, 2016). In 2017, diabetes cost the U.S. an estimated $327 billion: $237 billion in direct medical costs and $90 billion in reduced productivity. This is a 34 percent increase from the estimated $245 billion spent on diabetes in 2012 (American Diabetes Association, 2018a). Diabetic retinopathy is progressive damage to the small blood vessels in the retina that may result in loss of vision. It is the leading cause of blindness in adults between 20-74 years of age. Approximately 4.1 million adults are affected by diabetic retinopathy (CDC, 2015).

**Clinical Recommendation Statement:**
American Diabetes Association (2018b): - Adults with type 1 diabetes should have an initial dilated and comprehensive eye examination by an ophthalmologist or optometrist within 5 years after the onset of diabetes. (Level of evidence: B) - Patients with type 2 diabetes should have an initial dilated and comprehensive eye examination by an ophthalmologist or optometrist at the time of the diabetes diagnosis. (Level of evidence: B) - If there is no evidence of retinopathy for one or more annual eye exam and glycemia is well controlled, then exams every 1-2 years may be considered. If any level of diabetic retinopathy is present, subsequent dilated retinal examinations should be repeated at least annually by an ophthalmologist or optometrist. If retinopathy is progressing or sight threatening, then examinations will be required more frequently. (Level of evidence: B)

**Guidance**
Only patients with a diagnosis of Type 1 or Type 2 diabetes should be included in the denominator of this measure; patients with a diagnosis of secondary diabetes due to another condition should not be included. The eye exam must be performed by an ophthalmologist or optometrist.

**Initial Population:** Patients 18-75 years of age with diabetes with a visit during the measurement period.

**Denominator:** Equals Initial Population.

**Denominator Exclusions:** Exclude patients whose hospice care overlaps the measurement period. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.
### Numerator:
Patients with an eye screening for diabetic retinal disease. This includes diabetics who had one of the following:
- Diabetic with a diagnosis of retinopathy that overlaps the measurement period and a retinal or dilated eye exam by an eye care professional in the measurement period.
- Diabetic with no diagnosis of retinopathy overlapping the measurement period and a retinal or dilated eye exam by an eye care professional in the measurement period or the year prior to the measurement period.

### Numerator Exclusions:
Not Applicable.

### Denominator Exceptions:
None.

### Supplemental Data Elements:
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.10.2 Data Entry

**Description:** Percentage of patients 18-75 years of age with diabetes who had a retinal or dilated eye exam by an eye care professional during the measurement period or a negative retinal exam (no evidence of retinopathy) in the 12 months prior to the measurement period.

Visit or admit patient with any diabetes diagnosis (see value sets), age 18-75 at start of measurement period. Multiple encounter codes accepted:

- Inpatient (Value set 2.16.840.1.113883.3.666.5.307).
- Annual wellness visit (Value set 2.16.840.1.113883.3.526.3.1240).
- Ophthalmologic visit (Value set 2.16.840.1.113883.3.526.3.1285).
- Nursing facility visit
- Services in long-term residential facility

See Appendix A for information on using Value Sets.

- Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:
Include in numerator if any of the following:

- Diabetes eye exam (wellness tab), result = normal/negative within 1 year of start of measurement period:

- Ophthalmologic visit with documented dilated eye exam during measurement period:
- Diagnosis of diabetic retinopathy: 2.16.840.1.113883.3.526.3.327

3.11 CMS 134V8 Diabetes: Medical Attention for Nephropathy

3.11.1 Detail

| Description: | The percentage of patients 18-75 years of age with diabetes who had a nephropathy screening test or evidence of nephropathy during the measurement period. |
**Rationale:**

As the seventh leading cause of death in the U.S., diabetes kills approximately 79,500 people a year and affects more than 30 million Americans (9.4 percent of the U.S. population) (CDC, 2017a, 2017b). Diabetes is a long-lasting disease marked by high blood glucose levels, resulting from the body's inability to produce or use insulin properly (CDC, 2017c). People with diabetes are at increased risk of serious health complications including vision loss, heart disease, stroke, kidney failure, amputation of toes, feet or legs, and premature death (CDC, 2016). In 2017, diabetes cost the U.S. an estimated $327 billion: $237 billion in direct medical costs and $90 billion in reduced productivity. This is a 34 percent increase from the estimated $245 billion spent on diabetes in 2012 (American Diabetes Association, 2018a). High blood sugar levels in patients with diabetes put them at a higher risk of damaging their kidneys and causing chronic kidney disease, which can lead to kidney failure (CDC, 2016, 2017d). During 2011-2012 there were 36.5% new cases of chronic kidney disease (stages 1-4) among 297,000 diabetic patients 20 years and older (Murphy et al., 2016). In 2014, diabetes accounted for 44% of 118,000 new cases of end stage renal disease (United States Renal Data System, 2016).
### Clinical Recommendation Statement:

| American Diabetes Association (2018b): Screening | At least once a year, assess urinary albumin (e.g., spot urinary albumin-to-creatinine ratio [UACR]) and estimated glomerular filtration rate (eGFR) in patients with type 1 diabetes duration of greater than or equal to 5 years in all patients with type 2 diabetes, and in all patients with comorbid hypertension. (Level of evidence: B) Treatment - An angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) is not recommended for the primary prevention of diabetic kidney disease in patients with diabetes who have normal blood pressure, normal UACR (<30 mg/g creatinine), and normal estimated glomerular filtration rate. (Level of evidence: B) - Either an ACE inhibitor or ARB is suggested for the treatment of the nonpregnant patient with modestly elevated UACR (30-299 mg/g creatinine) (Level of evidence: B) and is strongly recommended for those with urinary albumin to creatinine ratio >=300 mg/g creatinine and/or estimated glomerular filtration rate < 60 mL/min/1.73.m2. (Level of evidence: A) - Periodically monitor serum creatinine and potassium levels for the development of increased creatinine or changes in potassium when ACE inhibitors, angiotensin receptor blockers, or diuretics are used. (Level of evidence: B) - Continued monitoring of UACR in patients with albuminuria treated with an ACE inhibitor or ARBs is reasonable to assess the response to treatment and progression of diabetic kidney disease. (Level of evidence: E) - When estimated glomerular filtration rate is <60 mL/min/1.73 m2, evaluate and manage potential complications of chronic kidney disease. (Level of evidence: E) - Patients should be referred for evaluation for renal replacement treatment if they have an estimated glomerular filtration rate <30 mL/min/1.73 m2. (Level of evidence: A) - Promptly refer to a physician experienced in the care of kidney disease for uncertainty about the etiology of kidney disease, difficult management issues, and rapidly progressing kidney disease. (Level of evidence: B) American Association of Clinical Endocrinologists & American College of Endocrinology (2015): - Beginning 5 years after diagnosis in patients with type 1 diabetes (if diagnosed before age 30) or at diagnosis in patients with type 2 diabetes and those with type 1 diabetes diagnosed after age 30, annual assessment of serum creatinine to determine the estimated glomerular filtration rate (eGFR) and urine albumin excretion rate (AER) should be performed to identify, stage, and monitor progression of diabetic nephropathy (Grade C; best evidence level 3). - Patients with nephropathy should be counseled regarding the need for optimal glycemic control, blood pressure control, dyslipidemia control, and smoking cessation (Grade B; best evidence level 2). - In addition, they should have routine monitoring of albuminuria, kidney function electrolytes, and lipids (Grade B; best evidence level 2). - Associated conditions such as anemia and bone and mineral disorders should be assessed as kidney function declines (Grade D; best evidence level 4). - Referral to a nephrologist is recommended well before the need for renal replacement therapy (Grade D; best evidence level 4). |
**Guidance:**
Only patients with a diagnosis of Type 1 or Type 2 diabetes should be included in the denominator of this measure; patients with a diagnosis of secondary diabetes due to another condition should not be included.

**Initial Population:**
Patients 18-75 years of age with diabetes with a visit during the measurement period.

**Denominator:**
Equals Initial Population.

**Denominator Exclusions:**
Exclude patients whose hospice care overlaps the measurement period. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.

**Numerator:**
Patients with a screening for nephropathy or evidence of nephropathy during the measurement period.

**Numerator Exclusions:**
Not Applicable.

**Denominator Exceptions:**
None

**Supplemental Data Elements:**
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.11.2 Data Entry

**Description:**
The percentage of patients 18-75 years of age with diabetes who had a nephropathy screening test or evidence of nephropathy during the measurement period.

Visit or admission, patient with diagnosis of diabetes, age 18–75 at start of measurement period. Multiple encounter codes accepted.

- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Annual wellness visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Nursing facility visit
- Long-term residential facility visit

See Appendix A for information about using Value Sets.
Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Palliative Care list](image)

Figure 3-15: Palliative Care list

Include in numerator:

- Rx for ACE Inhibitor or ARB (active, can be outside med or internal order, processed) (Value Set: 2.16.840.1.113883.3.526.3.1139).

- Nephropathy diagnosis (multiple options) including:

- Procedures including:

- Diagnoses including:
  - Dialysis care as POV.

- Education:


- Lab tests for urine protein (multiple including POC if set up) resulted during measurement period (Value Set: 2.16.840.1.113883.3.464.1003.109.12.1024).
3.12 CMS 137V8 Initiation and Engagement of Alcohol and Other Drug Dependence Treatment

3.12.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 13 years of age and older with a new episode of alcohol or other drug abuse or (AOD) dependence who received the following. Two rates are reported. a. Percentage of patients who initiated treatment including either an intervention or medication for the treatment of AOD abuse or dependence within 14 days of the diagnosis b. Percentage of patients who engaged in ongoing treatment including two additional interventions or a medication for the treatment of AOD abuse or dependence within 34 days of the initiation visit. For patients who initiated treatment with a medication, at least one of the two engagement events must be a treatment intervention.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stratification:</td>
<td>Report a total score, and each of the following strata: Stratum 1: Patients age 13-17. Stratum 2: Patients age &gt;=18.</td>
</tr>
<tr>
<td>Rationale:</td>
<td>There are more deaths, illnesses, and disabilities from substance abuse than from any other preventable health condition. Treatment of medical problems caused by substance use and abuse places a huge burden on the health care system (Schneider Institute for Health Policy, 2001). In 2011, an estimated 22.5 million persons (8.7 percent of the population aged 12 or older) were classified with substance dependence or abuse. Of these, 68 percent were dependent on or abused alcohol, but not illicit drugs, 13 percent abused or were dependent on both alcohol and illicit drugs, while 19 percent were dependent on or abused illicit drugs, but not alcohol (Substance Abuse and Mental Health Services Administration [SAMHSA], 2012). Of the 19.3 million persons aged 12 or older in 2011 who were classified as needing substance use treatment but did not receive treatment at a specialty facility in the past year, 912,000 persons (4.7 percent) reported that they felt they needed treatment for their illicit drug or alcohol use problem. Of these 912,000 persons who felt they needed treatment, only 281,000 (30.8 percent) reported that they made an effort to get treatment, while 631,000 (69.2 percent) reported making no effort to get treatment (SAMHSA, 2012).</td>
</tr>
</tbody>
</table>
Clinical Recommendation Statement:

American Psychiatric Association (2006) * Because many substance use disorders are chronic, patients usually require long-term treatment, although the intensity and specific components of treatment may vary over time [I rating]. * It is important to intensify the monitoring for substance use during periods when the patient is at a high risk of relapsing, including during the early stages of treatment, times of transition to less intensive levels of care, and the first year after active treatment has ceased [I rating]. * Outpatient treatment of substance use disorders is appropriate for patients whose clinical condition or environmental circumstances do not require a more intensive level of care [I rating]. As in other treatment settings, a comprehensive approach is optimal, using, where indicated, a variety of psychotherapeutic and pharmacological interventions along with behavioral monitoring [I rating]. * Naltrexone, injectable naltrexone, acamprosate, a -aminobutyric acid (GABA) are recommended for patients with alcohol dependence [I rating]. Disulfiram is also recommended for patients with alcohol dependence [II rating]. * Methadone and buprenorphine are recommended for patients with opioid dependence [I rating]. Naltrexone is an alternative strategy [I rating]. Michigan Quality Improvement Consortium (2017) *Patients with substance use disorder or risky substance use: Patient Education and Brief Intervention by PCP or Trained Staff (e.g., RN, MSW) *If diagnosed with substance use disorder or risky substance use, initiate an intervention within 14 days. *Frequent follow-up is helpful to support behavior change; preferably 2 visits within 30 days. *Refer to a substance abuse health specialist, an addiction physician specialist, or a physician experienced in pharmacologic management of addiction. U.S. Preventive Services Task Force recommendation statement (2012): *The USPSTF recommends that clinicians screen adults aged 18 years or older for alcohol misuse and provide persons engaged in risky or hazardous drinking with brief behavioral counseling interventions to reduce alcohol misuse.

Definitions

The initiation visit is the first visit for alcohol or other drug dependence treatment within 14 days after a diagnosis of alcohol or other drug dependence. Treatment includes inpatient AOD admissions, outpatient visits, intensive outpatient encounters or partial hospitalization. The Intake Period: January 1-November 14 of the measurement year. The Intake Period is used to capture new episodes of Alcohol or Drug Dependence. The November 14 cut-off date ensures that all services can occur before the measurement period ends.

Guidance:

The new episode of alcohol and other drug dependence should be the first episode of the measurement period that is not preceded in the 60 days prior by another episode of alcohol or other drug dependence.

Initial Population:

Patients age 13 years of age and older who were diagnosed with a new episode of alcohol, opioid, or other drug abuse or dependency during a visit between January 1 and November 14 of the measurement period.
### Denominator:
Equals Initial Population..

### Denominator Exclusions:
Patients with a previous active diagnosis of alcohol, opioid, or other drug abuse or dependence in the 60 days prior to the first episode of alcohol or drug dependence. Exclude patients whose hospice care overlaps the measurement period.

### Numerator:
Numerator 1: Initiation of treatment includes either an intervention or medication for the treatment of AOD abuse or dependence within 14 days of the diagnosis Numerator 2: Engagement in ongoing treatment includes two additional interventions or a medication for the treatment of AOD abuse or dependence within 34 days of the initiation visit. For patients who initiated treatment with a medication, at least one of the two engagement events must be a treatment intervention (i.e., engagement for these members cannot be satisfied with medication treatment alone).

### Numerator Exclusions:
Not Applicable.

### Denominator Exceptions:
None.

### Supplemental Data Elements:
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.12.2 Data Entry

**Description:** Percentage of patients 13 years of age and older with a new episode of AOD dependence who received the following. Two rates are reported:
- Percentage of patients who initiated treatment within 14 days of the diagnosis.
- Percentage of patients who initiated treatment and who had two or more additional services with an AOD diagnosis within 30 days of the initiation visit.

Visit or admit, patient 13 years or older at start of measurement period. Multiple encounter codes accepted:
- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Discharge services – hospital same day discharge (Value Set 2.16.840.1.113883.3.464.1003.101.12.1006).
- ER visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1010).
• Hospital observation care (Value Set 2.16.840.1.113883.3.464.1003.101.12.1002).
• Hospital inpatient visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1004).
• Telehealth services 2.16.840.1.113883.3.464.1003.101.12.1031
• Alcohol and Drug Dependence Treatment: 2.16.840.1.113883.3.464.1003.106.12.1005
• Psych visit, Psychotherapy: 2.16.840.1.113883.3.526.3.1496

See Appendix A for information on using Value Sets.

• Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Figure 3-16: Palliative Care list](image)

Diagnosis of alcohol or substance abuse must be the first time diagnosed during the current measurement period (any prior diagnosis, even in a different measurement period, must be at least 60 days earlier), otherwise exclude patient from denominator.

![Figure 3-17: Problem Management tab](image)

Diagnosis of alcohol or substance abuse must be within the first 319 days of the measurement period (Jan 1 to Nov 15) to allow time for follow up.

Document Alcohol and Drug Treatment on IPL (TREG) (Any of the following):

![Figure 3-18: Care Planning Activities](image)
Numerators (2) (if Medications are used for treatment, they must be finished by pharmacy):

- Encounter for psychotherapy or “alcohol and drug dependence treatment” within 14 days of first diagnosis:
  - Stratification 1: Age 13<18 at start of measurement period.
  - Stratification 2: Age 18+.
- Two or more encounters for psychotherapy or alcohol and drug dependence treatment within 30 days of first diagnosis:
  - Stratification 1: Age 13<18 at start of measurement period.
  - Stratification 2: Age 18+.

3.13 CMS 138V8 Preventive Care and Screening: Tobacco Use: Screening and Cessation Intervention

3.13.1 Detail

| Description:                                                                 | Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. Three rates are reported: |
|                                                                            | • Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months. |
|                                                                            | • Percentage of patients aged 18 years and older who were screened for tobacco use and identified as a tobacco user who received tobacco cessation intervention. |
|                                                                            | • Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. |
| Rationale:                                                               | This measure is intended to promote adult tobacco screening and tobacco cessation interventions for those who use tobacco products. There is good evidence that tobacco screening and brief cessation intervention (including counseling and/or pharmacotherapy) is successful in helping tobacco users quit. Tobacco users who are able to stop using tobacco lower their risk for heart disease, lung disease, and stroke. |
**Clinical Recommendation Statement:**

The USPSTF recommends that clinicians ask all adults about tobacco use, advise them to stop using tobacco, and provide behavioral interventions and U.S. Food and Drug Administration (FDA)-approved pharmacotherapy for cessation to adults who use tobacco (Grade A Recommendation) (U.S. Preventive Services Task Force, 2015). The USPSTF recommends that clinicians ask all pregnant women about tobacco use, advise them to stop using tobacco, and provide behavioral interventions for cessation to pregnant women who use tobacco (Grade A Recommendation) (U.S. Preventive Services Task Force, 2015). The USPSTF concludes that the current evidence is insufficient to recommend electronic nicotine delivery systems for tobacco cessation in adults, including pregnant women. The USPSTF recommends that clinicians direct patients who smoke tobacco to other cessation interventions with established effectiveness and safety (previously stated) (Grade I Statement) (U.S. Preventive Services Task Force, 2015).

**Improvement Notation:**

Higher score indicates better quality.

**Definition:**

Tobacco Use - Includes any type of tobacco Tobacco Cessation Intervention - Includes brief counseling (3 minutes or less), and/or pharmacotherapy -- Note: Concepts aligned with brief counseling (e.g., minimal and intensive advice/counseling interventions conducted both in person and over the phone) are included in the value set for the numerator. Other concepts such as written self-help materials (e.g., brochures, pamphlets) and complementary/alternative therapies are not included in the value set and do not qualify for the numerator. Brief counseling also may be of longer duration or be performed more frequently, as evidence shows there is a dose-response relationship between the intensity of counseling provided (either length or frequency) and tobacco cessation rates (U.S. Preventive Services Task Force, 2015).
| Guidance: | If a patient uses any type of tobacco (i.e., smokes or uses smokeless tobacco), the expectation is that they should receive tobacco cessation intervention: either counseling and/or pharmacotherapy. In order to promote a team-based approach to patient care, the tobacco cessation intervention can be performed by another healthcare provider; therefore, the tobacco use screening and tobacco cessation intervention do not need to be performed by the same provider or clinician. To satisfy the intent of this measure, a patient must have at least one tobacco use screening during the 24-month period. If a patient has multiple tobacco use screenings during the 24-month period, only the most recent screening, which has a documented status of tobacco user or tobacco non-user, will be used to satisfy the measure requirements. If tobacco use status of a patient is unknown, the patient does not meet the screening component required to be counted in the numerator and should be considered a measure failure. Instances where tobacco use status of "unknown" is recorded include: 1) the patient was not screened; or 2) the patient was screened, and the patient (or caregiver) was unable to provide a definitive answer. If the patient does not meet the screening component of the numerator but has an allowable medical exception, then the patient should be removed from the denominator of the measure and reported as a valid exception. The medical reason exception may be applied to either the screening data element OR to any of the applicable tobacco cessation intervention data elements (counseling and/or pharmacotherapy) included in the measure. If a patient has a diagnosis of limited life expectancy, that patient has a valid denominator exception for not being screened for tobacco use or for not receiving tobacco use cessation intervention (counseling and/or pharmacotherapy) if identified as a tobacco user. As noted above in a recommendation statement from the USPSTF, the current evidence is insufficient to recommend electronic nicotine delivery systems (ENDS) including electronic cigarettes for tobacco cessation. Additionally, ENDS are not currently classified as tobacco in the recent evidence review to support the update of the USPSTF recommendation given that the devices do not burn or use tobacco leaves. In light of the current lack of evidence, the measure does not currently capture e-cigarette usage as either tobacco use or a cessation aid. The requirement of two or more visits is to establish that the eligible professional or eligible clinician has an existing relationship with the patient for certain types of encounters. This measure contains three reporting rates which aim to identify patients who were screened for tobacco use (rate/population 1), patients who were identified as tobacco users and who received tobacco cessation intervention (rate/population 2), and a comprehensive look at the overall performance on tobacco screening and cessation intervention (rate/population 3). |
| Guidance: (cont)                                                                 | By separating this measure into various reporting rates, the eligible professional or eligible clinician will be able to better ascertain where gaps in performance exist and identify opportunities for improvement. The overall rate (rate/population 3) can be utilized to compare performance to published versions of this measure prior to the 2018 performance year, when the measure had a single performance rate. For accountability reporting in the CMS MIPS program, the rate for population 2 is used for performance. The denominator of population criteria 2 is a subset of the resulting numerator for population criteria 1, as population criteria 2 is limited to assessing if patients identified as tobacco users received an appropriate tobacco cessation intervention. For all patients, population criteria 1 and 3 are applicable, but population criteria 2 will only be applicable for those patients who are identified as tobacco users. Therefore, data for every patient that meets the initial population criteria will only be submitted for population 1 and 3, whereas data submitted for population 2 will be for a subset of patients who meet the initial population criteria, as the denominator has been further limited to those who were identified as tobacco users. |
| Initial Population:                                                            | All patients aged 18 years and older seen for at least two visits or at least one preventive visit during the measurement period |
| Denominator Exclusions:                                                        | None |
| Numerator:                                                                    | Population 1: Patients who were screened for tobacco use at least once within 24 months. Population 2: Patients who received tobacco cessation intervention. Population 3: Patients who were screened for tobacco use at least once within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. |
| Numerator Exclusions:                                                         | Not Applicable. |
| Denominator Exceptions:                                                       | Population 1: Documentation of medical reason(s) for not screening for tobacco use (e.g., limited life expectancy, other medical reason) Population 2: Documentation of medical reason(s) for not providing tobacco cessation intervention (e.g., limited life expectancy, other medical reason) Population 3: Documentation of medical reason(s) for not screening for tobacco use OR for not providing tobacco cessation intervention for patients identified as tobacco users (e.g., limited life expectancy, other medical reason) |
| Supplemental Data Elements:                                                    | For every patient evaluated by this measure also identify payer, race, ethnicity, and sex. |
3.13.2 Data Entry

**Description:**
Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user.

Three rates are reported:
- Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months.
- Percentage of patients aged 18 years and older who were screened for tobacco use and identified as a tobacco user who received tobacco cessation intervention.
- Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user.

There are three rates:
Multiple codes accepted for encounter type

- Annual Wellness visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Health & Behavioral Assessment, Reassessment (Value Set 2.16.840.1.113883.3.526.3.1245).
- Occupational Therapy Evaluation (Value Set 2.16.840.1.113883.3.526.3.1011).
- Ophthalmological Services (Value Set 2.16.840.1.113883.3.526.3.1285).
- Preventive Care Services (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1023, .....
- Preventive Care Services-Individual Counseling (2.16.840.1.113883.3.464.1003.101.12.1026).
- Preventive Care Services - Group Counseling (2.16.840.1.113883.3.464.1003.101.12.1027).
- Preventive Care Services - Other (2.16.840.1.113883.3.464.1003.101.12.1030).
- Psych Visit - Diagnostic Evaluation (Value Set 2.16.840.1.113883.3.526.3.1492).
- Psych Visit – Psychotherapy (Value Set 2.16.840.1.113883.3.526.3.1496).
Psychoanalysis (Value Set 2.16.840.1.113883.3.526.3.1141).

Speech and Hearing Evaluation (Value Set 2.16.840.1.113883.3.526.3.1530).

See Appendix A for information about using Value Sets.

Document smoking status in Health Factors:

![Select the appropriate health factor](image)

Optional free text comment

Figure 3-19: Add Health Factors

Document intervention if identified as a tobacco user:

Counseling using Patient Ed (TO-QT):
Order or active prescription for tobacco cessation therapy (Value Set: 2.16.840.1.113883.3.526.3.1190).

Change health factor to non-smoker/non-tobacco user:
Figure 3-22: Add Health Factor dialog

First rate, screening only:

- **Denominator 1:**
  - Office visit age 18 or older at start of measurement period.
  - Either two office visits, OR one preventive care visit (See Value Sets) during measurement period.

- **Numerator 1:**
  - Screened within 24 months of end of measurement period.
  - Denominator Exceptions for limited life expectancy diagnosis (IPL) (Value Set: 2.16.840.1.113883.3.526.3.1259).
  - Or not screened for medical reasons (Value Set: 2.16.840.1.113883.3.526.3.1007).

Second Rate, tobacco users who received tobacco cessation intervention.

- **Denominator 2:**
  - Office visit age 18 or older:
  - Either two office visits or one preventive care visit during measurement period:
  - Screened and identified as tobacco user:

- **Numerator 2:**
  - Receiving cessation counseling for tobacco user after positive screen during measurement period (Value Set: 2.16.840.1.113883.3.526.3.509).
  - Taking tobacco cessation pharmacotherapy during measurement period (Value Set: 2.16.840.1.113883.3.526.3.1190).
  - Receive prescription for tobacco cessation pharmacotherapy after positive screen Denominator exceptions for limited life expectancy, counseling or pharmacotherapy not ordered for medical reasons.
Third Rate, patients screened and receive intervention if positive tobacco use screen:

- **Denominator:**
  - Same as denominator 1 (total patients screened, age 18+, 2 office visits or 1 preventive care visit).

- **Numerator 3:**
  - If screen positive, cessation counseling for tobacco user starts after positive screen during measurement period.
  - If screen positive, actively taking medication for tobacco cessation during measurement period.
  - If screen positive, new order for tobacco use cessation after positive screen during measurement period.

- **Denominator exceptions:**
  - Limited life expectancy diagnosis.
  - Tobacco use cessation counseling not performed for medical reason.
  - Tobacco use screening nor performed for medical reason.
  - Tobacco use cessation pharmacotherapy not ordered for medical reason.

### 3.14 CMS 139V8 Falls: Screening for Future Fall Risk

#### 3.14.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 65 years of age and older who were screened for future fall risk during the measurement period.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rationale:</strong></td>
<td>As the leading cause of both fatal and nonfatal injuries for older adults, falls are one of the most common and significant health issues facing people aged 65 years or older (Schneider, Shubert and Harmon 2010). Moreover, the rate of falls increases with age (Dykes et al. 2010). Older adults are five times more likely to be hospitalized for fall-related injuries than any other cause-related injury. It is estimated that one in every three adults over 65 will fall each year (Centers for Disease Control and Prevention 2015). In those over age 80, the rate of falls increases to fifty percent (Doherty et al. 2009). Falls are also associated with substantial cost and resource use, approaching $30,000 per fall hospitalization Woolcott et al. 2011). Identifying at-risk patients is the most important part of management, as applying preventive measures in this vulnerable population can have a profound effect on public health (al-Aama 2011). Family physicians have a pivotal role in screening older patients for risk of falls, and applying preventive strategies for patients at risk (al-Aama 2011).</td>
</tr>
</tbody>
</table>
Clinical Recommendation Statement: All older persons who are under the care of a health professional (or their caregivers) should be asked at least once a year about falls. (AGS/BGS/AAOS 2010) Older persons who present for medical attention because of a fall, report recurrent falls in the past year, or demonstrate abnormalities of gait and/or balance should have a fall evaluation performed. This evaluation should be performed by a clinician with appropriate skills and experience, which may necessitate referral to a specialist (e.g., geriatrician). (AGS/BGS/AAOS 2010)

Definition: Screening for Future Fall Risk: Assessment of whether an individual has experienced a fall or problems with gait or balance. A specific screening tool is not required for this measure, however potential screening tools include the Morse Fall Scale and the timed Get-Up-And-Go test.

Fall: A sudden, unintentional change in position causing an individual to land at a lower level, on an object, the floor, or the ground, other than as a consequence of sudden onset of paralysis, epileptic seizure, or overwhelming external force.

Guidance: None

Initial Population: Patients aged 65 years and older with a visit during the measurement period.

Denominator: Equals Initial Population.

Denominator Exclusions: Exclude patients whose hospice care overlaps the measurement period.
Exclude patients who were non-ambulatory at some point in the measurement period.

Numerator: Patients who were screened for future fall risk at least once within the measurement period.

Numerator Exclusions: Not Applicable

Denominator Exceptions: None

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

3.14.2 Data Entry

Description: Percentage of patients 65 years of age and older who were screened for future fall risk during the measurement period.

Visit patient age 65 or older at start of measurement period.

Many qualified visits:
- Annual Wellness Visit (Value Set 2.16.840.1.113883.3.526.3.1240).
- Care Services in Long-Term Residential Facility
- Discharge Services - Nursing Facility
- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).
- Ophthalmological Services (Value Set 2.16.840.1.113883.3.526.3.1285).
- Preventive Care Services
- Preventive Care Services-Individual Counseling

See Appendix A for information on using Value Sets.

- Excludes patients on hospice, Comfort measures, or discharged to hospice
  (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered
  by Coding. Outpatient entered on IPL/TREG:

![Palliative Care](image)

Figure 3-23: Palliative Care list

- Exclude patients not ambulatory

Fall risk screen (Exam - fall risk) and result

3.15 CMS 144V8 Heart Failure: Beta-Blocker Therapy for Left Ventricular Systolic Dysfunction

3.15.1 Detail

| Description: | Percentage of patients aged 18 years and older with a diagnosis of heart failure (HF) with a current or prior left ventricular ejection fraction (LVEF) < 40% who were prescribed beta-blocker therapy either within a 12-month period when seen in the outpatient setting OR at each hospital discharge. |
### Rate Aggregation:

This measure is intended to have one reporting rate, which aggregates the following populations into a single performance rate for reporting purposes:
- **Population 1:** Patients who were prescribed beta-blocker therapy within a 12-month period when seen in the outpatient setting.
- **Population 2:** Patients who were prescribed beta-blocker therapy at each hospital discharge.

For the purposes of this measure, a single performance rate can be calculated as follows:

\[
\text{Performance Rate} = \frac{\text{Numerator 1} + \text{Numerator 2}}{\text{Denominator 1} - \text{Denominator Exceptions 1} + \text{Denominator 2} - \text{Denominator Exceptions 2}}
\]

### Rationale:

Beta-blockers are recommended for all patients with stable heart failure and left ventricular systolic dysfunction, unless contraindicated. Treatment should be initiated as soon as a patient is diagnosed with left ventricular systolic dysfunction and does not have low blood pressure, fluid overload, or recent treatment with an intravenous positive inotropic agent. Beta-blockers have been shown to lessen the symptoms of heart failure, improve the clinical status of patients, reduce future clinical deterioration, and decrease the risk of mortality and the combined risk of mortality and hospitalization.
Clinical Recommendation Statement:

Use of 1 of the 3 beta blockers proven to reduce mortality (e.g., bisoprolol, carvedilol, and sustained-release metoprolol succinate) is recommended for all patients with current or prior symptoms of HFrEF [heart failure with reduced ejection fraction], unless contraindicated, to reduce morbidity and mortality (Class I, Level of Evidence: A) (ACCF/AHA, 2013). Treatment with a beta blocker should be initiated at very low doses [see excerpt from guideline table below] followed by gradual increments in dose if lower doses have been well tolerated... Clinicians should make every effort to achieve the target doses of the beta blockers shown to be effective in major clinical trials. Even if symptoms do not improve, long-term treatment should be maintained to reduce the risk of major clinical events. Abrupt withdrawal of treatment with a beta blocker can lead to clinical deterioration and should be avoided (ACCF/AHA, 2013).

Drugs Commonly Used for Stage C HFrEF (abbreviated to align with focus of measure to include only Beta-blocker therapy)

<table>
<thead>
<tr>
<th>Drug Initial Daily Dose(s)</th>
<th>Maximum Dose(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean Doses Achieved in Clinical Trials</td>
<td></td>
</tr>
<tr>
<td>Beta Blockers</td>
<td></td>
</tr>
<tr>
<td>Bisoprolol 1.25 mg once 10 mg once 8.6 mg/d Carvedilol 3.125 mg twice 50 mg twice 37 mg/d Carvedilol CR 10 mg once 80 mg once N/A Metoprolol succinate 12.5 to 25 mg once 200 mg once 159 mg/d extended release (metoprolol CR/XL) For the hospitalized patient: In patients with HFrEF experiencing a symptomatic exacerbation of HF requiring hospitalization during chronic maintenance treatment with GDMT [guideline-directed medical therapy; GDMT represents optimal medical therapy as defined by ACCF/AHA guideline-recommended therapies (primarily Class I)], it is recommended that GDMT be continued in the absence of hemodynamic instability or contraindications (Class I, Level of Evidence: B) (ACCF/AHA, 2013). Initiation of beta-blocker therapy is recommended after optimization of volume status and successful discontinuation of intravenous diuretics, vasodilators, and inotropic agents. Beta-blocker therapy should be initiated at a low dose and only in stable patients. Caution should be used when initiating beta blockers in patients who have required inotropes during their hospital course (Class I, Level of Evidence: B) (ACCF/AHA, 2013).</td>
<td></td>
</tr>
</tbody>
</table>
**Guidance:**
A range value should satisfy the logic requirement for 'Ejection Fraction' as long as the ranged observation value clearly meets the less than 40% threshold noted in the denominator logic. A range that is inclusive of or greater than 40% would not meet the measure requirement.

To satisfy this measure, it must be reported for all heart failure patients at least once during the measurement period if seen in the outpatient setting. If the patient has an eligible inpatient discharge during the measurement period, as defined in the measure logic, it is expected to be reported at each hospital discharge.

Beta-blocker therapy: For patients with prior LVEF < 40%, beta-blocker therapy should include bisoprolol, carvedilol, or sustained release metoprolol succinate.

The requirement of two or more visits is to establish that the eligible professional or eligible clinician has an existing relationship with the patient.

<table>
<thead>
<tr>
<th>Initial Population:</th>
<th>All patients aged 18 years and older with a diagnosis of heart failure.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator:</td>
<td>Equals Initial Population with a current or prior LVEF &lt; 40%.</td>
</tr>
<tr>
<td>Denominator Exclusions:</td>
<td>None.</td>
</tr>
<tr>
<td>Numerator:</td>
<td>Patients who were prescribed beta-blocker therapy either within a 12-month period when seen in the outpatient setting OR at each hospital discharge.</td>
</tr>
<tr>
<td>Numerator Exclusions:</td>
<td>Not Applicable.</td>
</tr>
<tr>
<td>Denominator Exceptions:</td>
<td>Documentation of medical reason(s) for not prescribing beta-blocker therapy (e.g., low blood pressure, fluid overload, asthma, patients recently treated with an intravenous positive inotropic agent, allergy, intolerance, other medical reasons). Documentation of patient reason(s) for not prescribing beta-blocker therapy (e.g., patient declined, other patient reasons). Documentation of system reason(s) for not prescribing beta-blocker therapy (e.g., other reasons attributable to the healthcare system).</td>
</tr>
<tr>
<td>Supplemental Data Elements:</td>
<td>For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.</td>
</tr>
</tbody>
</table>

### 3.15.2 Data Entry

| Description: | Percentage of patients aged 18 years and older with a diagnosis of HF with a current or prior LVEF < 40 % who were prescribed beta-blocker therapy either within a 12-month period when seen in the outpatient setting OR at each hospital discharge. |

Two rates looking for beta-blocker use in OPD (w/in 12 months) or at hospital discharge for patients with heart failure and low LVEF.
Multiple encounter codes accepted:

- Care Services in Long-Term Residential Facility
  (Value Set 2.16.840.1.113883.3.464.1003.101.12.1014)
- Discharge Services - Hospital Inpatient
  (Value Set 2.16.840.1.113883.3.464.1003.101.12.1007)
- Home Healthcare Services (Value Set
  2.16.840.1.113883.3.464.1003.101.12.1016)
- Nursing Facility Visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1012)
- Office Visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1001)
- Outpatient Consultation (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1008)

See Appendix A for information on using Value Sets

Denominator 1 (outpatient):

- Patient, age 18 or older at start of measurement period, with at least 2 visits
  during measurement period, diagnosis “heart failure” on IPL.
- (Value Set: 2.16.840.1.113883.3.526.3.376) with moderate to severe LVSD
  (<40% ejection fraction) (Value Set: 2.16.840.1.113883.3.526.3.1090).
- Exceptions:
  - Beta blocker not ordered for medical reason
    (Value Set: 2.16.840.1.113883.3.526.3.1007).
  - patient reason (Value Set: 2.16.840.1.113883.3.526.3.1008).
  - or system reason (Value Set: 2.16.840.1.113883.3.526.3.1009) --
  - Arrhythmia, hypotension, asthma, bradycardia diagnosis in IPL at time of
    visit, allergy to beta blocker or intolerance to beta blocker

Denominator exceptions:

- Allergy/intolerance to beta blocker therapy ingredient
  (Value Set: 2.16.840.1.113883.3.526.3.1493)
- AV block without pacer (Value Set: 2.16.840.1.113883.3.526.3.367)

Numerator 1: Active order or prescription for Beta blocker medication

Denominator 2 (Inpatient): Patient 18+ with diagnosis of Heart Failure at time of
discharge and low LVSD (<40%)

Numerator 2: Order or active medication order for Beta Blocker therapy as Discharge Medication

Denominator exceptions:

- Bradycardia (<50) during and before hospitalization
- Medication not ordered for medical, patient, system reasons
- Arrhythmia, hypotension, asthma, bradycardia
- Allergy/intolerance to Beta Blocker therapy
- Allergy/intolerance to beta blocker therapy ingredient
- AV block without pacer

### 3.16 CMS 155V8 Weight Assessment and Counseling for Nutrition and Physical Activity for Children and Adolescents

#### 3.16.1 Detail

| Description: | Percentage of patients 3-17 years of age who had an outpatient visit with a PCP or Obstetrician/Gynecologist (OB/GYN) and who had evidence of the following during the measurement period. Three rates are reported.  
• Percentage of patients with height, weight, and BMI percentile documentation  
• Percentage of patients with counseling for nutrition  
• Percentage of patients with counseling for physical activity. |
| Stratification: | Report a total score, and each of the following strata:  
• Stratum 1 - Patients age 3-11.  
• Stratum 2 - Patients age 12-17. |
| Rationale: | Over the last three decades, childhood obesity has more than doubled in children and tripled in adolescents. Approximately 15 percent of children and adolescents in the United States are overweight and 17 percent are obese (Fryar, Carroll, & Ogden, 2014). Childhood obesity has both immediate and long-term effects on health and well-being. Children who are obese are more likely to be obese as adults and are therefore at risk for adult health problems, such as heart disease, type 2 diabetes, stroke and several types of cancer (Centers for Disease Control and Prevention, 2013. The direct medical costs associated with childhood obesity total about $19,000 per child, contributing to the $14 billion spent on care related to childhood obesity in the United States (Finkelstein, Graham, & Malhotra, 2014). Since obesity can become a lifelong health issue, it is important to screen for obesity in children and adolescents, and to provide interventions that promote weight loss (U.S. Preventive Services Task Force, 2017). |
**Clinical Recommendation Statement:**

U.S. Preventive Services Task Force (2017) - The task force recommends that clinicians screen for obesity in children and adolescents 6 years and older and offer or refer them to comprehensive, intensive behavioral interventions to promote improvements in weight status. (B recommendation) American Academy of Pediatrics - Bright Futures (Hagan, Shaw, & Duncan, 2017) - - Plot and assess BMI percentiles routinely for early recognition of overweight and obesity. - Assess barriers to healthy eating and physical activity. - Provide anticipatory guidance for nutrition and physical activity.

**Guidance:**

The visit must be performed by a PCP or OB/GYN. Because BMI norms for youth vary with age and sex, this measure evaluates whether BMI percentile, rather than an absolute BMI value, is assessed.

**Initial Population:**

Patients 3-17 years of age with at least one outpatient visit with a PCP or an OB/GYN during the measurement period.

<table>
<thead>
<tr>
<th>Denominator</th>
<th>Equals Initial Population.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator Exclusions</td>
<td>Patients who have a diagnosis of pregnancy during the measurement period. Exclude patients whose hospice care overlaps the measurement period.</td>
</tr>
</tbody>
</table>

**Numerator:**

Numerator 1: Patients who had a height, weight, and BMI percentile recorded during the measurement period.
Numerator 2: Patients who had counseling for nutrition during the measurement period.
Numerator 3: Patients who had counseling for physical activity during the measurement period.

<table>
<thead>
<tr>
<th>Numerator Exclusions</th>
<th>Not Applicable.</th>
</tr>
</thead>
</table>

**Denominator Exceptions:**

None.

**Supplemental Data Elements:**

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.16.2 Data Entry

**Description:**

Percentage of patients 3-17 years of age who had an outpatient visit with a PCP or OB/GYN and who had evidence of the following during the measurement period. Three rates are reported:

- Percentage of patients with height, weight, and BMI percentile documentation.
- Percentage of patients with counselling for nutrition.
- Percentage of patients with counselling for physical activity.
Three rates are reported:

- **Multiple encounter codes accepted:**
  - Preventive Care Services-Individual Counseling (Value Set 2.16.840.1.113883.3.464.1003.101.12.1026).
  - Preventive Care Services - Group Counseling (Value Set 2.16.840.1.113883.3.464.1003.101.12.1027).
  - Preventive Care (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1024, ….1022).
  - Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).

- **Denominator (same for all three):**
  - Visit or admission, child age 3-17 at start of measurement period.
  - Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Palliative Care](image)

**Figure 3-24: Palliative Care list**

  - Exclude patients who are pregnant (Value Set: 2.16.840.1.113883.3.526.3.378).

- **Numerator 1:**
  - Height, weight, BMI all recorded during measurement period.
  - Stratification 1 – age 3 to less than 11 at start of measurement period.
  - Stratification 2 – age 11 to less than 17 at start of measurement period.

- **Numerator 2:**
  - TREG edit for POV (education for both diet and exercise).
Stratification 1 – age 3 to less than 11 at start of measurement period.
Stratification 2 – age 11 to less than 17 at start of measurement period.

Numerator 3:
- Counseling for Physical Activity begins during measurement period (Value Set: 2.16.840.1.113883.3.464.1003.118.12.1035).
- TREG as above.
- Stratification 1 – age 3 to less than 11 at start of measurement period.
- Stratification 2 – age 11 to less than 17 at start of measurement period.

3.17 CMS 156V8 Use of High-Risk Medications in the Elderly

3.17.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients 65 years of age and older who were ordered high-risk medications. Two rates are reported.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Percentage of patients who were ordered at least one high-risk medication.</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients who were ordered at least two of the same high-risk medications.</td>
</tr>
<tr>
<td><strong>Rationale:</strong></td>
<td>Certain medications (MacKinnon &amp; Hepler, 2003) are associated with increased risk of harm from drug side-effects and drug toxicity and pose a concern for patient safety. There is clinical consensus that these drugs pose increased risks in the elderly (Kaufman, Brodin, &amp; Sarafian, 2005). Potentially Inappropriate Medication use in older adults has been connected to significantly longer hospital stay lengths and increased hospitalization costs (Hagstrom et al., 2015) as well as increased risk of death (Lau et al. 2004). Older adults receiving inappropriate medications are more likely to report poorer health status at follow-up, compared to those who receive appropriate medications (Fu, Liu, &amp; Christensen, 2004). A study of the prevalence of potentially inappropriate medication use in older adults found that 40 percent of individuals 65 and older filled at least one prescription for a potentially inappropriate medication and 13 percent filled two or more (Fick et al., 2008). While some adverse drug events are not preventable, studies estimate that between 30 and 80 percent of adverse drug events in the elderly are preventable (MacKinnon &amp; Hepler, 2003). Reducing the number of inappropriate prescriptions can lead to improved patient safety and significant cost savings. Conservative estimates of extra costs due to potentially inappropriate medications in the elderly average $7.2 billion a year (Fu et al., 2007). Medication use by older adults will likely increase further as the U.S. population ages, new drugs are developed, and new therapeutic and preventive uses for medications are discovered (Rothberg et al., 2008). The annual direct costs of preventable adverse drug events (ADEs) in the Medicare population have been estimated to exceed $800 million (Institute of Medicine, 2007). By the year 2030, nearly one in five U.S. residents is expected to be aged 65 years or older; this age group is projected to more than double in number from 38.7 million in 2008 to more than 88.5 million in 2050. Likewise, the population aged 85 years or older is expected to increase almost four-fold, from 5.4 million to 19 million between 2008 and 2050. As the elderly population continues to grow, the number of older adults who present with multiple medical conditions for which several medications are prescribed continues to increase, resulting in polypharmacy (Gray &amp; Gardner, 2009).</td>
</tr>
<tr>
<td><strong>Clinical Recommendation Statement:</strong></td>
<td>The measure is based on recommendations from the American Geriatrics Society Beers Criteria for Potentially Inappropriate Medication Use in Older Adults. The criteria were developed through key clinical expert consensus processes by Beers in 1997, Zhan in 2001 and an updated process by Fick et al. in 2003, 2012 and 2015. The Beers Criteria identifies lists of drugs that are potentially inappropriate for all older adults and drugs that are potentially inappropriate in the elderly based on various high-risk factors such as dosage, days supply and underlying diseases or conditions. NCQA’s Geriatric Measurement Advisory Panel selected a subset of drugs that should be used with caution in the elderly for inclusion in the proposed measure based upon the recommendations in the Beers Criteria.</td>
</tr>
</tbody>
</table>
**Definition:** A high-risk medication is identified by either of the following:
- A prescription for medications classified as high risk at any dose and for any duration.
- Prescriptions for medications classified as high risk at any dose with greater than a 90-day supply.

**Guidance:** The intent of Numerator 1 of the measure is to assess if the patient has been prescribed at least one high-risk medication. The intent of Numerator 2 of the measure is to assess if the patient has been prescribed at least two of the same high-risk medications on different days. The intent of the measure is to assess if the reporting provider ordered the high-risk medication(s). If the patient had a high-risk medication previously prescribed by another provider, they would not be counted towards the numerator unless the reporting provider also ordered a high-risk medication for them.

**Initial Population:** Patients 65 years and older who had a visit during the measurement period.

**Denominator:** Equals Initial Population.

**Denominator Exclusions:** Exclude patients whose hospice care overlaps the measurement period.

**Numerator:**

- **Numerator 1:** Patients with an order for at least one high-risk medication during the measurement period
- **Numerator 2:** Patients with at least two orders for the same high-risk medication on different days during the measurement period

**Numerator Exclusions:** Not Applicable.

**Denominator Exceptions:** None.

**Supplemental Data Elements:** For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.17.2 Data Entry

**Description:** Percentage of patients 65 years of age and older who were ordered high-risk medications. Two rates are reported.
- Percentage of patients who were ordered at least one high-risk medication.
- Percentage of patients who were ordered at least two of the same high-risk medications.

Visit or admission, patient age 65 or older at start of measurement period. Multiple encounter codes accepted:
- Inpatient (Value Set 2.16.840.1.113883.3.666.5.307).

• Discharge Services - Nursing Facility (Value Set 2.16.840.1.113883.3.464.1003.101.12.1013).

• Care Services in Long-Term Residential Facility (Value Set 2.16.840.1.113883.3.464.1003.101.12.1014).

• Annual Wellness Visit (Value Set 2.16.840.1.113883.3.526.3.1240).

• Nursing Facility Visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1012).

• Ophthalmologic Outpatient Visit (Value Set 2.16.840.1.113883.3.464.1003.101.11.1206).

• Preventive Care Services (Value Sets 2.16.840.1.113883.3.464.1003.101.12.1023, …, 1025).

See Appendix A for information on using Value Sets.

• Excludes patients on hospice, Comfort measures, or discharged to hospice (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Figure 3-26: Palliative Care list](image)

Numerator 1: To see if ANY high-risk meds prescribed:
  - Prescribed any medication on high risk medications for the elderly list (See value sets).
  - Given prescription for any of these meds with over 90-day supply.

Numerator 2: (any duplicate prescriptions, or two meds in same class with more than 90-day supply). Any of the following:
  - Two prescriptions for meds listed “list of single RX NORM code concepts” for High Risk drugs for the Elderly (value sets) during measurement period.
  - Sum of prescribed days >90 for 2 prescriptions for Nonbenzodiazepine hypnotics plus Nonbenzodiazepine hypnotics during measurement period.
  - Sum of prescribed days >90 for two prescriptions for “anti-infectives, other”, during measurement period. (See value sets for list of medications).
3.18 CMS 159V8 Depression Remission at Twelve Months

3.18.1 Detail

| Description: | The percentage of adolescent patients 12 to 17 years of age and adult patients 18 years of age or older with major depression or dysthymia who reached remission 12 months (+/- 60 days) after an index event. |
| Stratification: | Ages 12 to 17  
Ages 18 and older |
**Rationale:**

Adults: Depression is a common and treatable mental disorder. 8.1% of American adults age 20 and over had depression in a given 2 week period. Women (10.4%) were almost twice as likely as men (5.5%) to have had depression. The prevalence of depression among adults decreased as family income levels increased. About 80% of adults with depression reported at least some difficulty with work, home, or social activities because of their depression symptoms (Centers for Disease Control and Prevention, 2018). Persons with a current diagnosis of depression and a lifetime diagnosis of depression or anxiety were significantly more likely than persons without these conditions to have cardiovascular disease, diabetes, asthma, and obesity and to be a current smoker, to be physically inactive and to drink heavily (Strine et al., 2008). People who suffer from depression have lower incomes, lower educational attainment and fewer days working each year, leading to seven fewer weeks of work per year, a loss of 20% in potential income and a lifetime loss for each family who has a depressed family member of $300,000 (Smith & Smith, 2010). The incremental economic burden of individuals with major depressive disorder (MDD) increased by 21.5% (from $173.2 billion to $210.5 billion, inflation-adjusted dollars). The composition of these costs remained stable, with approximately 45% attributable to direct costs, 5% to suicide-related costs, and 50% to workplace costs. Only 38% of the total costs were due to MDD itself as opposed to comorbid conditions. (Greenberg, 2015). Adolescents and Adults: The Centers for Disease Control and Prevention states that during 2009-2012 an estimated 7.6% of the U.S. population aged 12 and over had depression, including 3% of Americans with severe depressive symptoms. Almost 43% of persons with severe depressive symptoms reported serious difficulties in work, home, and social activities, yet only 35% reported having contact with a mental health professional in the past year. Depression is associated with higher mortality rates in all age groups. People who are depressed are 30 times more likely to take their own lives than people who are not depressed and five times more likely to abuse drugs. Depression is the leading cause of medical disability for people aged 14 to 44. Depressed people lose 5.6 hours of productive work every week when they are depressed, fifty percent of which is due to absenteeism and short-term disability. Adolescents: In 2014, an estimated 2.8 million adolescents age 12 to 17 in the United States had at least one major depressive episode (MDE) in the past year. This represented 11.4% of the U.S. population. The same survey found that only 41.2 percent of those who had a MDE received treatment in the past year. The 2013 Youth Risk Behavior Survey of students grades 9 to 12 indicated that during the past 12 months 39.1% (F) and 20.8% (M) indicated feeling sad or hopeless almost every day for at least 2 weeks, planned suicide attempt 16.9% (F) and 10.3% (M), with attempted suicide 10.6% (F) and 5.4% (M). Adolescent-onset depression is associated with chronic depression in adulthood.
Many mental health conditions (anxiety, bipolar, depression, eating disorders, and substance abuse) are evident by age 14. The 12-month prevalence of MDEs increased from 8.7% in 2005 to 11.3% in 2014 in adolescents and from 8.8% to 9.6% in young adults (both P < .001). The increase was larger and statistically significant only in the age range of 12 to 20 years. The trends remained significant after adjustment for substance use disorders and sociodemographic factors. Mental health care contacts overall did not change over time; however, the use of specialty mental health providers increased in adolescents and young adults, and the use of prescription medications and inpatient hospitalizations increased in adolescents. In 2015, 9.7% of adolescents in MN who were screened for depression or other mental health conditions, screened positively.

**Clinical Recommendation Statement:**

Adults: Source: Institute for Clinical Systems Improvement (ICSI) Health Care Guideline for Adult Depression in Primary Care (Trangle et al., 2016) Major depression is a treatable cause of pain, suffering, disability and death, yet primary care clinicians detect major depression in only one-third to one-half of their patients with major depression (Williams et al. 2002; Schonfeld et al., 1997). Usual care for depression in the primary care setting has resulted in only about half of depressed adults getting treated (Kessler et al., 2005) and only 20-40% showing substantial improvement over 12 months (Unutzer et al., 2002; Katon et al., 1999). Recommendations and algorithm notations supporting depression outcomes and duration of treatment according to ICSI's Health Care Guideline: Recommendation: Clinicians should establish and maintain follow-up with patients. Appropriate, reliable follow-up is highly correlated with improved response and remission scores. It is also correlated with the improved safety and efficacy of medications and helps prevent relapse. Proactive follow-up contacts (in person, telephone) based on the collaborative care model have been shown to significantly lower depression severity (Unutzer et al., 2002). In the available clinical effectiveness trials conducted in real clinical practice settings, even the addition of a care manager leads to modest remission rates (Trivedi et al., 2006; Unutzer et al., 2002). Interventions are critical to educating the patient regarding the importance of preventing relapse, safety and efficacy of medications, and management of potential side effects. Establish and maintain initial follow-up contact intervals (office, phone, other) (Hunkeler et al., 2000; Simon et al., 2000). PHQ-9 as monitor and management tool. The PHQ-9 is an effective management tool, as well, and should be used routinely for subsequent visits to monitor treatment outcomes and severity. It can also help the clinician decide if/how to modify the treatment plan (Duffy et al., 2008; Lowe et al., 2004).
Using a measurement-based approach to depression care, PHQ-9 results and side effect evaluation should be combined with treatment algorithms to drive patients toward remission. A five-point drop in PHQ-9 score is considered the minimal clinically significant difference (Trivedi, 2009). Every time that the PHQ-9 is assessed, suicidality is assessed, as well. If the suicidality was indeed of high risk, urgent referral to crisis specialty health care is advised. In case of low suicide risk, the patient can proceed with treatment in the primary care practice (Huijbregts et al., 2013). Care Algorithm: Has patient reached remission? The goals of treatment should be to achieve remission, reduce relapse and recurrence, and return to previous level of occupational and psychosocial function. Full remission is defined as a 2-month period devoid of major depressive signs and symptoms (American Psychiatric Association, 2013). If using a PHQ-9 tool, remission translates to PHQ-9 score of less than 5 (Kroenke, 2001). Results from the STAR*D study showed that remission rates lowered with more treatment steps, but the overall cumulative rate was 67% (Rush et al., 2006). Response is defined as a 50% or greater reduction in symptoms (as measured on a standardized rating scale). Partial response is defined as a 25-50% reduction in symptoms. This definition is based on how the depression literature defines response. Response and remission take time. In the STAR*D study, longer times than expected were needed to reach response or remission. In fact, one-third of those who ultimately responded did so after six weeks. Of those who achieved remission by Quick Inventory of Depressive Symptomatology (QIDS), 50% did so only at or after six weeks of treatment (Trivedi et al., 2006). If the primary care clinician is seeing some improvement, continue working with that patient to augment or increase dosage to reach remission. This can take up to 3 months. A reasonable criterion for extending the initial treatment: assess whether the patient is experiencing a 25% or greater reduction in baseline symptom severity at six weeks of therapeutic dose. If the patient's symptoms are reduced by 25% or more, but the patient is not yet at remission, and if medication has been well tolerated, continue to prescribe. Raising the dose is recommended (Trivedi et al., 2006). Improvement with psychotherapy is often a bit slower than with pharmacotherapy. A decision regarding progress with psychotherapy and the need to change or augment this type of treatment may require 8 to 10 weeks before evaluation (Schulberg et al., 1998). Care Algorithm: Continuation and Maintenance Treatment Duration Based on Episode Acute therapy is the treatment phase focused on treating the patient to remission. Acute therapy typically lasts 6-12 weeks but technically lasts until remission is reached (American Psychiatric Association, 2010). Full remission is defined as a 2-month period devoid of major depressive signs and symptoms (American Psychiatric Association, 2013). Continuation therapy is the 4-to-9 month period beyond the acute treatment phase during which the patient is treated with antidepressants, psychotherapy, ECT or other somatic therapies to prevent relapse (American Psychiatric Association, 2010).
<table>
<thead>
<tr>
<th>Clinical Recommendation Statement: (cont)</th>
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<tbody>
<tr>
<td>Relapse is common within the first 6 months following remission from an acute depressive episode; as many as 20-85% of patients may relapse (American Psychiatric Association, 2010). This measure assesses achievement of remission, which is a desired outcome of effective depression treatment and monitoring. Adult Depression in Primary Care - Guideline Aims - Increase the percentage of patients with major depression or persistent depressive disorder who have improvement in outcomes from treatment for major depression or persistent depressive disorder. - Increase the percentage of patients with major depression or persistent depressive disorder who have follow-up to assess for outcomes from treatment. - Improve communication between the primary care physician and the mental health care clinician (if patient is co-managed). Adolescents: Source: American Academy of Child and Adolescent Psychiatry Practice Parameter for the Assessment and Treatment of Children and Adolescents with Depressive Disorders (2007) <a href="http://www.jaacap.com/article/S0890-8567(09)62053-0/pdf">http://www.jaacap.com/article/S0890-8567(09)62053-0/pdf</a> Recommendations: Recommendations supporting depression outcomes and duration of treatment according to AACAP guideline: - Treatment of depressive disorders should always include an acute and continuation phase; some children may also require maintenance treatment. The main goal of the acute phase is to achieve response and ultimately full symptomatic remission (definitions below). - Each phase of treatment should include psychoeducation, supportive management, and family and school involvement. - Education, support, and case management appear to be sufficient treatment for the management of depressed children and adolescents with an uncomplicated or brief depression or with mild psychosocial impairment. - For children and adolescents who do not respond to supportive psychotherapy or who have more complicated depressions, a trial with specific types of psychotherapy and/or antidepressants is indicated. Sources: Guidelines for Adolescent Depression in Primary Care (GLAD-PC) (2018) <a href="http://pediatrics.aappublications.org/content/141/3/e20174081">http://pediatrics.aappublications.org/content/141/3/e20174081</a> Guidelines for adolescent depression in primary care (GLAD-PC): II. Treatment and ongoing management <a href="http://pediatrics.aappublications.org/content/141/3/e20174082">http://pediatrics.aappublications.org/content/141/3/e20174082</a> Recommendations supporting depression outcomes and duration of treatment according to GLAD-PC: Recommendations for Ongoing Management of Depression: - Mild depression: consider a period of active support and monitoring before starting other evidence based treatment - Moderate or severe major clinical depression or complicating factors: -- consultation with mental health specialist with agreed upon roles -- evidence based treatment (CBT or IPT and/or antidepressant SSRI) - Monitor for adverse effects during antidepressant therapy -- clinical worsening, suicidality, unusual changes in behavior - Systematic and regular tracking of goals and outcomes -- improvement in functioning status and resolution of depressive symptoms Regardless of the length of treatment, all patients should be monitored on a monthly basis for 6 to 12 months after the full resolution of symptoms</td>
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</table>
**Definition:**

Denominator Identification Period: The period in which eligible patients can have an index event. The denominator identification period occurs prior to the measurement period and is defined as 14 months to 2 months prior to the start of the measurement period. For patients with an index event, there needs to be enough time following index for the patients to have the opportunity to reach remission 12 months +/- 60 days after the index event date. Index Event Date: The date in which the first instance of elevated PHQ-9 or PHQ-9M greater than nine and diagnosis of depression or dysthymia occurs during the denominator identification measurement period. Patients may be screened using PHQ-9 and PHQ-9M up to 7 days prior to the office visit (including the day of the office visit). Measure Assessment Period: The index event date marks the start of the measurement assessment period for each patient which is 14 months (12 months +/- 60 days) in length to allow for a follow-up PHQ-9 or PHQ-9M between 10 and 14 months following the index event. This assessment period is fixed and does not start over with a higher PHQ-9 or PHQ-9M that may occur after the index event date. Remission is defined as a PHQ-9 or PHQ-9M score of less than five. 12 months is defined as the point in time from the index event date extending out 12 months and then allowing a grace period of 60 days prior to and 60 days after this date. The most recent PHQ-9 or PHQ-9M score less than five obtained during this 4 month period is deemed as remission at 12 months, values obtained prior to or after this period are not counted as numerator compliant (remission).

**Guidance**

None

**Initial Population:**

Adolescent patients 12 to 17 years of age and adult patients 18 years of age and older with a diagnosis of major depression or dysthymia and an initial PHQ-9 or PHQ-9M score greater than nine during the index event. Patients may be screened using PHQ-9 and PHQ-9M up to 7 days prior to the office visit (including the day of the office visit).

**Denominator:**

Equals Initial Population

**Denominator Exclusions:**

Patients who died.
Patients who received hospice or palliative care services.
Patients who were permanent nursing home residents
Patients with a diagnosis of bipolar disorder
Patients with a diagnosis of personality disorder
Patients with a diagnosis of schizophrenia or psychotic disorder
Patients with a diagnosis of pervasive developmental disorder

**Numerator:**

Adolescent patients 12 to 17 years of age and adult patients 18 years of age and older who achieved remission at 12 months as demonstrated by a 12 month ( +/- 60 days) PHQ-9 or PHQ-9M score of less than five

**Numerator Exclusions:**

Not Applicable
Denominator Exceptions: None

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

3.18.2 Data Entry

Description: The percentage of adolescent patients 12 to 17 years of age and adult patients 18 years of age or older with major depression or dysthymia who reached remission 12 months (+/- 60 days) after an index event.

Index event (when PHQ-9 elevated), patient age 12 or more.

PHQ-9 >9 (Enter in Vital Signs).

Index event occurs between 14 months before start of measurement period and 2 months before start of measurement period (to allow time for 12 month follow up).

Diagnosis, wither of the following:
- “Depression” (Value Set: 2.16.840.113883.3.67.1.101.3.2444).
- “Dysthymia” (Value Set: 2.16.840.1.113883.3.67.1.101.1.254).

Denominator Exclusions:
- Palliative care order/encounter before interval (index event + 10-14 months).
- Care services in Long-term residential facility before same interval (Value Set: 2.16.840.1.113883.3.464.1003.101.12.1014).
- Patient expires before same interval.
- “Disorder diagnosis” before same interval.
  - Bipolar disorder (Value Set: 2.16.840.1.113883.3.67.1.101.1.128).
  - Personality disorder (Value Set: 2.16.840.1.113883.3.67.1.101.1.246).
  - Pervasive developmental disorder (Value Set: 2.16.840.1.113883.3.464.1003.105.12.1152).

Numerator:
PHQ-9 <5 score in same interval as above (12 months plus or minus 60 days).
- Stratification 1: Age 12-17 at index encounter with diagnosis depression.
- Stratification 2: Age 18+.
3.19 CMS 160V7 Depression Utilization of the PHQ-9 Tool

3.19.1 Detail

| Description: | The percentage of adolescent patients 12 to 17 years of age and adult patients age 18 and older with the diagnosis of major depression or dysthymia who have a completed PHQ-9 during each applicable 4-month period in which there was a qualifying depression encounter. |
| Stratification: | • Ages 12 to 17.  
• Ages 18 and older. |
| Rationale: | Adults:  
Depression is a common and treatable mental disorder. The Centers for Disease Control and Prevention states that an estimated 6.6% of the U.S. adult population (14.8 million people) experiences a major depressive disorder during any given 12-month period. Additionally, dysthymia accounts for an additional 3.3 million Americans. In 2006 and 2008, an estimated 9.1% of U.S. adults reported symptoms for current depression (Centers for Disease Control and Prevention, 2010). Persons with a current diagnosis of depression and a lifetime diagnosis of depression or anxiety were significantly more likely than persons without these conditions to have cardiovascular disease, diabetes, asthma, and obesity and to be a current smoker, to be physically inactive and to drink heavily (Strine, 2008). People who suffer from depression have lower incomes, lower educational attainment and fewer days working each year, leading to seven fewer weeks of work per year, a loss of 20% in potential income and a lifetime loss for each family who has a depressed family member of $300,000 (Smith, 2010). The cost of depression (lost productivity and increased medical expense) in the United States is $83 billion each year (Greenberg, 2003).  
Adolescents and Adults:  
The Centers for Disease Control and Prevention states that during 2009-2012 an estimated 7.6% of the U.S. population aged 12 and over had depression, including 3% of Americans with severe depressive symptoms. Almost 43% of persons with severe depressive symptoms reported serious difficulties in work, home and social activities, yet only 35% reported having contact with a mental health professional in the past year. Depression is associated with higher mortality rates in all age groups. People who are depressed are 30 times more likely to take their own lives than people who are not depressed and five times more likely to abuse drugs. Depression is the leading cause of medical disability for people aged 14 - 44. Depressed people lose 5.6 hours of productive work every week when they are depressed, fifty percent of which is due to absenteeism and short-term disability. |
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<th>Rationale (cont.):</th>
<th>Adolescents:</th>
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<tr>
<td></td>
<td>In 2014, an estimated 2.8 million adolescents age 12 to 17 in the United States had at least one major depressive episode in the past year. This represented 11.4% of the U.S. population. The same survey found that only 41.2 percent of those who had a Major Depressive Episode received treatment in the past year. The 2013 Youth Risk Behavior Survey of students grades 9 to 12 indicated that during the past 12 months 39.1% (F) and 20.8% (M) indicated feeling sad or hopeless almost every day for at least 2 weeks, planned suicide attempt 16.9% (F) and 10.3% (M), with attempted suicide 10.6% (F) and 5.4% (M). Adolescent-onset depression is associated with chronic depression in adulthood. Many mental health conditions (anxiety, bipolar, depression, eating disorders, and substance abuse) are evident by age 14. The 12-month prevalence of MDEs increased from 8.7% in 2005 to 11.3% in 2014 in adolescents and from 8.8% to 9.6% in young adults (both P &lt; .001). The increase was larger and statistically significant only in the age range of 12 to 20 years. The trends remained significant after adjustment for substance use disorders and sociodemographic factors. Mental health care contacts overall did not change over time; however, the use of specialty mental health providers increased in adolescents and young adults, and the use of prescription medications and inpatient hospitalizations increased in adolescents. In 2015, 9.7% of adolescents in MN who were screened for depression or other mental health conditions, screened positively. Please note that this process measure for administration of the PHQ-9 or PHQ-9M depression tool, a PROM that is validated for both the assessment and diagnosis of depression as well as for monitoring ongoing outcomes of treatment, is a PAIRED process measure with RELATED measures of depression remission (PHQ-9/PHQ-9M &lt; 5) and depression response (PHQ-9/PHQ-9M is improved by &gt; 50%) at 6 and 12 months. To quote a NQF Behavioral Steering Committee member as these measures were initially endorsed &quot;the best way to avoid being measured is to never give the PHQ-9&quot;. This process measure allows an understanding of the use of the tool in the target population, promotes frequent and follow-up contact with patients whose score indicates a need for treatment and serves as a catalyst in a collaborative care model for patients with major depression or dysthymia. It is estimated that up to 90% of patients diagnosed with depression and anxiety are treated solely in primary care (National Institute for Health and Care Excellence, 2011).</td>
</tr>
</tbody>
</table>
**Clinical Recommendation Statement:**

Adults:

Source: ICSI Health Care Guideline for Adult Depression in Primary Care (Trangle, 2016).

Major depression is a treatable cause of pain, suffering, disability and death, yet primary care clinicians detect major depression in only one-third to one-half of their patients with major depression (Williams Jr, 2002; Schonfeld, 1997).

Usual care for depression in the primary care setting has resulted in only about half of depressed adults getting treated (Kessler, 2005) and only 20-40% showing substantial improvement over 12 months (Unutzer, 2002; Katon, 1999).

Recommendations and algorithm notations supporting depression assessment and monitoring of depression outcomes according to ICSI's Health Care Guideline:

Recommendation: Clinicians should establish and maintain follow-up with patients. Appropriate, reliable follow-up is highly correlated with improved response and remission scores. It is also correlated with the improved safety and efficacy of medications and helps prevent relapse.

Proactive follow-up contacts (in person, telephone) based on the collaborative care model have been shown to significantly lower depression severity (Unutzer, 2002). In the available clinical effectiveness trials conducted in real clinical practice settings, even the addition of a care manager leads to modest remission rates (Trivedi, 2006; Unutzer, 2002). Interventions are critical to educating the patient regarding the importance of preventing relapse, safety and efficacy of medications, and management of potential side effects. Establish and maintain initial follow-up contact intervals (office, phone, other) (Hunkeler, 2000; Simon, 2000). PHQ-9 as monitor and management tool. The PHQ-9 is an effective management tool, as well, and should be used routinely for subsequent visits to monitor treatment outcomes and severity. It can also help the clinician decide if/how to modify the treatment plan (Duffy, 2008; Lowe, 2004). Using a measurement-based approach to depression care, PHQ-9 results and side effect evaluation should be combined with treatment algorithms to drive patients toward remission. A five-point drop in PHQ-9 score is considered the minimal clinically significant difference (Trivedi, 2009). Every time that the PHQ-9 is assessed, suicidality is assessed, as well. If the suicidality was indeed of high risk, urgent referral to crisis specialty health care is advised. In case of low suicide risk, the patient can proceed with treatment in the primary care practice (Huijbregts, 2013). Adult Depression in Primary Care - Guideline Aims

Increase the percentage of patients with major depression or persistent depressive disorder who have improvement in outcomes from treatment for major depression or persistent depressive disorder. Increase the percentage of patients with major depression or persistent depressive disorder who have follow-up to assess for outcomes from treatment. Improve communication between the primary care physician and the mental health care clinician (if patient is co-managed).
<table>
<thead>
<tr>
<th>Clinical Recommendation Statement (cont.):</th>
<th>Adolescents:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommendations supporting depression outcomes and duration of treatment according to AACAP guideline:</td>
<td></td>
</tr>
<tr>
<td>Treatment of depressive disorders should always include an acute and continuation phase; some children may also require maintenance treatment. The main goal of the acute phase is to achieve response and ultimately full symptomatic remission (definitions below). Each phase of treatment should include psychoeducation, supportive management, and family and school involvement</td>
<td></td>
</tr>
<tr>
<td>Education, support, and case management appear to be sufficient treatment for the management of depressed children and adolescents with an uncomplicated or brief depression or with mild psychosocial impairment</td>
<td></td>
</tr>
<tr>
<td>For children and adolescents who do not respond to supportive psychotherapy or who have more complicated depressions, a trial with specific types of psychotherapy and/or antidepressants is indicated</td>
<td></td>
</tr>
<tr>
<td>Definitions:</td>
<td></td>
</tr>
<tr>
<td>Response: No symptoms or a significant reduction in depressive symptoms for at least 2 weeks</td>
<td></td>
</tr>
<tr>
<td>Remission: A period of at least 2 weeks and &lt;2months with no or few depressive symptoms</td>
<td></td>
</tr>
<tr>
<td>Recovery: Absence of significant symptoms of depression (e.g., no more than 1 to 2 symptoms) for greater than 2 months</td>
<td></td>
</tr>
<tr>
<td>Relapse: A DSM episode of depression during the period of remission</td>
<td></td>
</tr>
<tr>
<td>Recurrence: The emergence of symptoms of depression during the period of recovery (a new episode)</td>
<td></td>
</tr>
<tr>
<td>Sources:</td>
<td></td>
</tr>
<tr>
<td>Guidelines for Adolescent Depression in Primary Care (GLAD-PC) (2018) <a href="http://pediatrics.aappublications.org/content/141/3/e20174081">http://pediatrics.aappublications.org/content/141/3/e20174081</a> Guidelines for adolescent depression in primary care (GLAD-PC): II. Treatment and ongoing management <a href="http://pediatrics.aappublications.org/content/141/3/e20174082">http://pediatrics.aappublications.org/content/141/3/e20174082</a> Recommendations supporting depression outcomes and duration of treatment according to GLAD-PC:</td>
<td></td>
</tr>
</tbody>
</table>
Clinical Recommendation Statement (cont.): Recommendations for Ongoing Management of Depression:
- Mild depression: consider a period of active support and monitoring before starting other evidence-based treatment
- Moderate or severe major clinical depression or complicating factors:
  - Consultation with mental health specialist with agreed upon roles.
  - Evidence based treatment (CBT or IPT and/or antidepressant SSRI)
  - Monitor for adverse effects during antidepressant therapy -- clinical worsening, suicidality, unusual changes in behavior
  - Systematic and regular tracking of goals and outcomes -- improvement in functioning status and resolution of depressive symptoms.
- Regardless of the length of treatment, all patients should be monitored on a monthly basis for 6 to 12 months after the full resolution of symptoms.

Definition: Completed PHQ-9 or PHQ-9M - The patient must answer ALL nine questions for the score to be valid.

Guidance: If a patient has a qualifying diagnosis and encounter in more than one of the 4-month periods within the measurement year, the patient must be counted (denominator and numerator) in each qualifying 4-month period. For example, a patient could be counted in the first and third 4-month periods.

Initial Population: Adolescent patients 12 to 17 years of age and adult patients 18 years of age and older with an office visit and the diagnosis of major depression or dysthymia during the 4-month period.

Denominator: Equals Initial Population.

Denominator Exclusions: Patients who died.
- Patients who received hospice or palliative care services.
- Patients who were permanent nursing home residents.
- Patients with a diagnosis of bipolar disorder.
- Patients with a diagnosis of personality disorder.
- Patients with a diagnosis of schizophrenia or psychotic disorder.
- Patients with a diagnosis of pervasive developmental disorder.

Numerator: Adolescent patients 12 to 17 years of age and adult patients 18 years of age and older who have a PHQ-9 or PHQ-9M tool administered at least once during the 4-month period.

Numerator Exclusions: Not applicable.

Denominator Exceptions: None.

Supplemental Data Elements: For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.
3.19.2 Data Entry

| Description: | The percentage of adolescent patients 12 to 17 years of age and adult patients age 18 and older with the diagnosis of major depression or dysthymia who have a completed PHQ-9 during each applicable 4-month period in which there was a qualifying depression encounter |

Three calculations for visits during measurement period in 4-month blocks:
- January –April.
- May-August.
- September-December.

For all three, visit age equal or >12 at time of encounter.
- Diagnosis = Depression (Value Set: 2.16.840.113883.3.67.1.101.3.2444).
- Dysthymia (Value Set: 2.16.840.1.113883.3.67.1.101.1.254).

Denominator exclusions:
- Palliative care (Value Set: 2.16.840.1.113883.3.600.1.1579).
- Patient expired.
- Pre-existing depression diagnosis at time of encounter.

Numerator:
- PHQ-9 done and reported during 4-month block (Enter in Vital Signs).
- Stratification 1: Age equal or > 12-<18 at time of encounter.
- Stratification 2: Age equal or > 18 at time of encounter.
- Includes telehealth encounters.

3.20 CMS 161V8 Adult Major Depressive Disorder: Suicide Risk Assessment

3.20.1 Detail

| Description: | Percentage of patients aged 18 years and older with a diagnosis of major depressive disorder (MDD) with a suicide risk assessment completed during the visit in which a new diagnosis or recurrent episode was identified. |
### Rationale:

This measure aims to improve rates of clinician assessment of suicide risk during an encounter where a new or recurrent episode of major depressive disorder is identified. In an epidemiologic study (2010) of mental illness in the United States with a large, representative sample, 69% of respondents with lifetime suicide attempts had also met diagnostic criteria for major depressive disorder. When considering other mood disorders related to depression, such as dysthymia and bipolar disorders, this rate increases to 74% (Bolton & Robinson, 2010). In a 2014 study conducted by Ahmedani et al., 50% of individuals who completed a suicide had been seen in a health care setting within four weeks prior. Better assessment and identification of suicide risk in the health care setting should lead to improved connection to treatment and reduction in suicide attempts and deaths by suicide.
Clinical Recommendation Statement: A careful and ongoing evaluation of suicide risk is necessary for all patients with major depressive disorder. Such an assessment includes specific inquiry about suicidal thoughts, intent, plans, means, and behaviors; identification of specific psychiatric symptoms (e.g., psychosis, severe anxiety, substance use) or general medical conditions that may increase the likelihood of acting on suicidal ideas; assessment of past and, particularly, recent suicidal behavior; delineation of current stressors and potential protective factors (e.g., positive reasons for living, strong social support); and identification of any family history of suicide or mental illness. As part of the assessment process, impulsivity and potential for risk to others should also be evaluated, including any history of violence or violent or homicidal ideas, plans, or intentions. The patient's risk of harm to him- or herself and to others should also be monitored as treatment proceeds. Admission generally indicated after a suicide attempt or aborted suicide attempt if: * Patient is psychotic * Attempt was violent, near-lethal, or premeditated * Precautions were taken to avoid rescue or discovery * Persistent plan and/or intent is present * Distress is increased or patient regrets surviving * Patient is male, older than age 45 years, especially with new onset of psychiatric illness or suicidal thinking * Patient has limited family and/or social support, including lack of stable living situation * Current impulsive behavior, severe agitation, poor judgment, or refusal of help is evident * Patient has change in mental status with a metabolic, toxic, infectious, or other etiology requiring further workup in a structured setting in the presence of suicidal ideation with: * Specific plan with high lethality * High suicidal intent Admission may be necessary after a suicide attempt or aborted suicide attempt, except in circumstances for which admission is generally indicated in the presence of suicidal ideation with: * Psychosis * Major psychiatric disorder * Past attempts, particularly if medically serious * Possibly contributing medical condition (e.g., acute neurological disorder, cancer, infection) * Lack of response to or inability to cooperate with partial hospital or outpatient treatment * Need for supervised setting for medication trial or ECT * Need for skilled observation, clinical tests, or diagnostic assessments that require a structured setting * Limited family and/or social support, including lack of stable living situation * Lack of an ongoing clinician-patient relationship or lack of access to timely outpatient follow-up.
<table>
<thead>
<tr>
<th>Clinical Recommendation Statement: (cont)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Evidence of putting one's affairs in order (e.g., giving away possessions, writing a will)] In the absence of suicide attempts or reported suicidal ideation/plan/intent but evidence from the psychiatric evaluation and/or history from others suggests a high level of suicide risk and a recent acute increase in risk Release from emergency department with follow-up recommendations may be possible After a suicide attempt or in the presence of suicidal ideation/plan when: * Suicidality is a reaction to precipitating events (e.g., exam failure, relationship difficulties), particularly if the patient's view of situation has changed since coming to emergency department * Plan/method and intent have low lethality * Patient has stable and supportive living situation * Patient is able to cooperate with recommendations for follow-up, with treater contacted, if possible, if patient is currently in treatment Outpatient treatment may be more beneficial than hospitalization Patient has chronic suicidal ideation and/or self-injury without prior medically serious attempts, if a safe and supportive living situation is available and outpatient psychiatric care is ongoing.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Definition:</th>
</tr>
</thead>
<tbody>
<tr>
<td>The specific type and magnitude of the suicide risk assessment is intended to be at the discretion of the individual clinician and should be specific to the needs of the patient. At a minimum, suicide risk assessment should evaluate: 1) Suicidal ideation 2) Patient's intent of initiating a suicide attempt AND, if either is present, 3) Patient plans for a suicide attempt 4) Whether the patient has means for completing suicide Low burden tools to track suicidal ideation and behavior such as the Columbia-Suicide Severity Rating Scale (C-SSRS) and the Suicide Assessment Five-Step Evaluation and Triage (SAFE-T) can also be used. Because no validated assessment tool or instrument fully meets the aforementioned requirements for the suicide risk assessment, individual tools or instruments have not been explicitly included in coding.</td>
</tr>
</tbody>
</table>
Guidance:
This measure is an episode-of-care measure and should be reported for each instance of a new or recurrent episode of major depressive disorder (MDD); every new or recurrent episode will count separately in the Initial Population. As the guidelines state, it is important to assess for additional factors which may increase or decrease suicide risk, such as presence of additional symptoms (e.g., psychosis, severe anxiety, hopelessness, severe chronic pain); presence of substance abuse, history and seriousness of previous attempts, particularly, recent suicidal behavior, current stressors and potential protective factors (e.g., positive reasons for living, strong social support), family history of suicide or mental illness or recent exposure to suicide, impulsivity and potential for risk to others, including history of violence or violent or homicidal ideas, plans, or intentions, and putting one's affairs in order (e.g., giving away possessions, writing a will). In addition, although the measure focuses on the initial visit, it is critical that suicide risk be monitored especially for the 90 days following the initial visit and throughout MDD treatment. It is expected that a suicide risk assessment will be completed at the visit during which a new diagnosis is made or at the visit during which a recurrent episode is first identified (i.e., at the initial evaluation). For the purposes of this measure, an episode of major depressive disorder (MDD) would be considered to be recurrent if a patient has not had an MDD-related encounter in the past 105 days. If there is a gap of 105 or more days between visits for major depressive disorder (MDD), that would imply a recurrent episode. The 105-day look-back period is an operational provision and not a clinical recommendation, or definition of relapse, remission, or recurrence. Suicide risk assessments completed via telehealth services can also meet numerator performance. Use of a standardized tool(s) or instrument(s) to assess suicide risk will meet numerator performance. Standardized tools can be mapped to the concept "Intervention, Performed"; "Suicide risk assessment (procedure)" included in the numerator logic below. The logic statement for the age requirement, as written, captures patients who turn 18 years old during the measurement period so that these patients are included in the measure, so long as the minimum criteria noted above is evaluated. To ensure all patients with major depressive disorder (MDD) are assessed for suicide risk, there are two clinical quality measures addressing suicide risk assessment; CMS 177 covers children and adolescents aged 6 through 17, and CMS 161 covers the adult population aged 18 years and older, as no individual suicide risk assessment tool or instrument would satisfy the requirements alone.

<table>
<thead>
<tr>
<th>Initial Population:</th>
<th>All patients aged 18 years and older with a diagnosis of MDD.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator:</td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td>Denominator Exclusions:</td>
<td>None.</td>
</tr>
<tr>
<td>Numerator:</td>
<td>Patients with a suicide risk assessment completed during the visit in which a new diagnosis or recurrent episode was identified.</td>
</tr>
<tr>
<td>-----------</td>
<td>---------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Numerator Exclusions:</td>
<td>Not Applicable.</td>
</tr>
<tr>
<td>Denominator Exceptions:</td>
<td>None.</td>
</tr>
<tr>
<td>Supplemental Data Elements:</td>
<td>For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.</td>
</tr>
<tr>
<td>Initial Population:</td>
<td>All patients aged 18 years and older with a diagnosis of MDD.</td>
</tr>
<tr>
<td>Denominator:</td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td>Denominator Exclusions:</td>
<td>None.</td>
</tr>
<tr>
<td>Numerator:</td>
<td>Patients with a suicide risk assessment completed during the visit in which a new diagnosis or recurrent episode was identified.</td>
</tr>
<tr>
<td>Numerator Exclusions:</td>
<td>Not Applicable.</td>
</tr>
<tr>
<td>Denominator Exceptions:</td>
<td>None.</td>
</tr>
<tr>
<td>Supplemental Data Elements:</td>
<td>For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.</td>
</tr>
</tbody>
</table>

### 3.20.2 Data Entry

**Description:** Percentage of patients aged 18 years and older with a diagnosis of MDD with a suicide risk assessment completed during the visit in which a new diagnosis or recurrent episode was identified.

Encounter, patient age 18 or older.

Multiple encounter codes accepted:
- Outpatient Consultation (Value Set 2.16.840.1.113883.3.464.1003.101.12.1008).
- Psych Visit - Diagnostic Evaluation (Value Set 2.16.840.1.113883.3.526.3.1492).
- Psych Visit – Psychotherapy (Value Set 2.16.840.1.113883.3.526.3.1496).
- Psychoanalysis (Value Set 2.16.840.1.113883.3.526.3.1141).
- Telehealth Services
2.16.840.1.113883.3.464.1003.101.12.1031

See Appendix A for information on using Value Sets.

New diagnosis of either of the following:

- Depression (Value Set: 2.16.840.1.113883.3.526.3.1491).
- Recurrent episode (defined as > 105 days before current diagnosis).

Numerator: Suicide Risk Assessment performed and recorded (Enter in Exams).

### 3.21 CMS 165V8 Controlling High Blood Pressure

#### 3.21.1 Detail

| Description | Percentage of patients 18-85 years of age who had a diagnosis of hypertension and whose blood pressure was adequately controlled (<140/90mmHg) during the measurement period. |
Rationale: High blood pressure (HBP), also known as hypertension, is when the pressure in blood vessels is higher than normal (Centers for Disease Control and Prevention [CDC], 2016). The causes of hypertension are multiple and multifaceted and can be based on genetic predisposition, environmental risk factors, being overweight and obese, sodium intake, potassium intake, physical activity, and alcohol use. High Blood Pressure is common, according to the National Health and Nutrition Examination Survey (NHANES), approximately 85.7 million adults >= 20 years of age had HBP (140/90 mm Hg) between 2011 to 2014 (Crim, 2012). Between 2011-2014 the prevalence of hypertension (>=140/90 mm Hg) among US adults 60 and older was approximately 67.2 percent (Benjamin et al., 2017). HBP, known as the "silent killer," increases risks of heart disease and stroke which are two of the leading causes of death in the U.S. (Yoon, Fryar, & Carroll, 2015). A person who has HBP is four times more likely to die from a stroke and three times more likely to die from heart disease (CDC, 2012). The National Vital Statistics Systems Center for Disease Control and Prevention reported that in 2014 there were approximately 73,300 deaths directly due to HBP and 410,624 deaths with any mention of HBP (CDC, 2014). Between 2004 and 2014 the number of deaths due to HBP rose by 34.1 percent (Benjamin et al., 2017). Managing and treating HBP would reduce cardiovascular disease mortality for males and females by 30.4 percent and 38.0 percent, respectively (Patel et al., 2015). The estimated annual average direct and indirect cost of HBP from 2012 to 2013 was $51.2 billion (Benjamin et al., 2017). Total direct costs of HBP is projected to increase to $200 billion by 2030 (Benjamin et al., 2017). A study on cost-effectiveness on treating hypertension found that controlling HBP in patients with cardiovascular disease and systolic blood pressures of >=160 mm Hg could be effective and cost-saving (Moran et al., 2015). Many studies have shown that controlling high blood pressure reduces cardiovascular events and mortality. The Systolic Blood Pressure Intervention Trial (SPRINT) investigated the impact of obtaining a SBP goal of <120 mm Hg compared to a SBP goal of <140 mm Hg among patients 50 and older with established cardiovascular disease and found that the patients with the former goal had reduced cardiovascular events and mortality (SPRINT Research Group et al., 2015). Controlling HBP will significantly reduce the risks of cardiovascular disease mortality and lead to better health outcomes like reduction of heart attacks, stroke, and kidney disease (James et al., 2014). Thus, the relationship between the measure (control of hypertension) and the long-term clinical outcomes listed is well established.
**Clinical Recommendation Statement:**

The U.S. Preventive Services Task Force (2015) recommends screening for high blood pressure in adults age 18 years and older. This is a grade A recommendation. American College of Cardiology/American Heart Association (2017) -For adults with confirmed hypertension and known CVD or 10-year ASCVD event risk of 10% or higher, a blood pressure target of less than 130/80 mmHg is recommended -For adults with confirmed hypertension, without additional markers of increased CVD risk, a blood pressure target of less than 130/80 mmHg may be reasonable (Note: clinical trial evidence is strongest for a target blood pressure of 140/90 mmHg in this population. However observational studies suggest that these individuals often have a high lifetime risk and would benefit from blood pressure control earlier in life) American College of Physicians and the American Academy of Family Physicians (2017): -Initiate intensifying pharmacologic treatment in adults aged 60 and older at high cardiovascular risk, based on individualized assessment, to achieve a target systolic blood pressure of less than 140 mmHg (Grade: weak recommendation, quality of evidence: low) -Initiate intensifying pharmacologic treatment in adults aged 60 and older with a history of stroke or transient ischemic attack to achieve a target systolic blood pressure of less than 140 mmHg to reduce the risk of recurrent stroke (Grade: weak recommendation, quality of evidence: moderate) American Diabetes Association (2018): Most patients with diabetes and hypertension should be treated to a systolic blood pressure goal of <140 mmHg and a diastolic blood pressure goal of <90 mmHg (Level of evidence: A) Report from the Eighth Joint National Committee (2014) -In the general population younger than 60 years, initiate pharmacologic treatment to lower blood pressure at diastolic blood pressure (DBP) of 90 mmHg or higher and treat to a goal of DBP of lower than 90 mmHg (Grade: A (for ages 30-59), Grade: E (for ages 18-29)) -In the general population younger than 60 years, initiate pharmacologic treatment to lower blood pressure at systolic blood pressure (SBP) to 140 mmHg or higher and treat to a goal of SBP of lower than 140 mmHg (Grade: E) -In the general population aged 60 years and older, initiate pharmacologic treatment to lower blood pressure at SBP of 150 mmHg or higher or a DBP of 90 mmHg or higher and treat to a goal of SBP lower than 150 mmHg and goal of DBP lower than 90 mmHg
**Guidance:**

In reference to the numerator element, only blood pressure readings performed by a clinician or a remote monitoring device are acceptable for numerator compliance with this measure. Do not include BP readings: -Taken during an acute inpatient stay or an ED visit -Taken on the same day as a diagnostic test or diagnostic or therapeutic procedure that requires a change in diet or change in medication on or one day before the day of the test or procedure, with the exception of fasting blood tests. -Reported by or taken by the member If no blood pressure is recorded during the measurement period, the patient’s blood pressure is assumed "not controlled." If there are multiple blood pressure readings on the same day, use the lowest systolic and the lowest diastolic reading as the most recent blood pressure reading.

**Initial Population:**
Patients 18-85 years of age who had a visit and diagnosis of essential hypertension overlapping the measurement period.

**Denominator:**
Equals Initial Population.

**Denominator Exclusions:**
Patients with evidence of end stage renal disease (ESRD), dialysis or renal transplant before or during the measurement period. Also exclude patients with a diagnosis of pregnancy during the measurement period. Exclude patients whose hospice care overlaps the measurement period. Exclude patients 66 and older who are living long term in an institution for more than 90 days during the measurement period. Exclude patients 66 and older with advanced illness and frailty because it is unlikely that patients will benefit from the services being measured.

**Numerator:**
Patients whose most recent blood pressure is adequately controlled (systolic blood pressure < 140 mmHg and diastolic blood pressure < 90 mmHg) during the measurement period.

**Numerator Exclusions:**
Not Applicable.

**Denominator Exceptions:**
None.

**Supplemental Data Elements:**
For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.21.2 Data Entry

**Description:**
Percentage of patients 18-85 years of age who had a diagnosis of hypertension and whose blood pressure was adequately controlled (<140/90mmHg) during the measurement period.

Visit or inpatient admission, age equal or greater than 18 and less than 85 at start of measurement period.
Diagnosis (on IPL) of Essential Hypertension
(Value Set: 2.16.840.1.113883.3.464.1003.104.12.1011) at start or within 6 months of start of measurement period.

“Qualifying encounter:”

- Annual Wellness Visit (2.16.840.1.113883.3.526.3.1240).
- Encounter Inpatient (2.16.840.1.113883.3.666.5.307).
- Preventive Care Services (2.16.840.1.113883.3.464.1003.101.12.1023, ….1025).
- Nursing facility visit
- Services in Long-term residential facility

Denominator Exclusions:

- Hospice order/intervention or hospital discharge to hospice during measurement period: (Value Set: 2.16.840.1.113762.1.4.1108.15). Inpatient hospice discharge entered by Coding. Outpatient entered on IPL/TREG:

![Palliative Care list](image)

Figure 3-27: Palliative Care list

- Pregnancy dx during measurement period
  (Value Set: 2.16.840.1.113883.3.526.3.378)
- End stage renal disease during measurement period
  (Value Set: 2.16.840.1.113883.3.526.3.353).
- Chronic kidney disease (stage 5) or kidney transplant recipient overlaps measurement period (Value Set: 2.16.840.1.113883.3.526.3.1002), (Value Set: 2.16.840.1.113883.3.464.1003.109.12.1029).
- Kidney transplant, dialysis services
- Vascular access for dialysis
- ESRD Monthly Outpatient Services  
  (Value Set: 2.16.840.1.113883.3.464.1003.109.12.1014) starts before end of measurement period.

Numerator: Last PE during measurement period BOTH diastolic BP < 90 and systolic BP <140.

### 3.22 CMS 177V8 Child and Adolescent MDD: Suicide Risk Assessment

#### 3.22.1 Detail

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder with an assessment for suicide risk.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rationale</td>
<td>Research has shown that patients with major depressive disorder are at a high risk for suicide attempts and completion - among the most significant and devastating sequelae of the disease. Suicide risk is a critical consideration in children and adolescents with MDD and an important aspect of care that should be assessed at each visit and subsequently managed to minimize that risk. Additionally, the importance of the assessments is underscored by research that indicates that many individuals who die by suicide do make contact with primary care providers and mental health services beforehand. More specifically, approximately 15% of suicide victims aged 35 years or younger had seen a mental health professional within 1 month of suicide while approximately 23% had seen a primary care provider within 1 month of suicide.</td>
</tr>
</tbody>
</table>
The evaluation must include assessment for the presence of harm to self or others (MS) (American Academy of Child and Adolescent Psychiatry, 2007). Suicidal behavior exists along a continuum from passive thoughts of death to a clearly developed plan and intent to carry out that plan. Because depression is closely associated with suicidal thoughts and behavior, it is imperative to evaluate these symptoms at the initial and subsequent assessments. For this purpose, low burden tools to track suicidal ideation and behavior such as the Columbia-Suicidal Severity Rating Scale can be used. Also, it is crucial to evaluate the risk (e.g., age, sex, stressors, comorbid conditions, hopelessness, impulsivity) and protective factors (e.g., religious belief, concern not to hurt family) that might influence the desire to attempt suicide. The risk for suicidal behavior increases if there is a history of suicide attempts, comorbid psychiatric disorders (e.g., disruptive disorders, substance abuse), impulsivity and aggression, availability of lethal agents (e.g., firearms), exposure to negative events (e.g., physical or sexual abuse, violence), and a family history of suicidal behavior (American Academy of Child and Adolescent Psychiatry, 2007).

A careful and ongoing evaluation of suicide risk is necessary for all patients with major depressive disorder (Category I). Such an assessment includes specific inquiry about suicidal thoughts, intent, plans, means, and behaviors; identification of specific psychiatric symptoms (e.g., psychosis, severe anxiety, substance use) or general medical conditions that may increase the likelihood of acting on suicidal ideas; assessment of past and, particularly, recent suicidal behavior; delineation of current stressors and potential protective factors (e.g., positive reasons for living, strong social support); and identification of any family history of suicide or mental illness (Category I) (American Psychiatric Association, 2010, reaffirmed 2015).

**Clinical Recommendation Statement**

Numerator Definition: The specific type and magnitude of the suicide risk assessment is intended to be at the discretion of the individual clinician and should be specific to the needs of the patient. At a minimum, suicide risk assessment should evaluate: 1. Risk (e.g., age, sex, stressors, comorbid conditions, hopelessness, impulsivity) and protective factors (e.g., religious belief, concern not to hurt family) that may influence the desire to attempt suicide. 2. Current severity of suicidality. 3. Most severe point of suicidality in episode and lifetime. Low burden tools to track suicidal ideation and behavior such as the Columbia-Suicidal Severity Rating Scale can also be used. Because no validated assessment tool or instrument fully meets the aforementioned requirements for the suicide risk assessment, individual tools or instruments have not been explicitly included in coding.
Guidance

A suicide risk assessment should be performed at every visit for major depressive disorder during the measurement period. Suicide risk assessments completed via telehealth services can also meet numerator performance. This measure is an episode-of-care measure; the level of analysis for this measure is every visit for major depressive disorder during the measurement period. For example, at every visit for MDD, the patient should have a suicide risk assessment. Use of a standardized tool(s) or instrument(s) to assess suicide risk will meet numerator performance, so long as the minimum criteria noted above is evaluated. Standardized tools can be mapped to the concept "Intervention, Performed": "Suicide risk assessment (procedure)" included in the numerator logic below, as no individual suicide risk assessment tool or instrument would satisfy the requirements alone.

<table>
<thead>
<tr>
<th>Initial Population</th>
<th>All patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator</td>
<td>Equals Initial Population.</td>
</tr>
<tr>
<td>Denominator Exclusions</td>
<td>None.</td>
</tr>
<tr>
<td>Numerator</td>
<td>Patient visits with an assessment for suicide risk.</td>
</tr>
<tr>
<td>Numerator Exclusions</td>
<td>Not Applicable.</td>
</tr>
<tr>
<td>Denominator Exceptions</td>
<td>None.</td>
</tr>
<tr>
<td>Supplemental Data Elements</td>
<td>For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.</td>
</tr>
</tbody>
</table>

3.22.2 Detail

**Description:** Percentage of patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder with an assessment for suicide risk.

Create qualifying encounter; denominator is total number of visits during measurement period:
- Age equal or > 6 and equal or <17 at start of measurement period.
- Diagnosis = major depressive disorder (Value Set: 2.16.840.1.113883.3.526.3.1491).

Encounters include:
- Group Psychotherapy (Value Set 2.16.840.1.113883.3.526.3.1187).
- Psych Visit - Diagnostic Evaluation (Value Set 2.16.840.1.113883.3.526.3.1492).
• Psych Visit - Family Psychotherapy (Value Set 2.16.840.1.113883.3.526.3.1018).
• Psych Visit - Psychotherapy (Value Set 2.16.840.1.113883.3.526.3.1496).
• Psychoanalysis (Value Set 2.16.840.1.113883.3.526.3.1141).
• Outpatient consultation (Value Set 2.16.840.1.113883.3.464.1003.101.12.1008).
• Telehealth Visit 2.16.840.1.113883.3.464.1003.101.12.1031

See Appendix A for information on using Value Sets.

“Major depressive disorder diagnoses” include 74 codes (Value Set: 2.16.840.1.113883.3.526.3.1491)

Numerator: Suicide risk assessment at every visit for a major depressive diagnosis during measurement period (if multiple visits, the numerator is total number with suicide risk assessed and documented) (Enter in Exams at every visit).

3.23 CMS 347V3 Statin Therapy for the Prevention and Treatment of Cardiovascular Disease

3.23.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of the following patients - all considered at high risk of cardiovascular events - who were prescribed or were on statin therapy during the measurement period:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Adults aged &gt;= 21 years who were previously diagnosed with or currently have an active diagnosis of clinical atherosclerotic cardiovascular disease (ASCVD).</td>
<td></td>
</tr>
<tr>
<td>• Adults aged &gt;= 21 years who have ever had a fasting or direct LDL-C level &gt;= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia.</td>
<td></td>
</tr>
<tr>
<td>• Adults aged 40-75 years with a diagnosis of diabetes with a fasting or direct LDL-C level of 70-189 mg/dL.</td>
<td></td>
</tr>
</tbody>
</table>
This measure is intended to have one reporting rate, which aggregates the following populations into a single performance rate for reporting purposes:

**Population 1:** Patients aged >= 21 years at the beginning of the measurement period with clinical ASCVD

**Population 2:** Patients aged >= 21 years at the beginning of the measurement period who have ever had a fasting or direct laboratory test result of LDL-C >= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia

**Population 3:** Patients aged 40 to 75 years at the beginning of the measurement period with Type 1 or Type 2 diabetes and with a LDL-C result of 70 -189 mg/dL recorded as the highest fasting or direct laboratory test result in the measurement year or during the two years prior to the beginning of the measurement period

For the purposes of this measure, a single performance rate can be calculated as follows:

\[
\text{Performance Rate} = \frac{(\text{Numerator 1} + \text{Numerator 2} + \text{Numerator 3})}{(\text{Denominator 1} - \text{Denominator Exclusions 1} - \text{Denominator Exceptions 1}) + (\text{Denominator 2} - \text{Denominator Exclusions 2} - \text{Denominator Exceptions 2}) + (\text{Denominator 3} - \text{Denominator Exclusions 3} - \text{Denominator Exceptions 3})}
\]
Rationale:

"Cardiovascular disease (CVD) is the leading cause of death in the United States, causing approximately 1 of every 3 deaths in the United States in 2015. In 2015, stroke caused approximately 1 of every 19 deaths in the United States and the estimated annual costs for CVD and stroke were $329.7 billion, including $199.2 billion in direct costs (hospital services, physicians and other professionals, prescribed medications, home health care, and other medical durables) and $130.5 billion in indirect costs from lost future productivity (cardiovascular and stroke premature deaths). CVD costs more than any other diagnostic group" (Benjamin et al., 2018). Data collected between 2011 and 2014 indicates that more than 94.6 million U.S. adults, 20 years or older, had total cholesterol levels equal to 200 mg/dL or more, while almost 28.5 million had levels 240 mg/dL or more (Benjamin et al., 2018). Elevated blood cholesterol is a major risk factor for CVD and statin therapy has been associated with a reduced risk of CVD. Numerous randomized trials have demonstrated that treatment with a statin reduces LDL-C, and reduces the risk of major cardiovascular events by approximately 20 percent (Ference, 2015). In 2013, guidelines on the treatment of blood cholesterol to reduce atherosclerotic cardiovascular risk in adults were published (see Stone et al., 2014). This guideline was published by an Expert Panel, which synthesized evidence from randomized controlled trials to identify people most likely to benefit from cholesterol-lowering therapy. The American College of Cardiology (ACC)/American Heart Association (AHA) Guideline recommendations are intended to provide a strong evidence-based foundation for the treatment of blood cholesterol for the primary and secondary prevention and treatment of Atherosclerotic Cardiovascular Disease (ASCVD) in adult men and women (21 years of age or older). The document concludes the addition of statin therapy reduces the risk of ASCVD among high-risk individuals, defined as follows: individuals with clinical ASCVD, with LDL-C >= 190 mg/dL, or with diabetes and LDL-C 70-189 mg/dL (Stone et al., 2014). One study that surveyed U.S. cardiology, primary care, and endocrinology practices found that 1 in 4 guideline-eligible patients were not on a statin and less than half were on the recommended statin intensity. Untreated and undertreated patients had significantly higher LDL-C levels than those receiving guideline-directed statin treatment (Navar et al., 2017). The Statin Safety Expert Panel that participated in an NLA Statin Safety Task Force meeting in October 2013 reaffirms the general safety of statin therapy. However, 1 in 10 people who try taking a statin will report some kind of intolerance, most commonly muscle aches. Other known low risk circumstances of statin intolerance includes side effects such as myopathy, cognitive dysfunction, increased hepatic transaminase levels, and new onset diabetes. Statin intolerance usually does not involve substantial risk for mortality or permanent disability (Guyton et al., 2014).
### Rationale

Ultimately, the panel members concluded that for most patients requiring statin therapy, the potential benefits of statin therapy outweigh the potential risks. In general terms, the benefits of statins to prevent non-fatal myocardial infarction, revascularization, stroke, and CVD mortality, far outweighs any potential harm related to the drug (Jacobson, 2014).

### Clinical Recommendation Statement:

This electronic clinical quality measure is intended to align with the 2013 ACC/AHA Guideline on the Treatment of Blood Cholesterol (Stone et al., 2014), which indicates the use of statins as the first line of cholesterol-lowering medication therapy to lower the risk of ASCVD among at-risk populations. Recommendations for Treatment of Blood Cholesterol to Reduce Atherosclerotic Cardiovascular Risk in Adults - Statin Treatment: Secondary Prevention:

1. High-intensity statin therapy should be initiated or continued as first-line therapy in women and men <=75 years of age who have clinical ASCVD, unless contraindicated. (Level of Evidence A), (Stone et al., 2014)

2. In individuals with clinical ASCVD in whom high-intensity statin therapy would otherwise be used, when high-intensity statin therapy is contraindicated or when characteristics predisposing to statin-associated adverse effects are present, moderate-intensity statin should be used as the second option, if tolerated. (Level of Evidence A), (Stone et al., 2014)

Primary Prevention in Individuals >=21 Years of Age With LDL-C >=190 mg/dL:

1. Adults >=21 years of age with primary LDL-C >=190 mg/dL should be treated with statin therapy. (10-year ASCVD risk estimation is not required.) (Level of Evidence B), (Stone et al., 2014)

Primary Prevention in Individuals With Diabetes and LDL-C 70-189 mg/dL:

1. Moderate-intensity statin therapy should be initiated or continued for adults 40-75 years of age with diabetes. (Level of Evidence A), (Stone et al., 2014)
**Definition:**

Clinical atherosclerotic cardiovascular disease (ASCVD) includes:
- Acute coronary syndromes
- History of myocardial infarction
- Stable or unstable angina
- Coronary or other arterial revascularization
- Stroke or transient ischemic attack (TIA)
- Peripheral arterial disease of atherosclerotic origin
- Lipoprotein density cholesterol (LDL-C) result: A fasting or direct LDL-C laboratory test performed and test result documented in the medical record.
- Statin therapy: Administration of one or more of a group of medications that are used to lower plasma lipoprotein levels in the treatment of hyperlipoproteinemia.

<table>
<thead>
<tr>
<th>Statin Medication Therapy List (NOTE: List does NOT include dosage):</th>
<th>[Generic name] (Brand or trade name) and (-) Medication type, if applicable:</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Atorvastatin] (Lipitor) - Statin</td>
<td>[Fluvastatin] (Lescol XL or Lescol) - Statin</td>
</tr>
<tr>
<td>[Lovastatin (Mevinolin)](Mevacor or Altoprev) - Statin</td>
<td>[Pitavastatin]Livalo [Pravastatin Sodium] (Pravachol) - Statin</td>
</tr>
<tr>
<td>[Rosuvastatin Calcium] (Crestor) - Statin</td>
<td>[Simvastatin] (Zocor) - Statin</td>
</tr>
<tr>
<td>[Amlodipine Besylate/Atorvastatin Calcium] (Caduet) - Combination</td>
<td>[Ezetimibe/Simvastatin] (Vytorin) - Combination</td>
</tr>
</tbody>
</table>

Some patients may not be appropriate to prescribe or use statin therapy (see exceptions and exclusions for a complete list). "Statin intolerance is the inability to tolerate a dose of statin required to reduce a person’s CV risk sufficiently from their baseline risk and could result from different statin related side effects including: muscle symptoms, headache, sleep disorders, dyspepsia, nausea, rash, alopecia, erectile dysfunction, gynecomastia, and/or arthritis" (Banach et al., 2015, p.2). Patients that experience symptoms such as these may prefer not to take or continue statin therapy and therefore may be exempt from the denominator.
| **Guidance:** | Numerator instructions and guidance: -Current statin therapy use must be documented in the patient's current medication list or ordered during the measurement period. -ONLY statin therapy meets the measure Numerator criteria (NOT other cholesterol lowering medications). -Prescription or order does not need to be linked to an encounter or visit; it may be called to the pharmacy. -Statin medication "samples" provided to patients can be documented as "current statin therapy" if documented in the medication list in health/medical record. -Patients who meet the denominator criteria for inclusion, but are not prescribed or using statin therapy, will NOT meet performance for this measure. There is only one performance rate calculated for this measure; the weighted average of the three populations. -Adherence to statin therapy is not calculated in this measure. Denominator Guidance: The denominator covers three distinct populations. Use the following process to prevent counting patients more than once. Denominator Population 1: Patients aged >= 21 years at the beginning of the measurement period with clinical ASCVD -If YES, meets Denominator Population 1 risk category -If NO, screen for next risk category Denominator Population 2: Patients aged >= 21 years at the beginning of the measurement period who have ever had a fasting or direct laboratory test result of LDL-C >= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia -If YES, meets Denominator Population 2 risk category -If NO, screen for next risk category Denominator Population 3: Patients aged 40 to 75 years at the beginning of the measurement period with Type 1 or Type 2 diabetes and with a LDL-C result of 70 -189 mg/dL recorded as the highest fasting or direct laboratory test result in the measurement year or during the two years prior to the beginning of the measurement period -If YES, meets Denominator Population 3 risk category -If NO, patient does NOT meet Denominator criteria and is NOT eligible for measure inclusion Denominator Guidance for Encounter: -In order for the patient to be included in the denominator, the patient must have ONE denominator-eligible visit, defined as follows: --Outpatient encounter visit type --Encounter, performed: initial or established office visit, face-to-face interaction, preventive care services, or annual wellness visit LDL-C Laboratory test result options: The measure can be reported for all patients with a documented fasting or direct LDL-C level recorded as follows: To meet Denominator Population 1: There is no LDL-C result required. To meet Denominator Population 2: If a patient has ANY previous fasting or direct laboratory result of LDL-C >= 190 mg/dL, report the highest value >= 190 mg/dL. To meet Denominator Population 3: If a patient has more than one LDL-C result during the measurement period or during the two years before the start of the measurement period, report the highest level recorded during either time. The Denominator Exception, "Patients with diabetes who have the most recent fasting or direct LDL-C laboratory test result < 70 mg/dL and are not taking statin therapy" applies only to Denominator Population 3. Intensity of statin therapy in primary and secondary prevention: |

### Measure Guidance Manual

Provider Measures for 2020 Reporting Period

December 2020
| **Guidance:** | The expert panel of the 2013 ACC/AHA Guidelines (Stone et al., 2014) defines recommended intensity of statin therapy on the basis of the average expected LDL-C response to specific statin and dose. Although intensity of statin therapy is important in managing cholesterol, this measure assesses prescription of ANY statin therapy, irrespective of intensity. Assessment of appropriate intensity and dosage documentation added too much complexity to allow inclusion of statin therapy intensity in the measure at this time. Lifestyle modification coaching: A healthy lifestyle is important for the prevention of cardiovascular disease. However, lifestyle modification monitoring and documentation added too much complexity to allow its inclusion in the measure at this time. |
| **Initial Population:** | All patients aged 21 years and older at the beginning of the measurement period with a patient encounter during the measurement period. |
| **Denominator:** | All patients who meet one or more of the following criteria (considered at "high risk" for cardiovascular events, under ACC/AHA guidelines): Patients aged >= 21 years at the beginning of the measurement period with clinical ASCVD diagnosis Patients aged >= 21 years at the beginning of the measurement period who have ever had a fasting or direct laboratory result of LDL-C >= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia Patients aged 40 to 75 years at the beginning of the measurement period with Type 1 or Type 2 diabetes and with an LDL-C result of 70-189 mg/dL recorded as the highest fasting or direct laboratory test result in the measurement year or during the two years prior to the beginning of the measurement period. |
| **Denominator Exclusions:** | Patients who have a diagnosis of pregnancy. Patients who are breastfeeding. Patients who have a diagnosis of rhabdomyolysis. |
| **Numerator:** | Patients who are actively using or who receive an order (prescription) for statin therapy at any point during the measurement period. |
| **Numerator Exclusions:** | None. |
| **Denominator Exceptions:** | Patients with adverse effect, allergy, or intolerance to statin medication Patients who are receiving palliative or hospice care Patients with active liver disease or hepatic disease or insufficiency Patients with end-stage renal disease (ESRD) Patients with diabetes who have the most recent fasting or direct LDL-C laboratory test result < 70 mg/dL and are not taking statin therapy |
| **Supplemental Data Elements:** | For every patient evaluated by this measure also identify payer, race, ethnicity, and sex. |
3.23.2 Data Entry

**Description:** Percentage of the following patients - all considered at high risk of cardiovascular events - who were prescribed or were on statin therapy during the measurement period:

- Adults aged >= 21 years who were previously diagnosed with or currently have an active diagnosis of clinical ASCVD.
- Adults aged >= 21 years who have ever had a fasting or direct low-density lipoprotein cholesterol (LDL-C) level >= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia.
- Adults aged 40-75 years with a diagnosis of diabetes with a fasting or direct LDL-C level of 70-189 mg/dL.

Three Measures based on age and risk

**Denominator 1:**

- Age equal or >21 at start of measurement period.
- Qualifying encounter during measurement period.
- Annual Wellness Visit (Value Set 2.16.840.1.113883.3.526.2.1363).
- Office Visit (Value Set 2.16.840.1.113883.3.464.1003.101.11.1005).
- Outpatient Consultation (Value Set 2.16.840.1.113883.3.464.1003.101.11.1040).
- Outpatient Encounters for Preventive Care (Value Set 2.16.840.1.113762.1.4.1047.9).
- Palliative care encounter (Value Set 2.16.840.1.113883.3.600.1.1575).
- Preventive Care Services (Value Set 2.16.840.1.113883.3.464.1003.101.11.1125).
- Preventive Care Services – Other (Value Set 2.16.840.1.113883.3.464.1003.101.12.1030).
- Preventive Care Services-Individual Counseling (Value Set 2.16.840.1.113883.3.464.1003.101.12.1026).
- Preventive Care Services-Initial Office Visit (Value Set 2.16.840.1.113883.3.464.1003.101.11.1115).
- Clinical diagnosis of ASCVD (Value Set: 2.16.840.1.113762.1.4.1047.21).

**Denominator exclusions:**

- Diagnosis of breastfeeding (Value Set: 2.16.840.1.113762.1.4.1047.73).
- Pregnancy (Value Set: 2.16.840.1.113883.3.600.1.1623).
- Rhabdomyolysis (Value Set: 2.16.840.1.113762.1.4.1047.102) overlapping measurement period.
Numerator 1:
- Medication order or active prescription for statin (low, moderate, or high intensity):
  (Value Set: 2.16.840.1.113762.1.4.1047.107.
  Value Set: 2.16.840.1.113762.1.4.1047.98.
  Value Set: 2.16.840.1.113762.1.4.1047.97).

Denominator Exceptions:
- Palliative care (Value Set: 2.16.840.1.113883.3.600.1.1579).
- ESRD (Value Set: 2.16.840.1.113883.3.526.3.353).
- Hepatitis B (Value Set: 2.16.840.1.113883.3.67.1.101.1.269).
- Liver Disease (Value Set: 2.16.840.1.113762.1.4.1047.42).
- Statin allergy (Value Set: 2.16.840.1.113883.3.117.1.7.1.423), overlapping measurement period.

Denominator 2:
- Age equal or >21 at start of measurement period.
- Qualifying encounter during measurement period (face to face, annual wellness visit, preventive care services, initial or established office visit. See value sets for details).
- LDL>190 or Diagnosis Hypercholesterolemia BUT without diagnosis of ASCVD or procedure before end of measurement period.

Denominator exclusions:
- Diagnosis of breastfeeding, pregnancy, or rhabdomyolysis overlapping measurement period.

Numerator 2: Medication order or active prescription for statin (low, moderate or high intensity).

Denominator Exceptions: Palliative care, ESRD, Hepatitis A, Hepatitis B, Liver Disease, or Statin allergy overlapping measurement period.

Denominator 3:
- Age equal or >40 and less than or equal to 75 at start of measurement period.
- Qualifying encounter during measurement period (face to face, annual wellness visit, preventive care services, initial or established office visit).
- Diagnosis = Diabetes (Value Set: 2.16.840.1.113883.3.464.1003.103.12.1001).
• LDL result equal to or >70 mg/dl AND <190 during 3 years before END of measurement period.
• No preexisting ASCVD Diagnosis or procedure, no diagnosis of hypercholesterolemia.
• No LDL result before end of measurement period greater than or equal to 190.

Denominator exclusions:
• Diagnosis of breastfeeding, pregnancy, or rhabdomyolysis overlapping measurement period.

Numerator 3: Medication order or active prescription for statin (low, moderate, or high intensity).

Denominator Exceptions: Palliative care, ESRD, Hepatitis A, Hepatitis B, Liver Disease, or Statin allergy overlapping measurement period.

Denominator exception: LDL within 3 years of end of measurement period <70mg/dl,

3.24 CMS 349V2 HIV Screening

3.24.1 Detail

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients aged 15-65 at the start of the measurement period who were between 15-65 years old when tested for HIV</th>
</tr>
</thead>
</table>
**Rationale:**

Human immunodeficiency virus (HIV) is a communicable infection that leads to a progressive disease with a long asymptomatic period. There were an estimated 38,500 new HIV infections in the United States in 2015 (Centers for Disease Control and Prevention, 2018a). Without treatment, most persons develop acquired immunodeficiency syndrome (AIDS) within 10 years of HIV infection. Antiretroviral therapy (ART) delays this progression and increases the length of survival, but it is most effective when initiated during the asymptomatic phase. Persons living with HIV who use ART and achieve viral suppression can have a nearly normal life expectancy (Samji et al., 2013). DHHS Guidelines for the Use of Antiretroviral Agents in HIV-1-Infected Adults and Adolescents recommends antiretroviral therapy for all HIV-infected individuals to reduce the risk of disease progression (regardless of CD4 cell count at diagnosis) (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2017). CDC estimates that almost 15% of the 1.1 million adults and adolescents living with HIV infection in the United States are unaware of their infection (Centers for Disease Control and Prevention, 2018b). Among persons diagnosed with HIV in 2011, one quarter were diagnosed with Stage 3 HIV (AIDS) at the time of HIV diagnosis (Centers for Disease Control and Prevention, 2018c), which is when the median CD4 count at diagnosis is less than 200 cells/mm3 for persons aged greater than or equal to 6 years (Centers for Disease Control and Prevention, 2018a). HIV screening identifies persons who were previously unaware of their infection, which enables them to seek medical and social services that can improve their health and the quality and length of their lives. Additionally, using ART with high levels of medication adherence has been shown to substantially reduce risk for HIV transmission (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2017). Based on the National Health Interview Survey, fewer than half of persons 18 and older reported ever having been tested for HIV as of 2017 (Blackwell et al., 2019).

**Clinical Recommendation Statement:**

The US Preventive Services Task Force recommends that clinicians screen for HIV infection in adolescents and adults aged 15 to 65 years. Younger adolescents and older adults who are at increased risk should also be screened (A Recommendation) (Moyer, 2013). Since 2006, the CDC has recommended routine opt-out HIV screening (i.e., patient is notified that testing will be performed unless the patient declines) in healthcare facilities of adolescents and adults 13-64 years of age and HIV diagnostic testing of adolescents and adults with clinical signs or symptoms consistent with HIV infection (Centers for Disease Control and Prevention, 2006).
**Guidance:**

This measure evaluates the proportion of patients aged 15 to 65 at the start of the measurement period who have documentation of having received an HIV test at least once on or after their 15th birthday and before their 66th birthday. In order to satisfy the measure, the reporting provider must have documentation of the administration of the laboratory test present in the patient's medical record. In cases where the HIV test was performed elsewhere, providers cannot rely on patient attestation or self-report to meet the measure requirements, as previous research has shown that patient self-report is an unreliable indicator of previous HIV testing history. Rather, providers must request documentation of those test results. If such documentation is not available, the patient should be considered still eligible for HIV screening. If such documentation is available, but cannot be provided in a standardized, structured format (such that the lab test and results can be readily incorporated as structured data within the EHR), providers should enter the information into their EHR as a laboratory test in a manner consistent with the EHR in use. If the specific Human Immunodeficiency Virus (HIV) Laboratory Test LOINC code of the test is not known, the entry should use the more generic code LOINC panel code [75622-1].

**Initial Population:**

Patients 15 to 65 years of age at the start of the measurement period AND who had at least one outpatient visit during the measurement period.

**Denominator:**

Equals Initial Population.

**Denominator Exclusions:**

Patients diagnosed with HIV prior to the start of the measurement period.

**Numerator:**

Patients with documentation of an HIV test performed on or after their 15th birthday and before their 66th birthday.

**Numerator Exclusions:**

Not Applicable.

**Denominator Exceptions:**

None.

**Supplemental Data Elements:**

For every patient evaluated by this measure also identify payer, race, ethnicity, and sex.

### 3.24.2 Data Entry

**Description:**

Percentage of patients 15-65 years of age who have been tested for HIV within that age range.

Initial population: Age equal to or greater than 15, and less than 66 at start of measurement period

Qualifying encounter:

- Office Visit (Value Set 2.16.840.1.113883.3.464.1003.101.12.1001)
• Preventive Care Services
  (Value Set 2.16.840.1.113883.3.464.1003.101.12.1022, ….1023, ….1024, …1025)

Denominator exclusions: Diagnosis of “Conditions Due to HIV” before start of measurement period (Value Set: 2.16.840.1.113762.1.4.1056.54)

Numerator (either of the following):
• Lab test HIV codes Ag and Ab (Value Set: 2.16.840.1.113762.1.4.1056.50)
• HIV Tests 1 and 2 (https://r.details.loinc.org/LOINC/75622-1.html), resulted before end of measurement period and when patient age 15–66.
Appendix A  Using the Value Set Authority Center

Using the VSAC to review approved codes for each measure. Codes displayed may be ICD codes, SNOMED codes, CPT codes or others depending on the measure logic. There are value sets for diagnoses, encounter codes, procedures. To receive credit for any particular measure, you must be using one of these codes.

To log on to your existing account, do the following:

1. Log on to the VSAC (https://vsac.nlm.nih.gov/welcome) (or create a new, free account Figure A-1).
2. Click the **CMS eCQM Value Sets** search box (Figure A-2).

![Figure A-2: Value Set Authority Center Welcome page](image)

The following search page displays:

![Figure A-3: Value Set Authority Center search page](image)
3. In the query field, enter the Value Set identification number. In this example enter the code for depression screening encounters (2.16.840.1.113883.3.600.1916), then click Search. Value Set identification numbers are found for each measure in the section, “Value Sets.”

Figure A-4: Value Set Authority Center search page

4. Click the underlined hyperlink on the right side (Figure A-5).

Figure A-5: Value Set Authority Center search results page
5. Scroll to view all of the codes (106 of them in this case).
Appendix B  Using the United States Health Information Knowledgebase

The United States Health Information Knowledgebase (USHIK) is an on-line, publicly accessible registry and repository of healthcare-related metadata, specifications, and standards. This includes information about changes in CQM and changes in Value Sets that allow comparison.

Establish an account (Free):


   ![Figure B-1: United States Health Information Knowledgebase main page](image1)

2. Click Quality Reporting in the top menu bar.

   ![Figure B-2: USHIK Quality Reporting](image2)
3. Click **Clinical Quality Measures** on the left.

![Figure B-3: eCQM Reporting](image)

4. This is a **very** long list, but you can filter it by selecting the most recent version from the **release package** dropdown. You can also compare two or more years by checking more than one in that dropdown. Or you can select hospital measures from the **Eligibility** dropdown (this screen is for providers).

5. Here two different years are selected:

![Figure B-4: eCQM Updates](image)

6. Apply these filters to see the following: Note different versions for different years.
7. You can select different versions to compare by checking the boxes on the left, then click the compare button.

8. You can see all of the recent versions and check any you want to view (useful for troubleshooting when the value sets have changed). Scroll down to see a narrative, and then click the QDM Data Elements and Codes button.
Figure B-7: CQM Data Elements and Codes

Version 7 codes are on the left. Version 8 on the right. You can hover over any code to get a description. Codes that are highlighted in blue have not changed and are shared by both versions. Codes that have been changed, added, or deleted, are grey, as below:

Figure B-8: CQM Data Elements and Codes

9. Codes for telehealth services are new in this version and were not in a prior version.
Appendix C  Using The Electronic Clinical Quality Improvement Center

C.1  Access eCQI


2. “Electronic Clinical Quality Improvement (eCQI) Resource Center - The one-stop shop for the most current resources to support electronic clinical quality improvement.” This replaces the USHIK site to some extent but is different. Comparing different years is easier on the USHIK site. There is a lot of information on this site. To see details, log on using the same information used for VSAC or USHIK.

3. Click the eCQM menu to select hospital or provider measures.

4. Scroll down to select the performance period (2019 selected).
Figure C-3: Eligible Professional/Clinician eCQMs

5. Then select from the extensive list of links to view items such as eCQM specifications. Clicking Value Sets addendum displays the VASC. eCQM flows show the wireframes or flow diagrams for each measure.
6. Scroll to the bottom to see the list of measures (for 2019 and earlier):
### Electronic Clinical Quality Measures (ECQM) Version 3.0

**Measure Guidance Manual Using The Electronic Clinical Quality Improvement Center**

**December 2020**

#### Figure C-5: Performance Period Links

7. Click one of these (e.g., Breast Cancer Screening) to see the specifications.

#### Breast Cancer Screening

**eCQMs for 2019 Performance Period**

<table>
<thead>
<tr>
<th>CMS Measure ID</th>
<th>Version</th>
<th>HQR Number</th>
<th>Measure Description</th>
<th>Initial Population</th>
<th>Denominator Statement</th>
<th>Denominator Exclusions</th>
<th>Numerator Statement</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS125v7</td>
<td>7</td>
<td>2372</td>
<td>Percentage of women 50-74 years of age who had a mammogram to screen for breast cancer.</td>
<td>Women 51-74 years of age with a visit during the measurement period.</td>
<td>Equals Initial Population.</td>
<td>Women who had a bilateral mastectomy or who have a history of a bilateral mastectomy or for whom there is evidence of a right and a left unilateral mastectomy.</td>
<td>Women with one or more mammograms during the measurement period or the 16 months prior to the measurement period.</td>
</tr>
</tbody>
</table>

---

**Figure C-6: Performance Period detail**
8. Scrolling down on this page to “Specifications” allows you to download details of the measure in various views (Word or PDF).

C.2 Shortcut to Wireframes

1. Launch one of the following:
   - For Provider Measures: https://ecqi.healthit.gov/ep-ec-ecqms
   - For Hospital Measures: https://ecqi.healthit.gov/eh-cah-ecqms

2. Scroll through the displayed list to find the link **eCQM Flows (ZIP)**.

3. Click the link. The associated ZIP file is displayed at the bottom left of the page (Figure C-7).

4. Click the ZIP file name. The contents of the file are displayed.

5. Double click the PDF file to be displayed.
Appendix D Using Treatment/Regimen/Follow Up in RPMS EHR

There are multiple ways to capture data in the RPMS EHR, but one of the fastest for certain data types is TREG (Treatment/Regimen/Follow-up). Data entered here is saved to the visit file where it can be used by eCQM reports. Not all data from TREG can be used for eCQM, however, including lab tests and medication orders. TREG only saves in SNOMED format, while lab tests are usually stored as LOINC codes, and medications as RxNORM codes. These data must be saved in these formats to be used for eCQM.

To access TREG:
1. Create a visit in EHR.
2. Enter a POV.
3. Highlight the POV and click Edit:

![Figure D-1: Problem Management tab](image)

4. On the Edit screen (Figure D-2), select Add Visit Instruction/Care Plans/Goal Activities from the lower right side:

![Figure D-2: Edit Screen](image)
5. This dialog box displays. Choose from among several education options in the upper half or click **Treatment/Regimen/Follow-up** and click **OK**.

![Patient Education Provided dialog](image)

Figure D-3: Patient Education Provided dialog

6. There are multiple options on this screen. Click the plus sign (+) to expand a list:

![Treatment/Regimen list](image)

Figure D-4: Treatment/Regimen list

![Anticoag DVT Prevention list](image)

Figure D-5: Anticoag DVT Prevention list
Figure D-6: Asthma list

Figure D-7: Behavioral Health list

Figure D-8: Case Management list

Figure D-9: Controlled Substances list

Figure D-10: Dialysis list

Figure D-11: Disposition list
Figure D-12: Follow-up list

Figure D-13: Massage Therapy list

Figure D-14: Nursing list

Figure D-15: Palliative Care list (These save codes are used by eCQM)
Figure D-16: Rehab Services list (These save codes are used by eCQM)

Figure D-17: Substance Abuse list (These save codes are used by eCQM)

Figure D-18: Tobacco list (Medications on this list do NOT work for eCQM as they do not save in the required RxNORM format)

Figure D-19: Weight Management list
Appendix E  Value Sets

**Hospital Measures:**
- CMS 9v8 Exclusive Breast Milk Feeding
- CMS 26v6 Home Management Plan of Care
- CMS 31v7 Hearing Screen before hospital discharge
- CMS 32v8 Time from Arrival in ER until Departure for Discharged ED patients
- CMS 53v7 Primary PCI Received Within 90 Minutes of Hospital Arrival
- CMS 55v7 Median Time from ED Arrival to ED Departure for Admitted ED Patients
- CMS 71v9 Anticoagulation Therapy for Atrial Fibrillation
- CMS 72v8 Antithrombotic Therapy By End of Hospital Day 2
- CMS 102v7 Assessed for Rehabilitation
- CMS 104v8 Discharged on Antithrombotic Therapy
- CMS 105v8 Discharged on Statin Medication
- CMS 107v7 Stroke Education
- CMS 111v8 Median Admit Decision Time to ED Departure Time for Admitted Patients
- CMS 113v7 Elective Delivery
- CMS 190v8 Intensive Care Unit Venous Thromboembolism Prophylaxis

**Provider Measures:**
- CMS 2v9 Preventive Care and Screening for Depression
- CMS 50v8 Closing the Referral Loop
- CMS 69v8 BMI Preventive Care and Screening
- CMS 117 v8 Childhood Immunization Status
- CMS 122v8 Diabetes HbA1c Poor Control
- CMS 124v8 Cervical Cancer Screening
- CMS 125v8 Breast Cancer Screening
- CMS 127 v8 Pneumococcal Vaccination Status for Older Adults
- CMS 130 v8 Colorectal Cancer Screening
- CMS 131v8 Diabetes Eye Exam
Electronic Clinical Quality Measures (ECQM) Version 3.0

- CMS 134v8 Diabetes Nephropathy
- CMS 137v8 Initiation and Engagement of Alcohol and Other Drug Dependence Treatment
- CMS 138v8 Preventive Care and Screening Tobacco Use
- CMS 139v8 Falls Risk
- CMS 144v8 Heart Failure Beta Blocker Use
- CMS 155v8 Weight Assessment and Counseling for Nutrition and Physical Activity for Children and Adolescents
- CMS 156v8 Use of High Risk Meds in Elderly
- CMS 159v8 Depression Remission at Twelve Months
- CMS 160v8 Depression Utilization of the PHQ-9 Tool
- CMS 161v8 Adult Major Depressive Disorder
- CMS 165v8 Controlling High Blood Pressure
- CMS 177v8 Child and Adolescent Major Depressive Disorder
- CMS 347v3 Statin Therapy for the Prevention and Treatment of Cardiovascular Disease
- CMS 349v2 HIV Screening

**Value Sets: CMS 9v8 Exclusive Breast Feeding**

| Description | Exclusive breast milk feeding during the newborn's entire hospitalization. The measure is reported as an overall rate which includes all newborns that were exclusively fed breast milk during the entire hospitalization. |

- "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)
- valueset "Discharge To Acute Care Facility" using "2.16.840.1.113883.3.117.1.7.1.87"
- valueset "Neonatal Intensive Care Unit (NICU)" using "2.16.840.1.113883.3.117.1.7.1.75"
- valueset "Patient Expired" using "2.16.840.1.113883.3.117.1.7.1.309"
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)
- "Procedure, Performed: Parenteral Nutrition" (2.16.840.1.113883.3.117.1.7.1.38)
• "Substance, Administered: Breast Milk" (2.16.840.1.113883.3.117.1.7.1.30)
• Substance, Administered: Dietary Intake Other than Breast Milk" (2.16.840.1.113883.3.117.1.7.1.27)

Value Sets: CMS 26v6 Home Management Plan of Care (HMPC) Document Given to Patient/Caregiver

<table>
<thead>
<tr>
<th>Description</th>
<th>An assessment that there is documentation in the medical record that a Home Management Plan of Care (HMPC) document was given to the pediatric asthma patient/caregiver</th>
</tr>
</thead>
<tbody>
<tr>
<td>Encounter, Performed: Encounter Inpatient (2.16.840.1.113883.3.666.5.307)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Ethnicity: Ethnicity (2.16.840.1.114222.4.11.837)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Payer: Payer (2.16.840.1.114222.4.11.3591)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Race: Race&quot;(2.16.840.1.114222.4.11.836)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Sex: ONC Administrative Sex (2.16.840.1.113762.1.4.1)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Communication: From Provider To Patient, Not Done: Asthma action plan (LOINC version 2.63 Code 69981-9)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Communication: From Provider To Patient: Asthma action plan (LOINC version 2.63 Code 69981-9)&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Asthma&quot; using &quot;2.16.840.1.113883.3.117.1.7.1.271&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Discharge To Home Or Police Custody&quot; using &quot;2.16.840.1.113883.3.117.1.7.1.82&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Patient Refusal&quot; using &quot;2.16.840.1.113883.3.117.1.7.1.93&quot;</td>
<td></td>
</tr>
</tbody>
</table>

Value Sets: CMS 31v7 Hearing Screening Prior to Hospital Discharge

<table>
<thead>
<tr>
<th>Description</th>
<th>This measure assesses the proportion of births that have been screened for hearing loss before hospital discharge.</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Diagnosis: Live Birth Newborn Born in Hospital&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.113762.1.4.1046.6)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnostic Study, Not Performed: Newborn Hearing Screen Left&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.114222.4.1.214079.1.1.3)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnostic Study, Not Performed: Newborn Hearing Screen Right&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.114222.4.1.214079.1.1.4)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnostic Study, Performed: Newborn Hearing Screen Left&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.114222.4.1.214079.1.1.3)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnostic Study, Performed: Newborn Hearing Screen Right&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.114222.4.1.214079.1.1.4)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Encounter, Performed: Encounter Inpatient&quot;</td>
<td></td>
</tr>
<tr>
<td>(2.16.840.1.113883.3.666.5.307)&quot;</td>
<td></td>
</tr>
</tbody>
</table>
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1.1)"
• valueset "Medical Reasons" using "2.16.840.1.114222.4.1.214079.1.1.7"
• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Patient deceased during stay (discharge status = dead) (finding)" using "SNOMEDCT version 2017-09 Code (371828006)"

### Value Sets: CMS 32v8 Time from Arrival to ED Departure for Discharged ED Patients

<table>
<thead>
<tr>
<th>Description</th>
<th>Medicaid elapsed time from ED arrival to ER departure for patients discharged from the ED.</th>
</tr>
</thead>
</table>
• "Encounter, Order: Observation Services" (2.16.840.1.113762.1.4.1111.143)"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1.1)"
• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Patient deceased during stay (discharge status = dead) (finding)" using "SNOMEDCT version 2017-09 Code (371828006)"
• valueset "Discharge To Acute Care Facility" using "2.16.840.1.113883.3.117.1.7.1.87"

### Value Sets: CMS 53v7 Primary PCI Received Within 90 Minutes of Hospital Arrival

<table>
<thead>
<tr>
<th>Description</th>
<th>AMI patients with a diagnostic ECG and ST-segment elevation, who receive a primary PCI during their hospital stay, and the time from hospital arrival to PCI is 90 minutes or less.</th>
</tr>
</thead>
</table>
• "Diagnosis: Cardiopulmonary arrest" (2.16.840.1.113883.3.666.5.748)"
• "Diagnostic Study, Performed: Electrocardiogram (ECG)" (2.16.840.1.113883.3.666.5.735)"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
- "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
- "Medication, Administered: Fibrinolytic Therapy" (2.16.840.1.113883.3.666.5.736)"
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
- "Procedure, Performed: Aortic balloon pump insertion" (2.16.840.1.113883.3.666.5.1151)"
- "Procedure, Performed: Endotracheal Intubation" (2.16.840.1.113762.1.4.1045.69)"
- SNOMEDCT, ICD10PCS
- "Procedure, Performed: PCI" (2.16.840.1.113762.1.4.1045.67)"
- "Procedure, Performed: Ventricular Assist Device placement" (2.16.840.1.113883.3.666.5.3015)"

valueset "Transfer From Outpatient" using "2.16.840.1.113883.3.67.1.101.950"
valueset "Transfer From Inpatient" using "2.16.840.1.113883.3.666.5.3013"
valueset "Transfer From ED Locations" using "2.16.840.1.113883.3.666.5.3006"
valueset "Acute or Evolving MI" using "2.16.840.1.113883.3.666.5.3022"
valueset "Ambulatory surgical center" using "2.16.840.1.113883.3.666.5.687"

**Value Sets: CMS 55V7 Median Time from ED Arrival to ED Departure for Admitted ED Patients**

<table>
<thead>
<tr>
<th>Description</th>
<th>Median time from ED arrival to time of departure from the ER for patients admitted to the facility from the ED</th>
</tr>
</thead>
</table>
- "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
- "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
- valueset "Hospital Settings" using "2.16.840.1.113762.1.4.1111.126"
• valueset "Psychiatric/Mental Health Diagnosis" using "2.16.840.1.113883.3.117.1.7.1.299"

Value Sets: CMS 71v9 Anticoagulation Therapy for Atrial Fibrillation/Flutter

<table>
<thead>
<tr>
<th>Description</th>
<th>Ischemic stroke patients with atrial fibrillation/flutter who are prescribed or continuing to take anticoagulation therapy at hospital discharge.</th>
</tr>
</thead>
</table>

• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Non-Elective Inpatient Encounter (2.16.840.1.113883.3.117.1.7.1.424)"
• “EncounterPerformed, Observation Services” 2.16.840.1.113762.1.4.1111.143
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Medication, Discharge: Anticoagulant Therapy" (2.16.840.1.113883.3.117.1.7.1.200)"
• "Medication, Not Discharged: Anticoagulant Therapy" (2.16.840.1.113883.3.117.1.7.1.200)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: Atrial Ablation" (2.16.840.1.113883.3.117.1.7.1.203)"
• Diagnosis: Atrial Fibrillation/Flutter" (2.16.840.1.113883.3.117.1.7.1.202)"
• valueset "Discharge To Acute Care Facility" using "2.16.840.1.113883.3.117.1.7.1.87"
• valueset "Discharged to Health Care Facility for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.207"
• valueset "Discharged to Home for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.209"
• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"
• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"
• valueset "Left Against Medical Advice" using "2.16.840.1.113883.3.117.1.7.1.308"
• valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
• valueset "Patient Expired" using "2.16.840.1.113883.3.117.1.7.1.309"
• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

Value Sets: CMS 72v8 Antithrombotic Therapy By End of Hospital Day 2

<table>
<thead>
<tr>
<th>Description</th>
<th>Ischemic stroke patients administered antithrombotic therapy by the end of hospital day 2.</th>
</tr>
</thead>
</table>
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Non-Elective Inpatient Encounter" (2.16.840.1.113883.3.117.1.7.1.424)"
• “Encounter Performed, Observation: 2.16.840.1.113762.1.4.1111.143
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Medication, Administered: Antithrombotic Therapy" (2.16.840.1.113883.3.117.1.7.1.201)"
• "Medication, Administered: Thrombolytic (t-PA) Therapy" (2.16.840.1.113883.3.117.1.7.1.226)"
• "Medication, Not Administered: Antithrombotic Therapy" (2.16.840.1.113883.3.117.1.7.1.201)"
• "Medication, Not Ordered: Antithrombotic Therapy" (2.16.840.1.113883.3.117.1.7.1.201)"
• Lab Test, INR: 2.16.840.1.113883.3.117.1.7.1.213
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: Intravenous or Intra-arterial Thrombolytic (t-PA) Therapy" (2.16.840.1.113762.1.4.1045.21)"
• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"
• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"
• valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

Value Sets: CMS 102v7 Assessed for Rehabilitation

<table>
<thead>
<tr>
<th>Description</th>
<th>Ischemic or hemorrhagic stroke patients who were assessed for rehabilitation services.</th>
</tr>
</thead>
</table>
• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"
• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"
• valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Non-Elective Inpatient Encounter"
  (2.16.840.1.113883.3.117.1.7.1.424)"
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Not Performed: Rehabilitation Assessment"
  (2.16.840.1.113762.1.4.1045.18)"
• "Procedure, Not Performed: Rehabilitation Therapy"
  (2.16.840.1.113762.1.4.1045.19)"
• "Procedure, Performed: Rehabilitation Assessment
  (2.16.840.1.113762.1.4.1045.18)"
• "Procedure, Performed: Rehabilitation Therapy"
  (2.16.840.1.113762.1.4.1045.19)"
• valueset "Discharge To Acute Care Facility" using
  "2.16.840.1.113883.3.117.1.7.1.87"
• valueset "Discharged to Health Care Facility for Hospice Care" using
  "2.16.840.1.113883.3.117.1.7.1.207"
• valueset "Discharged to Home for Hospice Care" using
  "2.16.840.1.113883.3.117.1.7.1.209"
• valueset "Discharged to Rehabilitation Facility" using
  "2.16.840.1.113883.3.117.1.7.1.132"
• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"
• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"
• valueset "Left Against Medical Advice" using
  "2.16.840.1.113883.3.117.1.7.1.308"
• valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
• valueset "Patient Expired" using "2.16.840.1.113883.3.117.1.7.1.309"
• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"
Value Sets: CMS 104v8 Discharged on Antithrombotic Therapy

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ischemic stroke patients prescribed or continuing to take</td>
</tr>
<tr>
<td>antithrombotic therapy at hospital discharge.</td>
</tr>
</tbody>
</table>

- "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
- "Encounter, Performed: Non-Elective Inpatient Encounter" (2.16.840.1.113883.3.117.1.7.1.424)"
- Encounter, Performed: Observation Services: (2.16.840.1.113762.1.4.1.1111.143)
- "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
- "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
- "Medication, Discharge: Antithrombotic Therapy" (2.16.840.1.113883.3.117.1.7.1.201)"
- "Medication, Not Discharged: Antithrombotic Therapy" 2.16.840.1.113883.3.117.1.7.1.201)"
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
- valueset "Discharge To Acute Care Facility" using "2.16.840.1.113883.3.117.1.7.1.87""
- valueset "Discharged to Health Care Facility for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.207""
- valueset "Discharged to Home for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.209""
- valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212""
- valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247""
- valueset "Left Against Medical Advice" using "2.16.840.1.113883.3.117.1.7.1.308""
- valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473""
- valueset "Patient Expired" using "2.16.840.1.113883.3.117.1.7.1.309""
- valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

Value Sets: CMS 105v8 Discharged on Statin Medication

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ischemic stroke patients who are prescribed or continuing to take</td>
</tr>
<tr>
<td>statin medication at hospital discharge.</td>
</tr>
</tbody>
</table>
• "Allergy/Intolerance: Statin Allergen" (2.16.840.1.113883.3.117.1.7.1.423)"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Non-Elective Inpatient Encounter" (2.16.840.1.113883.3.117.1.7.1.424)"
• Encounter Performed: Observation Services (2.16.840.1.113762.1.4.1111.143)
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Laboratory Test, Performed: LDL-C" (2.16.840.1.113883.3.117.1.7.1.215)"
• "Medication, Discharge: Statin Grouper" (2.16.840.1.113762.1.4.1110.19)"
• "Medication, Not Discharged: Statin Grouper" (2.16.840.1.113762.1.4.1110.19)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• valueset "Discharge To Acute Care Facility" using "2.16.840.1.11"3883.3.117.1.7.1.87"
• valueset "Discharged to Health Care Facility for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.207"
• valueset "Discharged to Home for Hospice Care" using "2.16.840.1.113883.3.117.1.7.1.209"
• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"
• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"
• valueset "Left Against Medical Advice" using "2.16.840.1.113883.3.117.1.7.1.308"
• valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
• valueset "Patient Expired" using "2.16.840.1.113883.3.117.1.7.1.309"
• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

Value Set: CMS 107v7 Stroke Education

| Description | Ischemic or hemorrhagic stroke patients or their caregivers who were given educational materials during the hospital stay addressing all of the following: activation of emergency medical system, need for follow-up after discharge, medications prescribed at discharge, risk factors for stroke, and warning signs and symptoms of stroke. |
• "Communication: From Provider To Patient, Not Done: Written Information Given" (2.16.840.1.113883.3.117.1.7.1.415)"

• "Communication: From Provider To Patient: Activation of Emergency Medical System Education" (2.16.840.1.113883.3.117.1.7.1.377)"

• "Communication: From Provider To Patient: Instructions for Follow Up After Discharge" (2.16.840.1.113883.3.117.1.7.1.378)"

• "Communication: From Provider To Patient: Prescribed Medications Education" (2.16.840.1.113883.3.117.1.7.1.379)"

• "Communication: From Provider To Patient: Risk Factors Education" (2.16.840.1.113883.3.117.1.7.1.380)"

• "Communication: From Provider To Patient: Warning Signs and Symptoms Education" (2.16.840.1.113883.3.117.1.7.1.381)"

• "Communication: From Provider To Patient: Written Information Given" (2.16.840.1.113883.3.117.1.7.1.415)"

• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"

• "Encounter, Performed: Non-Elective Inpatient Encounter" (2.16.840.1.113883.3.117.1.7.1.424)"

• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"

• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"

• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)

• "Patient Characteristic Payer: Payer (2.16.840.1.114222.4.11.3591)"

• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"

• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"

• valueset "Discharge To Home Or Police Custody" using "2.16.840.1.113883.3.117.1.7.1.82"

• valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"

• valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"

• valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

**Value Sets: CMS 108v8 Venous Thromboembolism Prophylaxis**

| Description | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after hospital admission or surgery end date for surgeries that start the day of or the day after hospital admission. |
• "Device, Applied: Graduated compression stockings (GCS)"
  (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Applied: Intermittent pneumatic compression devices (IPC)"
  (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Applied: Venous foot pumps (VFP)"
  (2.16.840.1.113883.3.117.1.7.1.230)"
• "Device, Not Applied: Graduated compression stockings (GCS)"
  (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Not Applied: Intermittent pneumatic compression devices (IPC)"
  (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Not Applied: Venous foot pumps (VFP)"
  (2.16.840.1.113883.3.117.1.7.1.230)"
• "Device, Not Ordered: Graduated compression stockings (GCS)"
  (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Not Ordered: Intermittent pneumatic compression devices (IPC)"
  (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Not Ordered: Venous foot pumps (VFP)"
  (2.16.840.1.113883.3.117.1.7.1.230)"
• "Diagnosis: Atrial Fibrillation/Flutter" (2.16.840.1.113883.3.117.1.7.1.202)"
• "Diagnosis: Obstetrics VTE" (2.16.840.1.113883.3.117.1.7.1.264)"
• "Diagnosis: Venous Thromboembolism" (2.16.840.1.113883.3.117.1.7.1.279)"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
• “Encounter Performed: Observation Services” (2.16.840.1.113762.1.4.1111.143)
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Laboratory Test, Performed: INR" (2.16.840.1.113883.3.117.1.7.1.213)"
• "Medication, Administered: Direct Thrombin Inhibitor"
  (2.16.840.1.113883.3.117.1.7.1.205)"
• "Medication, Administered: Glycoprotein IIb/IIIa Inhibitors" using "Glycoprotein IIb/IIIa Inhibitors" (2.16.840.1.113762.1.4.1045.41)"
• "Medication, Administered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)"
• "Medication, Administered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)"
- "Medication, Administered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)
- "Medication, Administered: Oral Factor Xa Inhibitor for VTE Prophylaxis or VTE Treatment" (2.16.840.1.113883.3.117.1.7.1.134)
- "Medication, Administered: Unfractionated Heparin" (2.16.840.1.113883.3.117.1.7.1.218)
- "Medication, Administered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)
- "Medication, Not Administered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)
- "Medication, Not Administered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)
- "Medication, Not Administered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)
- "Medication, Not Administered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)
- "Medication, Not Ordered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)
- "Medication, Not Ordered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)
- "Medication, Not Ordered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)
- "Medication, Not Ordered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1.1)
- "Procedure, Performed: General or Neuraxial Anesthesia" (2.16.840.1.113883.3.666.5.1743)
- "Procedure, Performed: General Surgery" (2.16.840.1.113883.3.117.1.7.1.255)
- "Procedure, Performed: Gynecological Surgery" (2.16.840.1.113883.3.117.1.7.1.257)
- "Procedure, Performed: Hip Fracture Surgery" (2.16.840.1.113883.3.117.1.7.1.258)
- "Procedure, Performed: Hip Replacement Surgery" (2.16.840.1.113883.3.117.1.7.1.259)
"Procedure, Performed: Intracranial Neurosurgery" (2.16.840.1.113883.3.117.1.7.1.260)"

"Procedure, Performed: Knee Replacement Surgery" (2.16.840.1.113883.3.117.1.7.1.261)"

"Procedure, Performed: Urological Surgery" (2.16.840.1.113883.3.117.1.7.1.272)"

"Assessment, Performed: Risk for venous thromboembolism" (LOINC version 2.63 Code 72136-5)"

codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"

valueset "Hemorrhagic Stroke" using "2.16.840.1.113883.3.117.1.7.1.212"

valueset "Intensive Care Unit" using "2.16.840.1.113762.1.4.1110.23"

valueset "Intravenous route" using "2.16.840.1.113883.3.117.1.7.1.222"

valueset "Ischemic Stroke" using "2.16.840.1.113883.3.117.1.7.1.247"

valueset "Low Risk" using "2.16.840.1.113883.3.117.1.7.1.400"

valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"

valueset "Mental Health Diagnoses" using "2.16.840.1.113883.3.464.1003.105.12.1004"

valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"

valueset "Principal" using "2.16.840.1.113883.3.117.1.7.1.14"

valueset "Subcutaneous route" using "2.16.840.1.113883.3.117.1.7.1.223"

**Value Sets: CMS 111V8 Median Admit Decision Time to ED Departure Time for Admitted Patients**

<table>
<thead>
<tr>
<th>Description:</th>
<th>Median time (in minutes) from admit decision time to time of departure from the ED for ED patients admitted to inpatient status.</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Encounter, Order: Decision to Admit to Hospital Inpatient&quot; (2.16.840.1.113883.3.117.1.7.1.295)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Encounter, Performed: ED Visit&quot;  (2.16.840.1.113883.3.117.1.7.1.292)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Encounter, Performed: Encounter Inpatient&quot; (2.16.840.1.113883.3.666.5.307)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Ethnicity: Ethnicity&quot; (2.16.840.1.114222.4.114222.4.11.837)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Payer: Payer&quot; (2.16.840.1.114222.4.11.3591)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Race: Race&quot; (2.16.840.1.114222.4.11.836)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Patient Characteristic Sex: ONC Administrative Sex&quot; (2.16.840.1.113762.1.4.1)&quot;</td>
<td></td>
</tr>
</tbody>
</table>
- valueset "Hospital Settings" using "2.16.840.1.113762.1.4.1111.126"
- valueset "Psychiatric/Mental Health Diagnosis" using "2.16.840.1.113883.3.117.1.7.1.299"

**Value Sets: CMS 113v7 Elective Delivery**

<table>
<thead>
<tr>
<th>Description</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patients with elective vaginal deliveries or elective cesarean births at &gt;= 37 and &lt; 39 weeks of gestation completed.</strong></td>
<td></td>
</tr>
</tbody>
</table>

- "Assessment, Performed: EGA at Delivery" (2.16.840.1.113762.1.4.1045.26)"
- "Assessment, Performed: Labor" (2.16.840.1.113883.3.117.1.7.1.281)"
- "Assessment, Performed: Time of Delivery" (2.16.840.1.113762.1.4.1045.28)"
- "Diagnosis: Conditions Possibly Justifying Elective Delivery Prior to 39 Weeks Gestation" (2.16.840.1.113883.3.117.1.7.1.286)"
- "Diagnosis: Cornual Ectopic Pregnancy" (2.16.840.1.113762.1.4.1110.12)"
- "Diagnosis: Perforation of Uterus" (2.16.840.1.113762.1.4.1110.14)"
- "Diagnosis: Uterine Rupture" using (2.16.840.1.113762.1.4.1110.16)"
- "Diagnosis: Uterine Window" (2.16.840.1.113883.3.117.1.7.1.137)"
- "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
- "Medication, Administered: Dinoprostone" (2.16.840.1.113762.1.4.1045.56)"
- "Medication, Administered: Oxytocin" (2.16.840.1.113762.1.4.1045.55)"
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
- "Procedure, Performed: Artificial Rupture of Membranes" (2.16.840.1.113762.1.4.1045.57)"
- "Procedure, Performed: Cesarean Birth" (2.16.840.1.113883.3.117.1.7.1.282)"
- "Procedure, Performed: Classical Cesarean Birth" (2.16.840.1.113883.3.117.1.7.1.421)"
- "Procedure, Performed: Delivery Procedures" (2.16.840.1.113762.1.4.1045.59)"
- "Procedure, Performed: Medical Induction of Labor" (2.16.840.1.113883.3.117.1.7.1.288)"
- "Procedure, Performed: Metroplasty" (2.16.840.1.113762.1.4.1110.25)"
- "Procedure, Performed: Myomectomy" (2.16.840.1.113883.3.117.1.7.1.422)"
• "Procedure, Performed: Transabdominal Cerclage" (2.16.840.1.113762.1.4.1110.18)"
• "Procedure, Performed: Uterine Horn" (2.16.840.1.113762.1.4.1110.24)"

Value Sets: CMS 190v8 Intensive Care Unit Venous Thromboembolism Prophylaxis

<table>
<thead>
<tr>
<th>Description</th>
<th>This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after the initial admission (or transfer) to the ICU or surgery end date for surgeries that start the day of or the day after ICU admission (or transfer).</th>
</tr>
</thead>
</table>

• "Device, Applied: Graduated compression stockings (GCS)" (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Applied: Intermittent pneumatic compression devices (IPC)" (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Applied: Venous foot pumps (VFP)" (2.16.840.1.113883.3.117.1.7.1.230)"
• "Device, Not Applied: Graduated compression stockings (GCS)" (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Not Applied: Intermittent pneumatic compression devices (IPC)" (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Not Applied: Venous foot pumps (VFP)" (2.16.840.1.113883.3.117.1.7.1.230)"
• "Device, Not Ordered: Graduated compression stockings (GCS)" (2.16.840.1.113883.3.117.1.7.1.256)"
• "Device, Not Ordered: Intermittent pneumatic compression devices (IPC)" (2.16.840.1.113883.3.117.1.7.1.214)"
• "Device, Not Ordered: Venous foot pumps (VFP)" (2.16.840.1.113883.3.117.1.7.1.230)"
• "Diagnosis: Atrial Fibrillation/Flutter" (2.16.840.1.113883.3.117.1.7.1.202)"
• "Diagnosis: Venous Thromboembolism" (2.16.840.1.113883.3.117.1.7.1.279)"
• "Encounter, Performed: ED Visit" (2.16.840.1.113883.3.117.1.7.1.292)"
• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
• “Encounter Performed: Observation Services” (2.16.840.1.113762.1.4.1111.143)
• "Intervention, Order: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Intervention, Performed: Comfort Measures" (1.3.6.1.4.1.33895.1.3.0.45)"
• "Laboratory Test, Performed: INR" (2.16.840.1.113883.3.117.1.7.1.213)"
• "Medication, Administered: Direct Thrombin Inhibitor" (2.16.840.1.113883.3.117.1.7.1.205)"
• "Medication, Administered: Glycoprotein IIb/IIIa Inhibitors" (2.16.840.1.113762.1.4.1045.41)"
• "Medication, Administered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)"
• "Medication, Administered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)"
• "Medication, Administered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)"
• "Medication, Administered: Oral Factor Xa Inhibitor for VTE Prophylaxis or VTE Treatment" (2.16.840.1.113883.3.117.1.7.1.134)"
• "Medication, Administered: Unfractionated Heparin" (2.16.840.1.113883.3.117.1.7.1.218)"
• "Medication, Administered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)"
• "Medication, Not Administered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)"
• "Medication, Not Administered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)"
• "Medication, Not Administered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)"
• "Medication, Not Administered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)"
• "Medication, Not Ordered: Injectable Factor Xa Inhibitor for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.211)"
• "Medication, Not Ordered: Low Dose Unfractionated Heparin for VTE Prophylaxis" (2.16.840.1.113762.1.4.1045.39)"
• "Medication, Not Ordered: Low Molecular Weight Heparin for VTE Prophylaxis" (2.16.840.1.113883.3.117.1.7.1.219)"
• "Medication, Not Ordered: Warfarin" (2.16.840.1.113883.3.117.1.7.1.232)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: General or Neuraxial Anesthesia" (2.16.840.1.113883.3.666.5.1743)"
"Procedure, Performed: General Surgery" (2.16.840.1.113883.3.117.1.7.1.255)
"Procedure, Performed: Gynecological Surgery" (2.16.840.1.113883.3.117.1.7.1.257)
"Procedure, Performed: Hip Fracture Surgery" (2.16.840.1.113883.3.117.1.7.1.258)
"Procedure, Performed: Hip Replacement Surgery" (2.16.840.1.113883.3.117.1.7.1.259)
"Procedure, Performed: Intracranial Neurosurgery" (2.16.840.1.113883.3.117.1.7.1.260)
"Procedure, Performed: Knee Replacement Surgery" (2.16.840.1.113883.3.117.1.7.1.261)
"Procedure, Performed: Urological Surgery" (2.16.840.1.113883.3.117.1.7.1.272)
"Assessment, Performed: Risk for venous thromboembolism" (LOINC version 2.63 Code 72136-5)
codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
code "Risk for venous thromboembolism" using "LOINC version 2.63 Code (72136-5)"
valueset "Intensive Care Unit" using "2.16.840.1.113762.1.4.1110.23"
valueset "Intravenous route" using "2.16.840.1.113883.3.117.1.7.1.222"
valueset "Low Risk" using "2.16.840.1.113883.3.117.1.7.1.400"
valueset "Medical Reason" using "2.16.840.1.113883.3.117.1.7.1.473"
valueset "Patient Refusal" using "2.16.840.1.113883.3.117.1.7.1.93"
valueset "Principal" using "2.16.840.1.113883.3.117.1.7.1.14"
valueset "Subcutaneous route" using "2.16.840.1.113883.3.117.1.7.1.223"

Value Sets: CMS 2v9 Preventive Care and Screening for Depression

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage of patients aged 12 years and older screened for depression on the date of the encounter using an age appropriate standardized depression screening tool AND if positive, a follow-up plan is documented on the date of the positive screen.</td>
</tr>
</tbody>
</table>

"Diagnosis: Bipolar Diagnosis" (2.16.840.1.113883.3.600.450)"
"Diagnosis: Depression diagnosis" (2.16.840.1.113883.3.600.145)"
"Encounter, Performed: Depression Screening Encounter Codes" (2.16.840.1.113883.3.600.1916)"
• "Intervention, Order: Referral for Depression Adolescent" (2.16.840.1.113883.3.600.537)"
• "Intervention, Order: Referral for Depression Adult" (2.16.840.1.113883.3.600.538)"
• "Intervention, Performed: Additional evaluation for depression - adolescent" (2.16.840.1.113883.3.600.1542)"SNOMEDCT"
• "Intervention, Performed: Additional evaluation for depression - adult" (2.16.840.1.113883.3.600.1545)"
• "Intervention, Performed: Follow-up for depression - adolescent" (2.16.840.1.113883.3.600.467)"
• "Intervention, Performed: Follow-up for depression - adult" (2.16.840.1.113883.3.600.468)"
• "Assessment, Performed: Suicide Risk Assessment" (2.16.840.1.113883.3.600.559)"
• "Medication, Order: Depression medications - adolescent" (2.16.840.1.113883.3.600.469)"
• "Medication, Order: Depression medications - adult" (2.16.840.1.113883.3.600.470)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Assessment, Not Performed: Adolescent depression screening assessment" (LOINC version 2.63 Code 73831-0)"
• "Assessment, Not Performed: Adult depression screening assessment" (LOINC version 2.63 Code 73832-8)"
• "Assessment, Performed: Adolescent depression screening assessment" (LOINC version 2.63 Code 73831-0)"
• "Assessment, Performed: Adult depression screening assessment" (LOINC version 2.63 Code 73832-8)"
• codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
• valueset "Medical or Other reason not done" using "2.16.840.1.113883.3.600.1.1502"
• valueset "Negative Depression Screening" using "2.16.840.1.113883.3.600.2451"
• valueset "Patient Reason refused" using "2.16.840.1.113883.3.600.791"
• valueset "Positive Depression Screening" using "2.16.840.1.113883.3.600.2450"

**Value Sets: CMS 50v7 Closing the Referral Loop: Receipt of Specialist Report**

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients with referrals, regardless of age, for which the referring provider receives a report from the provider to whom the patient was referred.</th>
</tr>
</thead>
</table>

• "Communication: From Provider To Provider: Consultant Report" (2.16.840.1.113883.3.464.1003.121.12.1006)"

• "Encounter, Performed: Office Visit" (2.16.840.1.113883.3.464.1003.101.12.1001)"

• "Encounter, Performed: Ophthalmological Services" (2.16.840.1.113883.3.526.3.1285)"

• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1025)"

• "Encounter, Performed: Preventive Care Services, Initial Office Visit, 0 to 17" (2.16.840.1.113883.3.464.1003.101.12.1022)"

• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1023)"

• "Encounter, Performed: Preventive Care, Established Office Visit, 0 to 17" (2.16.840.1.113883.3.464.1003.101.12.1024)"

• "Intervention, Performed: Referral" (2.16.840.1.113883.3.464.1003.101.12.1046)"

• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"

• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"

• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"

• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"

**Value Sets: CMS 69v8, BMI Preventive Care and Screening**

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients aged 18 years and older with a BMI documented during the current encounter or during the previous 12 months AND with a BMI outside of normal parameters, a follow-up plan is documented during the encounter or during the previous 12 months of the current encounter. Normal Parameters: Age 18 years and older BMI =&gt; 18.5 and &lt; 25 kg/m2.</th>
</tr>
</thead>
</table>

• "Diagnosis: Pregnancy Dx" (2.16.840.1.113883.3.600.1.1623)"
• "Encounter, Performed: BMI Encounter Code Set" 
  (2.16.840.1.113883.3.600.1.1751)"

• "Encounter, Performed: Palliative care encounter" 
  (2.16.840.1.113883.3.600.1.1575)"

• "Intervention, Not Ordered: Above Normal Follow-up" 
  (2.16.840.1.113883.3.600.1.1525)"

• "Intervention, Not Ordered: Below Normal Follow up" 
  (2.16.840.1.113883.3.600.1.1528)"

• "Intervention, Not Ordered: Referrals where weight assessment may occur" 
  (2.16.840.1.113883.3.600.1.1527)"

• “Intervention, Not Performed: Below Normal Follow up: 
  (2.16.840.1.113883.3.600.1.1528)"

• “Intervention, Not Performed: Above Normal Follow up: 
  (2.16.840.1.113883.3.600.1.1525)"

• "Intervention, Order: Above Normal Follow-up" 
  (2.16.840.1.113883.3.600.1.1525)"

• "Intervention, Order: Below Normal Follow up" 
  (2.16.840.1.113883.3.600.1.1528)"

• "Intervention, Order: Palliative or Hospice Care" 
  (2.16.840.1.113883.3.600.1.1579)"

• "Intervention, Order: Referrals where weight assessment may occur" 
  (2.16.840.1.113883.3.600.1.1527)"

• Intervention, Performed: Below Normal Follow up: 
  (2.16.840.1.113883.3.600.1.1528)"

• Intervention, Performed: Above Normal Follow-up: 
  (2.16.840.1.113883.3.600.1.1525)"

• "Medication, Not Ordered: Above Normal Medications" 
  (2.16.840.1.113883.3.600.1.1498)"

• "Medication, Not Ordered: Below Normal Medications" 
  (2.16.840.1.113883.3.600.1.1499)"

• "Medication, Order: Above Normal Medications" 
  (2.16.840.1.113883.3.600.1.1498)"

• "Medication, Order: Below Normal Medications" 
  (2.16.840.1.113883.3.600.1.1499)"

• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"

• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Physical Exam, Not Performed: BMI [Ratio]" using "BMI [Ratio] (LOINC version 2.63 Code 39156-5)"
• "Physical Exam, Performed: BMI [Ratio]" using "BMI [Ratio] (LOINC version 2.63 Code 39156-5)"
• codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
• code "BMI [Ratio]" using "LOINC version 2.63 Code (39156-5)"
• valueset "Medical or Other reason not done" using "2.16.840.1.113883.3.600.1.1502"
• valueset "Overweight or Obese" using "2.16.840.1.113762.1.4.1047.502"
• valueset "Patient Reason refused" using "2.16.840.1.113883.3.600.791"
• valueset "Underweight" using "2.16.840.1.113883.3.600.2388"

Value Sets: CMS 117 V8 Childhood Immunization Status

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of children 2 years of age who had four DTaP; three polio (IPV), one MMR; three H influenza type B (HiB); three hepatitis B (Hep B); one chicken pox (VZV); four pneumococcal conjugate (PCV); one hepatitis A (Hep A); two or three rotavirus (RV); and two influenza (flu) vaccines by their second birthday.</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Diagnosis: Anaphylactic Reaction to Common Baker's Yeast&quot; (2.16.840.1.113883.3.464.1003.199.12.1032)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Anaphylactic Reaction to DTaP Vaccine&quot; (2.16.840.1.113883.3.464.1003.199.12.1031)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Anaphylactic Reaction to Hepatitis A Vaccine&quot; (2.16.840.1.113883.3.464.1003.199.12.1026)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Disorders of the Immune System&quot; (2.16.840.1.113883.3.464.1003.120.12.1001)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Encephalopathy due to Childhood Vaccination&quot; (2.16.840.1.113883.3.464.1003.114.12.1007)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Hepatitis A&quot; (2.16.840.1.113883.3.464.1003.110.12.1024)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Hepatitis B&quot; (2.16.840.1.113883.3.464.1003.110.12.1025)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: HIV&quot; (2.16.840.1.113883.3.464.1003.120.12.1003)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Intussusception&quot; (2.16.840.1.113883.3.464.1003.199.12.1056)&quot;</td>
<td></td>
</tr>
<tr>
<td>&quot;Diagnosis: Malignant Neoplasm of Lymphatic and Hematopoietic Tissue&quot; (2.16.840.1.113883.3.464.1003.108.12.1009)&quot;</td>
<td></td>
</tr>
</tbody>
</table>
• "Diagnosis: Measles" (2.16.840.1.113883.3.464.1003.110.12.1053)"
• "Diagnosis: Mumps" (2.16.840.1.113883.3.464.1003.110.12.1032)"
• "Diagnosis: Rubella" (2.16.840.1.113883.3.464.1003.110.12.1037)"
• "Diagnosis: Severe Combined Immunodeficiency"
  (2.16.840.1.113883.3.464.1003.120.12.1007)"
• "Diagnosis: Varicella Zoster" (2.16.840.1.113883.3.464.1003.110.12.1039)"
• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
• "Encounter, Performed: Home Healthcare Services"
  (2.16.840.1.113883.3.464.1003.101.12.1016)"
• "Encounter, Performed: Office Visit"
  (2.16.840.1.113883.3.464.1003.101.12.1001)"
• "Encounter, Performed: Preventive Care Services, Initial Office Visit, 0 to 17"
  (2.16.840.1.113883.3.464.1003.101.12.1022)"
• "Encounter, Performed: Preventive Care, Established Office Visit, 0 to 17"
  (2.16.840.1.113883.3.464.1003.101.12.1024)"
• "Immunization, Administered: DTaP Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1214)"
• "Immunization, Administered: Haemophilus Influenzae Type B (HiB) Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1217)"
• "Immunization, Administered: Hepatitis A Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1215)"
• "Immunization, Administered: Hepatitis B Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1216)"
• "Immunization, Administered: Inactivated Polio Vaccine (IPV)"
  (2.16.840.1.113883.3.464.1003.196.12.1219)"
• "Immunization, Administered: Influenza Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1218)"
• "Immunization, Administered: MMR Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1224)"
• "Immunization, Administered: Pneumococcal Conjugate Vaccine"
  (2.16.840.1.113883.3.464.1003.196.12.1221)"
• "Immunization, Administered: Rotavirus Vaccine (3 dose schedule)"
  (2.16.840.1.113883.3.464.1003.196.12.1223)"
• "Immunization, Administered: Varicella Zoster Vaccine (VZV)"
  (2.16.840.1.113883.3.464.1003.196.12.1170)"
• "Intervention, Order: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)"
• "Intervention, Performed: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)"
• "Laboratory Test, Performed: Anti Hepatitis A IgG Antigen Test" (2.16.840.1.113762.3.464.1003.198.12.1033)"
• "Laboratory Test, Performed: Anti Hepatitis B Virus Surface Ab" (2.16.840.1.113762.3.464.1003.198.12.1073)"
• "Laboratory Test, Performed: Measles Antibody Test (IgG Antibody presence)" (2.16.840.1.113883.3.464.1003.198.12.1060)"
• "Laboratory Test, Performed: Measles Antibody Test (IgG Antibody Titer)" (2.16.840.1.113883.3.464.1003.198.12.1059)"
• "Laboratory Test, Performed: Mumps Antibody Test (IgG Antibody presence)" (2.16.840.1.113883.3.464.1003.198.12.1062)"
• "Laboratory Test, Performed: Mumps Antibody Test (IgG Antibody Titer)" (2.16.840.1.113883.3.464.1003.198.12.1061)"
• "Laboratory Test, Performed: Rubella Antibody Test (IgG Antibody presence)" (2.16.840.1.113883.3.464.1003.198.12.1064)"
• "Laboratory Test, Performed: Rubella Antibody Test (IgG Antibody Titer)" (2.16.840.1.113883.3.464.1003.198.12.1063)"
• "Laboratory Test, Performed: Varicella Zoster Antibody Test (IgG Antibody Presence)" (2.16.840.1.113883.3.464.1003.198.12.1067)"
• "Laboratory Test, Performed: Varicella Zoster Antibody Test (IgG Antibody Titer)" (2.16.840.1.113883.3.464.1003.198.12.1066)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: DTaP Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1022)"
• "Procedure, Performed: Hepatitis A Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1041)"
• "Procedure, Performed: Hepatitis B Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1042)"
• "Procedure, Performed: Inactivated Polio Vaccine (IPV) Administered" (2.16.840.1.113883.3.464.1003.110.12.1045)"
• "Procedure, Performed: Influenza Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1044)"
• "Procedure, Performed: MMR Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1031)"
• "Procedure, Performed: Pneumococcal Conjugate Vaccine Administered" (2.16.840.1.113883.3.464.1003.110.12.1046)"
• "Procedure, Performed: Varicella Zoster Vaccine (VZV) Administered" (2.16.840.1.113883.3.464.1003.110.12.1040)"
• "Diagnosis: Anaphylaxis due to Haemophilus influenzae type b vaccine (disorder)" (SNOMEDCT version 2017-09 Code 433621000124101)"
• "Diagnosis: Anaphylaxis due to Hepatitis B vaccine (disorder)" (SNOMEDCT version 2017-09 Code 428321000124101)"
• "Diagnosis: Anaphylaxis due to rotavirus vaccine (disorder)" (SNOMEDCT version 2017-09 Code 428331000124103)"
• "Diagnosis: Influenza virus vaccine adverse reaction (disorder)" SNOMEDCT version 2017-09 Code 420113004)"
• "Diagnosis: Neomycin adverse reaction (disorder)" (SNOMEDCT version 2017-09 Code 292927007)"
• "Diagnosis: Pneumococcal vaccine adverse reaction (disorder)" (SNOMEDCT version 2017-09 Code 293116002)"
• "Diagnosis: Poliomyelitis vaccine adverse reaction (disorder)" (SNOMEDCT version 2017-09 Code 293117006)"
• "Diagnosis: Polymyxin B adverse reaction (disorder)" (SNOMEDCT version 2017-09 Code 292992006)"
• "Diagnosis: Streptomycin adverse reaction (disorder)" (SNOMEDCT version 2017-09 Code 292925004)"
• "Immunization, Administered: rotavirus, live, monovalent vaccine" (CVX version 2017-11 Code 119)"
• "Procedure, Performed: Rotavirus vaccine, human, attenuated (RV1), 2 dose schedule, live, for oral use" use (CPT version 2018 Code 90681)"
• "Procedure, Performed: Rotavirus vaccine, pentavalent (RV5), 3 dose schedule, live, for oral use" (CPT version 2018 Code 90680)"
• codesystem "CPT" using "2.16.840.1.113883.6.12 version 2018"
• codesystem "CVX" using "2.16.840.1.113883.12.292 version 2017-11"
• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMED CT version 2017-09 Code (428371000124100)"

• code "Discharge to home for hospice care (procedure)" using "SNOMED CT version 2017-09 Code (428361000124107)"

• valueset "Positive Finding" using "2.16.840.1.113883.3.464.1003.121.12.1016"

**Value Sets: CMS 122V8 Diabetes: Hemoglobin A1c (HbA1c) Poor Control (> 9%)**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 18-75 years of age with diabetes who had hemoglobin A1c &gt; 9.0% during the measurement period.</th>
</tr>
</thead>
</table>

• "Diagnosis: Diabetes" (2.16.840.1.113883.3.464.1003.103.12.1001)"

• "Encounter, Performed: Annual Wellness Visit" (2.16.840.1.113883.3.526.3.1240)"

• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"

• "Encounter, Performed: Home Healthcare Services" (2.16.840.1.113883.3.464.1003.101.12.1016)"

• "Encounter, Performed: Office Visit" (2.16.840.1.113883.3.464.1003.101.12.1001)"

• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1025)"

• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1023)"

• “Encounter, Performed: Care Services in Long-Term Residential Facility: (2.16.840.1.113883.3.464.1003.101.12.1014)

• “Encounter, Performed: Nursing Facility Visit: (2.16.840.1.113883.3.464.1003.101.12.1012)

• "Intervention, Order: Hospice care ambulatory" (2.16.840.1.113762.1.4.1.1108.15)"

• "Intervention, Performed: Hospice care ambulatory" (2.16.840.1.113762.1.4.1.1108.15)"

• "Laboratory Test, Performed: HbA1c Laboratory Test" (2.16.840.1.113883.3.464.1003.198.12.1013)"

• "Patient Characteristic Ethnicity: Ethnicity" (16.840.1.114222.4.11.837)"

• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"

• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"

• Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"

• codesystem "SNOMED CT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"

• code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"

Value Sets: CMS 124V8 Cervical Cancer Screening

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of women 21-64 years of age who were screened for cervical cancer using either of the following criteria: Women age 21-64 who had cervical cytology performed every 3 years. Women age 30-64 who had cervical cytology/HPV co-testing performed every 5 years.</th>
</tr>
</thead>
</table>

• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"
• "Encounter, Performed: Home Healthcare Services" (2.16.840.1.113883.3.464.1003.101.12.1016)"
• "Encounter, Performed: Office Visit" (2.16.840.1.113883.3.464.1003.101.12.1001)"
• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1025)"
• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1023)"
• "Intervention, Order: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)"
• "Intervention, Performed: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)"
• "Laboratory Test, Performed: HPV Test" (2.16.840.1.113883.3.464.1003.110.12.1059)"
• "Laboratory Test, Performed: Pap Test" (2.16.840.1.113883.3.464.1003.108.12.1017)"
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: Female" (2.16.840.1.113883.3.560.100.2)"
• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: Hysterectomy with No Residual Cervix" Cervix (2.16.840.1.113883.3.464.1003.198.12.1014)"
• "Diagnosis: Congenital absence of cervix (disorder)" using "Congenital absence of cervix (disorder) (SNOMEDCT version 2017-09 Code 37687000)"
- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"

**Value Sets:** CMS 125V7 Breast Cancer Screening

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of women 50-74 years of age who had a mammogram to screen for breast cancer</th>
</tr>
</thead>
</table>

- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
- valueset "Annual Wellness Visit" "2.16.840.1.113883.3.526.3.1240"
- valueset "Bilateral Mastectomy" "2.16.840.1.113883.3.464.1003.198.12.1005"
- valueset "Encounter Inpatient" "2.16.840.1.113883.3.666.5.307"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Female" using "2.16.840.1.113883.3.560.100.2"
- valueset "History of bilateral mastectomy" "2.16.840.1.113883.3.464.1003.198.12.1068"
- valueset "Home Healthcare Services" "2.16.840.1.113883.3.464.1003.101.12.1016"
- valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15"
- valueset "Left" using "2.16.840.1.113883.3.464.1003.122.12.1036"
- valueset "Mammography" using "2.16.840.1.113883.3.464.1003.108.12.1018"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113883.3.464.1003.12.1025"
- valueset "Payer" using "2.16.840.1.114222.4.11.3591"
- valueset "Preventive Care Services - Established Office Visit, 18 and Up" "2.16.840.1.113883.3.464.1003.101.12.1025"
- valueset "Preventive Care Services-Initial Office Visit, 18 and Up" "2.16.840.1.113883.3.464.1003.101.12.1023"
- valueset "Race" "2.16.840.1.114222.4.11.836"
- valueset "Right" "2.16.840.1.113883.3.464.1003.122.12.1035"
- valueset "Status Post Left Mastectomy" "2.16.840.1.113883.3.464.1003.198.12.1069"
- valueset "Status Post Right Mastectomy" "2.16.840.1.113883.3.464.1003.198.12.1070"
- valueset "Unilateral Mastectomy" "2.16.840.1.113883.3.464.1003.198.12.1020"
- valueset "Unilateral Mastectomy, Unspecified Laterality" (2.16.840.1.113883.3.464.1003.198.12.1068)
- "Diagnosis: Status Post Left Mastectomy" (2.16.840.1.113883.3.464.1003.198.12.1069)
- "Diagnosis: Status Post Right Mastectomy" (2.16.840.1.113883.3.464.1003.198.12.1070)
- "Diagnosis: Unilateral Mastectomy, Unspecified Laterality" (2.16.840.1.113883.3.464.1003.198.12.1071)
- "Diagnostic Study, Performed: Mammography" (2.16.840.1.113883.3.464.1003.108.12.1018)
- "Encounter, Performed: Annual Wellness Visit" (2.16.840.1.113883.3.526.3.1240)
- "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)
- "Encounter, Performed: Home Healthcare Services" (2.16.840.1.113883.3.464.1003.101.12.1016)
- "Encounter, Performed: Office Visit" (2.16.840.1.113883.3.464.1003.101.12.1001)
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1025)
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1023)
- “Encounter, Performed: Care Services in Long-Term Residential Facility:” (2.16.840.1.1.113883.3.464.1003.101.12.1014)
- “Encounter, Performed: Nursing Facility Visit:” (2.16.840.1.1.113883.3.464.1003.101.12.1012)
- "Intervention, Order: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)
- "Intervention, Performed: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)
- "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)
- "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: Female" Female (2.16.840.1.113883.3.560.100.2)"
- "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"
- "Procedure, Performed: Bilateral Mastectomy" (2.16.840.1.113883.3.464.1003.198.12.1005)"
- "Procedure, Performed: Unilateral Mastectomy" (2.16.840.1.113883.3.464.1003.198.12.1020)"
- Procedure, Performed: Unilateral Mastectomy, unspecified"
  2.16.840.1.113883.3.464.1003.198.12.1071"

**Value Sets: CMS 127V7 Pneumococcal Vaccination Status for Older Adults**

<table>
<thead>
<tr>
<th>Description</th>
<th></th>
<th>Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine.</th>
</tr>
</thead>
</table>
- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09" |
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)" |
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)" |
- valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240" |
- valueset "Care Services in Long-Term Residential Facility" using "2.16.840.1.113883.3.464.1003.101.12.1014" |
- valueset "Discharge Services - Nursing Facility" using "2.16.840.1.113883.3.464.1003.101.11.1065" |
- valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307" |
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837" |
- valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016" |
- valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15" |
- valueset "Nursing Facility Visit" using "2.16.840.1.113883.3.464.1003.101.12.1012" |
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001" |
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1" |
- valueset "Payer" using "2.16.840.1.114222.4.11.3591" |
• valueset "Pneumococcal Vaccine Administered" using "2.16.840.1.113883.3.464.1003.110.12.1034"

• valueset "Pneumococcal Vaccine" using "2.16.840.1.113883.3.464.1003.110.12.1027"

• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"

• valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"

• valueset "Race" using "2.16.840.1.114222.4.11.836"

• "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883.3.526.3.1240)"

• "Encounter, Performed: Care Services in Long-Term Residential Facility" using (2.16.840.1.113883.3.464.1003.101.12.1014)"

• "Encounter, Performed: Discharge Services - Nursing Facility" (2.16.840.1.113883.3.464.1003.101.11.1065)"

• "Encounter, Performed: Encounter Inpatient" (2.16.840.1.113883.3.666.5.307)"

• "Encounter, Performed: Home Healthcare Services" (2.16.840.1.113883.3.464.1003.101.12.1016)"

• "Encounter, Performed: Nursing Facility Visit" (2.16.840.1.113883.3.464.1003.101.12.1012)"

• "Encounter, Performed: Office Visit" (2.16.840.1.113883.3.464.1003.101.12.1001)"

• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" (2.16.840.1.113883.3.464.1003.101.12.1025)"

• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1023)"

• "Immunization, Administered: Pneumococcal Vaccine" using (2.16.840.1.113883.3.464.1003.110.12.1027)"

• "Intervention, Order: Hospice care ambulatory" using "Hospice care ambulatory (2.16.840.1.113762.1.4.1108.15)"

• "Intervention, Performed: Hospice care ambulatory" (2.16.840.1.113762.1.4.1108.15)"

• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"

• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"

• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: Pneumococcal Vaccine Administered" using (2.16.840.1.113883.3.464.1003.110.12.1034)"

Value Sets: CMS 130V7 Colorectal Cancer Screening

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>codesystem &quot;SNOMEDCT&quot; using &quot;2.16.840.1.113883.6.96 version 2017-09&quot;</td>
<td></td>
</tr>
<tr>
<td>code &quot;Discharge to healthcare facility for hospice care (procedure)&quot; using &quot;SNOMEDCT version 2017-09 Code (428371000124100)&quot;</td>
<td></td>
</tr>
<tr>
<td>code &quot;Discharge to home for hospice care (procedure)&quot; using &quot;SNOMEDCT version 2017-09 Code (428361000124107)&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Annual Wellness Visit&quot; using &quot;2.16.840.1.113883.3.526.3.1240&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Colonoscopy&quot; using &quot;2.16.840.1.113883.3.464.1003.108.12.1020&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;CT Colonography&quot; using &quot;2.16.840.1.113883.3.464.1003.108.12.1038&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Encounter Inpatient&quot; using &quot;2.16.840.1.113883.3.666.5.307&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Ethnicity&quot; using &quot;2.16.840.1.114222.4.11.837&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;FOBT&quot; using &quot;2.16.840.1.113883.3.464.1003.198.12.1011&quot;</td>
<td></td>
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<tr>
<td>valueset &quot;FIT DNA&quot; using &quot;2.16.840.1.113883.3.464.1003.108.12.1039&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Flexible Sigmoidoscopy&quot; using &quot;2.16.840.1.113883.3.464.1003.198.12.1010&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Home Healthcare Services&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1016&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Hospice care ambulatory&quot; using &quot;2.16.840.1.113762.1.4.1108.15&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Malignant Neoplasm of Colon&quot; using &quot;2.16.840.1.113883.3.464.1003.108.12.1001&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Office Visit&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1001&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;ONC Administrative Sex&quot; using &quot;2.16.840.1.113762.1.4.1&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Payer&quot; using &quot;2.16.840.1.114222.4.11.3591&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Preventive Care Services - Established Office Visit, 18 and Up&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1025&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Preventive Care Services-Initial Office Visit, 18 and Up&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1023&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Race&quot; using &quot;2.16.840.1.114222.4.11.836&quot;</td>
<td></td>
</tr>
</tbody>
</table>
• valueset "Total Colectomy" using "2.16.840.1.113883.3.464.1003.198.12.1019"

• "Diagnosis: Malignant Neoplasm of Colon" using
  (2.16.840.1.113883.3.464.1003.108.12.1001)"

• Diagnostic Study, Performed: CT
  Colonography:(2.16.840.1.113883.3.464.1003.108.12.1038)

• "Encounter, Performed: Annual Wellness Visit" using
  (2.16.840.1.113883.3.526.3.1240)"

• "Encounter, Performed: Encounter Inpatient" using
  (2.16.840.1.113883.3.666.5.307)"

• "Encounter, Performed: Home Healthcare Services" using
  (2.16.840.1.113883.3.464.1003.101.12.1016)"

• "Encounter, Performed: Nursing Facility Visit"
  (2.16.840.1.113883.3.464.1003.101.12.1012)"

• "Encounter, Performed: Office Visit"
  (2.16.840.1.113883.3.464.1003.101.12.1001)"

• “Encounter, Performed: Care Services in Long-Term Residential Facility:
  (2.16.840.1.113883.3.464.1003.101.12.1014)

• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18
  and Up" using (2.16.840.1.113883.3.464.1003.101.12.1025)"

• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up"
  using (2.16.840.1.113883.3.464.1003.101.12.1023)"

• "Intervention, Order: Hospice care ambulatory" using
  (2.16.840.1.113762.1.4.1108.15)"

• "Intervention, Performed: Hospice care ambulatory"
  (2.16.840.1.113762.1.4.1108.15)"

• "Laboratory Test, Performed: FOBT" using
  (2.16.840.1.113883.3.464.1003.198.12.1011)"

• "Laboratory Test, Performed: FIT DNA" using
  (2.16.840.1.113883.3.464.1003.108.12.1039)"

• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"

• "Patient Characteristic Payer: Payer" (2.16.840.1.114222.4.11.3591)"

• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"

• "Patient Characteristic Sex: ONC Administrative Sex" (2.16.840.1.113762.1.4.1)"

• "Procedure, Performed: Colonoscopy" using
  (2.16.840.1.113883.3.464.1003.108.12.1020)"
- "Procedure, Performed: CT Colonography" using (2.16.840.1.113883.3.464.1003.108.12.1038)"
- "Procedure, Performed: Flexible Sigmoidoscopy" using (2.16.840.1.113883.3.464.1003.198.12.1010)"
- "Procedure, Performed: Total Colectomy" using (2.16.840.1.113883.3.464.1003.198.12.1019)"

**Value Sets: CMS 131V8 Diabetes: Eye Exam**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 18-75 years of age with diabetes who had a retinal or dilated eye exam by an eye care professional during the measurement period or a negative retinal exam (no evidence of retinopathy) in the 12 months prior to the measurement period.</th>
</tr>
</thead>
</table>

- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
- valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240"
- valueset "Diabetes" using "2.16.840.1.113883.3.464.1003.103.12.1001"
- valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
- valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15"
- valueset "Negative Finding" using "2.16.840.1.113883.3.464.1003.195.12.1002"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
- valueset "Ophthalmological Services" using "2.16.840.1.113883.3.526.3.1285"
- valueset "Payer" using "2.16.840.1.114222.4.11.3591"
- valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
- valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- valueset "Retinal or Dilated Eye Exam" using "2.16.840.1.113883.3.464.1003.115.12.1088"
• "Diagnosis: Diabetes" using "Diabetes (2.16.840.1.113883.3.464.1003.103.12.1001)"
• “Diagnosis: Diabetic Retinopathy: 2.16.840.1.113883.3.526.3.327
• "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883
Encounter Inpatient (2.16.840.1.113883.3.666.5.307)"
• "Encounter, Performed: Home Healthcare Services" using
(2.16.840.1.113883.3.464.1003.101.12.1016)"
• "Encounter, Performed: Office Visit" using
(2.16.840.1.113883.3.464.1003.101.12.1001)" Ophthalmological Services
(2.16.840.1.113883.3.526.3.1285)"
• "Encounter, Performed: Preventive Care Services - Established Office Preventive Care Services - Established Office Visit, 18 and Up
(2.16.840.1.113883.3.464.1003.101.12.1025)"
• "Encounter, Performed: Preventive Care Services-Initial Preventive Care Services-Initial Office Visit, 18 and Up
(2.16.840.1.113883.3.464.1003.101.12.1023)"
• “Encounter, Performed: Care Services in Long-Term Residential Facility:
(2.16.840.1.113883.3.464.1003.101.12.1014)
• “Encounter, Performed: Nursing Facility Visit:
(2.16.840.1.113883.3.464.1003.101.12.1012)
• "Intervention, Order: Hospice care ambulatory" using
(2.16.840.1.113762.1.4.1108.15)"
• "Intervention, Performed: Hospice care ambulatory" using
(2.16.840.1.113762.1.4.1108.15)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using
(2.16.840.1.113762.1.4.1)"
• "Physical Exam, Performed: Retinal or Dilated Eye Exam" using
(2.16.840.1.113883.3.464.1003.115.12.1088)"

Value Sets: CMS 134V8 Diabetes: Medical Attention for Nephropathy

<table>
<thead>
<tr>
<th>Description:</th>
<th>The percentage of patients 18-75 years of age with diabetes who had a nephropathy screening test or evidence of nephropathy during the measurement period.</th>
</tr>
</thead>
</table>

• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
• code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
• valueset "ACE Inhibitor or ARB" using "2.16.840.1.113883.3.526.3.1139"
• valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240"
• valueset "Diabetes" using "2.16.840.1.113883.3.464.1003.103.12.1001"
• valueset "Diabetic Nephropathy" using "2.16.840.1.113883.3.464.1003.109.12.1004"
• valueset "Dialysis Education" using "2.16.840.1.113883.3.464.1003.109.12.1016"
• valueset "Dialysis Services" using "2.16.840.1.113883.3.464.1003.109.12.1013"
• valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307"
• valueset "ESRD Monthly Outpatient Services" using "2.16.840.1.113883.3.464.1003.109.12.1014"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Glomerulonephritis and Nephrotic Syndrome" using "2.16.840.1.113883.3.464.1003.109.12.1018"
• valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
• valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15"
• valueset "Hypertensive Chronic Kidney Disease" using "2.16.840.1.113883.3.464.1003.109.12.1017"
• valueset "Kidney Failure" using "2.16.840.1.113883.3.464.1003.109.12.1028"
• valueset "Kidney Transplant" using "2.16.840.1.113883.3.464.1003.109.12.1012"
• valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Other Services Related to Dialysis" using "2.16.840.1.113883.3.464.1003.109.12.1015"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
• valueset "Preventive Care Services- Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
• valueset "Proteinuria" using "2.16.840.1.113883.3.526.3.1003"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- valueset "Urine Protein Tests" using "2.16.840.1.113883.3.464.1003.109.12.1024"
- "Diagnosis: Diabetes" using (2.16.840.1.113883.3.464.1003.103.12.1001)"
- "Diagnosis: Diabetic Nephropathy" using "Di" (2.16.840.1.113883.3.464.1003.109.12.1004)"
- "Diagnosis: Glomerulonephritis and Nephrotic Syndrome" using (2.16.840.1.113883.3.464.1003.109.12.1018)"
- Hypertensive Chronic Kidney Disease (2.16.840.1.113883.3.464.1003.109.12.1017)"
- "Diagnosis: Kidney Failure" using (2.16.840.1.113883.3.464.1003.109.12.1028)"
- "Diagnosis: Proteinuria" using " (2.16.840.1.113883.3.526.3.1003)"
- "Encounter, Performed: Annual Wellness Visit" using "(2.16.840.1.113883.3.526.3.1240)"
- "Encounter, Performed: Encounter Inpatient" using "(2.16.840.1.113883.3.666.5.307)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1025)"
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1023)"
- “Encounter, Performed: Care Services in Long-Term Residential Facility:" (2.16.840.1.1.113883.3.464.1003.101.12.1014)
- “Encounter, Performed: Nursing Facility Visit:" (2.16.840.1.1.113883.3.464.1003.101.12.1012)
- "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Intervention, Performed: Dialysis Education" using (2.16.840.1.113883.3.464.1003.109.12.1016)"
• "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
• "Intervention, Performed: Other Services Related to Dialysis" using (2.16.840.1.113883.3.464.1003.109.12.1015)"
• "Laboratory Test, Performed: Urine Protein Tests" using (2.16.840.1.113883.3.464.1003.109.12.1024)"
• "Medication, Active: ACE Inhibitor or ARB" using (2.16.840.1.113883.3.526.3.1139)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: Dialysis Services" using (2.16.840.1.113883.3.464.1003.109.12.1013)"
• "Procedure, Performed: Kidney Transplant" using (2.16.840.1.113883.3.464.1003.109.12.1012)"

Value Sets: CMS 137V8 Initiation and Engagement of Alcohol and Other Drug Dependence Treatment

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 13 years of age and older with a new episode of AOD dependence who received the following. Two rates are reported.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Percentage of patients who initiated treatment within 14 days of the diagnosis.</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients who initiated treatment and who had two or more additional services with an AOD diagnosis within 30 days of the initiation visit.</td>
</tr>
</tbody>
</table>

• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
• code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
• "Diagnosis: Alcohol and Drug Dependence" using (2.16.840.1.113883.3.464.1003.106.12.1001)"
- "Intervention, Performed: Alcohol and Drug Dependence Treatment" (2.16.840.1.113883.3.464.1003.106.12.1005)"
- "Encounter, Performed: Discharge Services - Hospital Inpatient Same Day Discharge" using (2.16.840.1.113883.3.464.1003.101.12.1006)"
- "Encounter, Performed: Discharge Services - Hospital Inpatient" using (2.16.840.1.113883.3.464.1003.101.12.1007)"
- "Encounter, Performed: ED Visit" using (2.16.840.1.113883.3.464.1003.101.12.1010)"
- "Encounter, Performed: Encounter Inpatient" using (2.16.840.1.113883.3.666.5.307)"
- "Encounter, Performed: Hospital Inpatient Visit - Initial" using "(2.16.840.1.113883.3.464.1003.101.12.1004)"
- "Encounter, Performed: Hospital Observation Care - Initial" using (2.16.840.1.113883.3.464.1003.101.12.1002)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Intervention, Performed: Psych Visit - Psychotherapy" using (2.16.840.1.113883.3.526.3.1496)"
- "Encounter, Performed: Telehealth Services: 2.16.840.1.113883.3.464.1003.101.12.1031"
- "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
### Value Sets: CMS 138V8 Preventive Care and Screening: Tobacco Use: Screening and Cessation Intervention

<table>
<thead>
<tr>
<th>Description:</th>
<th>Value Sets</th>
</tr>
</thead>
</table>
| Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. | • valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Health & Behavioral Assessment - Individual" using "2.16.840.1.113883.3.526.3.1020"
• valueset "Health and Behavioral Assessment - Initial" using "2.16.840.1.113883.3.526.3.1245"
• valueset "Health and Behavioral Assessment, Reassessment" using "2.16.840.1.113883.3.526.3.1529"
• valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
• valueset "Limited Life Expectancy" using "2.16.840.1.113883.3.526.3.1259"
• valueset "Medical Reason" using "2.16.840.1.113883.3.526.3.1007"
• valueset "Occupational Therapy Evaluation" using "2.16.840.1.113883.3.526.3.1011"
• valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Ophthalmological Services" using "2.16.840.1.113883.3.526.3.1285"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025" |
| Three rates are reported:                                                                            |                                                                           |
| • Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months. |                                                                           |
| • Percentage of patients aged 18 years and older who were screened for tobacco use and identified as a tobacco user who received tobacco cessation intervention. |                                                                           |
| • Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. |                                                                           |
• valueset "Preventive Care Services - Group Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1027"
• valueset "Preventive Care Services - Other" using "2.16.840.1.113883.3.464.1003.101.12.1030"
• valueset "Preventive Care Services-Individual Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1026"
• valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
• valueset "Psych Visit - Diagnostic Evaluation" using "2.16.840.1.113883.3.526.3.1492"
• valueset "Psych Visit - Psychotherapy" using "2.16.840.1.113883.3.526.3.1496"
• valueset "Psychoanalysis" using "2.16.840.1.113883.3.526.3.1141"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• valueset "Speech and Hearing Evaluation" using "2.16.840.1.113883.3.526.3.1530"
• valueset "Tobacco Non-User" using "2.16.840.1.113883.3.526.3.1189"
• valueset "Tobacco Use Cessation Counseling" using "2.16.840.1.113883.3.526.3.509"
• valueset "Tobacco Use Cessation Pharmacotherapy" using "2.16.840.1.113883.3.526.3.1190"
• valueset "Tobacco Use Screening" using "2.16.840.1.113883.3.526.3.1278"
• valueset "Tobacco User" using "2.16.840.1.113883.3.526.3.1170"
• "Assessment, Not Performed: Tobacco Use Screening" using (2.16.840.1.113883.3.526.3.1278)"
• "Assessment, Performed: Tobacco Use Screening" using (2.16.840.1.113883.3.526.3.1278)"
• "Diagnosis: Limited Life Expectancy" using (2.16.840.1.113883.3.526.3.1259)"
• "Encounter, Performed: Annual Wellness Visit" (2.16.840.1.113883.3.526.3.1240)"
• "Encounter, Performed: Health & Behavioral Assessment - Individual" using (2.16.840.1.113883.3.526.3.1020)"
• "Encounter, Performed: Health and Behavioral Assessment - Initial" using (2.16.840.1.113883.3.526.3.1245)"
• "Encounter, Performed: Health and Behavioral Assessment, Reassessment" using (2.16.840.1.113883.3.526.3.1529)"
• "Encounter, Performed: Home Healthcare Services" using (2.16.840.1.113883.3.464.1003.101.12.1016"
• "Encounter, Performed: Occupational Therapy Evaluation" using (2.16.840.1.113883.3.526.3.1011"
• "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001"
• "Encounter, Performed: Ophthalmological Services" using (2.16.840.1.113883.3.526.3.1285"
• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1025"
• "Encounter, Performed: Preventive Care Services - Group Counseling" using (2.16.840.1.113883.3.464.1003.101.12.1027"
• "Encounter, Performed: Preventive Care Services - Other" using (2.16.840.1.113883.3.464.1003.101.12.1030"
• "Encounter, Performed: Preventive Care Services-Individual Counseling" using (2.16.840.1.113883.3.464.1003.101.12.1026"
• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using " (2.16.840.1.113883.3.464.1003.101.12.1023"
• "Encounter, Performed: Psych Visit - Diagnostic Evaluation" using 2.16.840.1.113883.3.526.3.1492"
• "Encounter, Performed: Psych Visit - Psychotherapy" using " (2.16.840.1.113883.3.526.3.1496"
• "Encounter, Performed: Psychoanalysis " using (2.16.840.1.113883.3.526.3.1141"
• "Encounter, Performed: Speech and Hearing Evaluation" using (2.16.840.1.113883.3.526.3.1530"
• "Intervention, Not Performed: Tobacco Use Cessation Counseling" using (2.16.840.1.113883.3.526.3.509"
• "Intervention, Performed: Tobacco Use Cessation Counseling" using (2.16.840.1.113883.3.526.3.509"
• "Medication, Active: Tobacco Use Cessation Pharmacotherapy" using (2.16.840.1.113883.3.526.3.1190"
• "Medication, Not Ordered: Tobacco Use Cessation Pharmacotherapy" using (2.16.840.1.113883.3.526.3.1190"
• "Medication, Order: Tobacco Use Cessation Pharmacotherapy" using (2.16.840.1.113883.3.526.3.1190"
- "Patient Characteristic Ethnicity: Ethnicity" using "Ethnicity (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"

**Value Sets: CMS 139V8 Falls: Screening for Future Fall Risk**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 65 years of age and older who were screened for future fall risk during the measurement period.</th>
</tr>
</thead>
<tbody>
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<td>codesystem &quot;SNOMEDCT&quot; using &quot;2.16.840.1.113883.6.96 version 2017-09&quot;</td>
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<tr>
<td>code &quot;Discharge to healthcare facility for hospice care (procedure)&quot; using &quot;SNOMEDCT version 2017-09 Code (428371000124100)&quot;</td>
<td></td>
</tr>
<tr>
<td>code &quot;Discharge to home for hospice care (procedure)&quot; using &quot;SNOMEDCT version 2017-09 Code (428361000124107)&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Ambulatory Status&quot; using &quot;2.16.840.1.113883.3.464.1003.118.11.1219&quot;</td>
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</tr>
<tr>
<td>valueset &quot;Annual Wellness Visit&quot; using &quot;2.16.840.1.113883.3.526.3.1240&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Audiology Visit&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1066&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Care Services in Long-Term Residential Facility&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1014&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Discharge Services - Nursing Facility&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1013&quot;</td>
<td></td>
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<tr>
<td>valueset &quot;Encounter Inpatient&quot; using &quot;2.16.840.1.113883.3.666.5.307&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Ethnicity&quot; using &quot;2.16.840.1.114222.4.11.837&quot;</td>
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<td>valueset &quot;Falls Screening&quot; using &quot;2.16.840.1.113883.3.464.1003.118.12.1028&quot;</td>
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<tr>
<td>valueset &quot;Home Healthcare Services&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1016&quot;</td>
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<tr>
<td>valueset &quot;Hospice care ambulatory&quot; using &quot;2.16.840.1.113762.1.4.1108.15&quot;</td>
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<tr>
<td>valueset &quot;Nursing Facility Visit&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1012&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Office Visit&quot; using &quot;2.16.840.1.113883.3.464.1003.101.12.1001&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;ONC Administrative Sex&quot; using &quot;2.16.840.1.113762.1.4.1&quot;</td>
<td></td>
</tr>
<tr>
<td>valueset &quot;Ophthalmological Services&quot; using &quot;2.16.840.1.113883.3.526.3.1285&quot;</td>
<td></td>
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<tr>
<td>valueset &quot;Patient not ambulatory&quot; using &quot;2.16.840.1.113883.3.464.1003.118.12.1009&quot;</td>
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</tr>
</tbody>
</table>
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
• valueset "Preventive Care Services-Individual Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1026"
• valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• "Assessment, Performed: Ambulatory Status" using (2.16.840.1.113883.3.464.1003.118.11.1219)"
• "Assessment, Performed: Falls Screening" using (2.16.840.1.113883.3.464.1003.118.12.1028)"
• "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883.3.526.3.1240)"
• "Encounter, Performed: Audiology Visit" using (2.16.840.1.113883.3.464.1003.101.12.1066)"
• "Encounter, Performed: Care Services in Long-Term Residential Facility" using (2.16.840.1.113883.3.464.1003.101.12.1014)"
• "Encounter, Performed: Discharge Services - Nursing Facility" using (2.16.840.1.113883.3.464.1003.101.12.1013)"
• "Encounter, Performed: Encounter Inpatient" using (2.16.840.1.113883.3.666.5.307)"
• "Encounter, Performed: Home Healthcare Services" using (2.16.840.1.113883.3.464.1003.101.12.1016)"
• "Encounter, Performed: Nursing Facility Visit" using "Nursing Facility Visit (2.16.840.1.113883.3.464.1003.101.12.1012)"
• "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
• "Encounter, Performed: Ophthalmological Services" using " (2.16.840.1.113883.3.526.3.1285)"
• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1025)"
• "Encounter, Performed: Preventive Care Services-Individual Counseling" using " (2.16.840.1.113883.3.464.1003.101.12.1026)"
• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using " (2.16.840.1.113883.3.464.1003.101.12.1023)"
• "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)
• "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)
• "Patient Characteristic Ethnicity: Ethnicity" (2.16.840.1.114222.4.11.837)
• "Patient Characteristic Payer: Payer" using " (2.16.840.1.114222.4.11.3591)
• "Patient Characteristic Race: Race" using Race (2.16.840.1.114222.4.11.836)
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)

Value Sets: CMS 144V8 HF: Beta-Blocker Therapy for Left Ventricular Systolic Dysfunction (LVSD)

<table>
<thead>
<tr>
<th>Description:</th>
<th>Percentage of patients aged 18 years and older with a diagnosis of HF with a current or prior LVEF &lt; 40% who were prescribed beta-blocker therapy either within a 12-month period when seen in the outpatient setting OR at each hospital discharge.</th>
</tr>
</thead>
</table>

• codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
• code "Heart rate" using "LOINC version 2.63 Code (8867-4)"
• valueset "Allergy to Beta Blocker Therapy" using "2.16.840.1.113883.3.526.3.1177"
• valueset "Arrhythmia" using "2.16.840.1.113883.3.526.3.366"
• valueset "Asthma" using "2.16.840.1.113883.3.526.3.362"
• valueset "Atrioventricular Block" using "2.16.840.1.113883.3.526.3.367"
• valueset "Beta Blocker Therapy for LVSD" using "2.16.840.1.113883.3.526.3.1184"
• valueset "Beta Blocker Therapy Ingredient" using "2.16.840.1.113883.3.526.3.1493"
• valueset "Bradycardia" using "2.16.840.1.113883.3.526.3.412"
• valueset "Cardiac Pacer in Situ" using "2.16.840.1.113883.3.526.3.368"
• valueset "Cardiac Pacer" using "2.16.840.1.113883.3.526.3.1193"
• valueset "Care Services in Long-Term Residential Facility" using "2.16.840.1.113883.3.464.1003.101.12.1014"
• valueset "Discharge Services - Hospital Inpatient" using "2.16.840.1.113883.3.464.1003.101.12.1007"
• valueset "Ejection Fraction" using "2.16.840.1.113883.3.526.3.1134"
• valueset "Ethnicity" using "2.16.840.1.114222.4.1.1.837"
- valueset "Heart Failure" using "2.16.840.1.113883.3.526.3.376"
- valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
- valueset "Hypotension" using "2.16.840.1.113883.3.526.3.370"
- valueset "Intolerance to Beta Blocker Therapy" using "2.16.840.1.113883.3.526.3.1178"
- valueset "Left Ventricular Systolic Dysfunction" using "2.16.840.1.113883.3.526.3.1091"
- valueset "Medical Reason" using "2.16.840.1.113883.3.526.3.1007"
- valueset "Moderate or Severe LVSD" using "2.16.840.1.113883.3.526.3.1090"
- valueset "Moderate or Severe" using "2.16.840.1.113883.3.526.3.1092"
- valueset "Nursing Facility Visit" using "2.16.840.1.113883.3.464.1003.101.12.1012"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
- valueset "Outpatient Consultation" using "2.16.840.1.113883.3.464.1003.101.12.1008"
- valueset "Patient Provider Interaction" using "2.16.840.1.113883.3.526.3.1012"
- valueset "Patient Reason" using "2.16.840.1.113883.3.526.3.1008"
- valueset "Payer" using "2.16.840.1.114222.4.11.3591"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- valueset "System Reason" using "2.16.840.1.113883.3.526.3.1009"
- "Allergy/Intolerance: Beta Blocker Therapy Ingredient" (2.16.840.1.113883.3.526.3.1493)"
- "Device, Applied: Cardiac Pacer" (2.16.840.1.113883.3.526.3.1193)"
- "Diagnosis: Allergy to Beta Blocker Therapy" using (2.16.840.1.113883.3.526.3.1177)"
- "Diagnosis: Arrhythmia" using (2.16.840.1.113883.3.526.3.366)"
- "Diagnosis: Asthma" using (2.16.840.1.113883.3.526.3.362)"
- "Diagnosis: Atrioventricular Block" using (2.16.840.1.113883.3.526.3.367)"
- "Diagnosis: Bradycardia" (2.16.840.1.113883.3.526.3.412)"
- "Diagnosis: Cardiac Pacemaker in Situ" using (2.16.840.1.113883.3.526.3.368)"
- "Diagnosis: Heart Failure" using (2.16.840.1.113883.3.526.3.376)"
- "Diagnosis: Hypotension" using (2.16.840.1.113883.3.526.3.370)"
- "Diagnosis: Intolerance to Beta Blocker Therapy" using (2.16.840.1.113883.3.526.3.1178)"
- "Diagnosis: Left Ventricular Systolic Dysfunction" using "(2.16.840.1.113883.3.526.3.1091)"
- "Diagnosis: Moderate or Severe LVSD" using (2.16.840.1.113883.3.526.3.1090)"
- "Diagnosis: Moderate or Severe LVSD" using (2.16.840.1.113883.3.526.3.1090)"
- "Diagnostic Study, Performed: Ejection Fraction" using (2.16.840.1.113883.3.526.3.1134)"
- "Encounter, Performed: Care Services in Long-Term Residential Facility" using "(2.16.840.1.113883.3.464.1003.101.12.1014)"
- "Encounter, Performed: Discharge Services - Hospital Inpatient" using (2.16.840.1.113883.3.464.1003.101.12.1007)"
- "Encounter, Performed: Nursing Facility Visit" using (2.16.840.1.113883.3.464.1003.101.12.1012)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Encounter, Performed: Outpatient Consultation" using (2.16.840.1.113883.3.464.1003.101.12.1008)"
- "Encounter, Performed:" using n (2.16.840.1.113883.3.526.3.1012)"
- "Medication, Active: Beta Blocker Therapy for LVSD" using (2.16.840.1.113883.3.526.3.1184)"
- "Medication, Not Ordered: Beta Blocker Therapy for LVSD" using (2.16.840.1.113883.3.526.3.1184)"
- "Medication, Order: Beta Blocker Therapy for LVSD" using (2.16.840.1.113883.3.526.3.1184)"
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
- "Physical Exam, Performed: Heart rate" using (LOINC version 2.63 Code 8867-4)"
Value Sets: CMS 155V8 Weight Assessment and Counseling for Nutrition and Physical Activity for Children and Adolescents

| Description | Percentage of patients 3-17 years of age who had an outpatient visit with a PCP or OB/GYN and who had evidence of the following during the measurement period. Three rates are reported.
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Percentage of patients with height, weight, and BMI percentile documentation</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients with counseling for nutrition</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients with counseling for physical activity.</td>
</tr>
</tbody>
</table>

- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
- valueset "BMI percentile" using "2.16.840.1.113883.3.464.1003.121.12.1012"
- valueset "Counseling for Nutrition" using "2.16.840.1.113883.3.464.1003.195.12.1003"
- valueset "Counseling for Physical Activity" using "2.16.840.1.113883.3.464.1003.118.12.1035"
- valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Height" using "2.16.840.1.113883.3.464.1003.121.12.1014"
- valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
- valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1.4.1"
- valueset "Payer" using "2.16.840.1.114222.4.1.3591"
- valueset "Pregnancy" using "2.16.840.1.113883.3.526.3.378"
- valueset "Preventive Care - Established Office Visit, 0 to 17" using "2.16.840.1.113883.3.464.1003.101.12.1024"
- valueset "Preventive Care Services - Group Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1027"
- valueset "Preventive Care Services-Individual Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1026"
- valueset "Preventive Care- Initial Office Visit, 0 to 17" using "2.16.840.1.113883.3.464.1003.101.12.1022"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- valueset "Weight" using "2.16.840.1.113883.3.464.1003.121.12.1015"
- "Diagnosis: Pregnancy" using "Pregnancy (2.16.840.1.113883.3.526.3.378)"
- "Encounter, Performed: Encounter Inpatient" using (2.16.840.1.113883.3.666.5.307)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Encounter, Performed: Preventive Care - Established Office Visit, 0 to 17" using (2.16.840.1.113883.3.464.1003.101.12.1024)"
- "Encounter, Performed: Preventive Care Services - Group Counseling" using (2.16.840.1.113883.3.464.1003.101.12.1027)"
- "Encounter, Performed: Preventive Care Services-Individual Counseling" using (2.16.840.1.113883.3.464.1003.101.12.1026)"
- "Encounter, Performed: Preventive Care- Initial Office Visit, 0 to 17" using (2.16.840.1.113883.3.464.1003.101.12.1022)"
- "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1.1108.15)"
- "Intervention, Performed: Counseling for Nutrition" using (2.16.840.1.113883.3.464.1003.195.12.1003)"
- "Intervention, Performed: Counseling for Physical Activity" using (2.16.840.1.113883.3.464.1003.118.12.1035)"
- "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1.1108.15)"
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
- "Physical Exam, Performed: BMI percentile" using (2.16.840.1.113883.3.464.1003.121.12.1012)"
• "Physical Exam, Performed: Height" using (2.16.840.1.113883.3.464.1003.121.12.1014)"

• "Physical Exam, Performed: Weight" using (2.16.840.1.113883.3.464.1003.121.12.1015)"

**Value Sets: CMS 156V8 Use of High-Risk Medications in the Elderly**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 65 years of age and older who were ordered high-risk medications. Two rates are reported.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Percentage of patients who were ordered at least one high-risk medication.</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients who were ordered at least two of the same high-risk medications.</td>
</tr>
</tbody>
</table>

- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
- code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
- valueset "Acetaminophen / Butalbital / Caffeine / Codeine" using "2.16.840.1.113883.3.464.1003.196.12.1326"
- valueset "Acetaminophen / Butalbital / Caffeine" using "2.16.840.1.113883.3.464.1003.196.12.1363"
- valueset "Acetaminophen / Butalbital" using "2.16.840.1.113883.3.464.1003.196.12.1317"
- valueset "Acetaminophen / Chlorpheniramine / Dextromethorphan / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1310"
- valueset "Acetaminophen / Chlorpheniramine / Dextromethorphan / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1321"
- valueset "Acetaminophen / Chlorpheniramine / Dextromethorphan" using "2.16.840.1.113883.3.464.1003.196.12.1421"
- valueset "Acetaminophen / Chlorpheniramine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1328"
- valueset "Acetaminophen / Chlorpheniramine / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1316"
- valueset "Acetaminophen / Dextromethorphan" using "2.16.840.1.113883.3.464.1003.196.12.1405"
- valueset "Acetaminophen / Dextromethorphan / Diphenhydramine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1399"
• valueset "Acetaminophen / Dextromethorphan / Doxylamine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1287"
• valueset "Acetaminophen / Dextromethorphan / Doxylamine" using "2.16.840.1.113883.3.464.1003.196.12.1338"
• valueset "Acetaminophen / Diphenhydramine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1318"
• valueset "Acetaminophen / Diphenhydramine" using "2.16.840.1.113883.3.464.1003.196.12.1350"
• valueset "Amitriptyline / Chlordiazepoxide" using "2.16.840.1.113883.3.464.1003.196.12.1286"
• valueset "Amitriptyline / Perphenazine" using "2.16.840.1.113883.3.464.1003.196.12.1428"
• valueset "Amitriptyline Hydrochloride" using "2.16.840.1.113883.3.464.1003.196.12.1373"
• valueset "Amoxapine" using "2.16.840.1.113883.3.464.1003.196.12.1273"
• valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240"
• valueset "Anti Infectives, other" using "2.16.840.1.113883.3.464.1003.196.12.1481"
• valueset "Aspirin / Butalbital / Caffeine" using "2.16.840.1.113883.3.464.1003.196.12.1347"
• valueset "Aspirin / Caffeine / Orphenadrine" using "2.16.840.1.113883.3.464.1003.196.12.1302"
• valueset "Atropine / Diphenoxylate" using "2.16.840.1.113883.3.464.1003.196.12.1274"
• valueset "Atropine / Hyoscyamine / Phenobarbital / Scopolamine" using "2.16.840.1.113883.3.464.1003.196.12.1355"
• valueset "Benztropine" using "2.16.840.1.113883.3.464.1003.196.12.1361"
• valueset "Brompheniramine / Codeine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1409"
• valueset "Brompheniramine / Codeine / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1450"
• valueset "Brompheniramine / Dextromethorphan / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1325"
• valueset "Brompheniramine / Dextromethorphan / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1339"
• valueset "Brompheniramine / Pseudoephedrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1332"
• valueset "Brompheniramine" using "2.16.840.1.113883.3.464.1003.196.12.1427"
• valueset "Butabarbital" using "2.16.840.1.113883.3.464.1003.196.12.1402"
• valueset "Carbinoxamine" using "2.16.840.1.113883.3.464.1003.196.12.1306"
• valueset "Care Services in Long-Term Residential Facility" using 
  "2.16.840.1.113883.3.464.1003.101.12.1014"
• valueset "Carisoprodol" using "2.16.840.1.113883.3.464.1003.196.12.1369"
• valueset "Chlophedianol / Chlorpheniramine / Phenylephrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1412"
• valueset "Chlophedianol / Dexchlorpheniramine / Pseudoephedrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1447"
• valueset "Chlorpheniramine / Dextromethorphan / Phenylephrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1312"
• valueset "Chlorpheniramine / Dextromethorphan / Pseudoephedrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1337"
• valueset "Chlorpheniramine / Hydrocodone / Pseudoephedrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1299"
• valueset "Chlorpheniramine / Hydrocodone" using 
  "2.16.840.1.113883.3.464.1003.196.12.1330"
• valueset "Chlorpheniramine / Phenylephrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1343"
• valueset "Chlorpheniramine / Pseudoephedrine" using 
  "2.16.840.1.113883.3.464.1003.196.12.1315"
• valueset "Chlorpheniramine" using "2.16.840.1.113883.3.464.1003.196.12.1352"
• valueset "Chlorpropamide" using "2.16.840.1.113883.3.464.1003.196.12.1303"
• valueset "Chlorzoxazone" using "2.16.840.1.113883.3.464.1003.196.12.1362"
• valueset "Clemastine" using "2.16.840.1.113883.3.464.1003.196.12.1308"
• valueset "Clomipramine" using "2.16.840.1.113883.3.464.1003.196.12.1336"
• valueset "Conjugated Estrogens / Medroxyprogesterone" using 
  "2.16.840.1.113883.3.464.1003.196.12.1324"
• valueset "Conjugated Estrogens" using 
  "2.16.840.1.113883.3.464.1003.196.12.1357"
• valueset "Cyclobenzaprine Hydrochloride" using 
  "2.16.840.1.113883.3.464.1003.196.12.1372"
• valueset "Cyproheptadine" using "2.16.840.1.113883.3.464.1003.196.12.1277"
• valueset "Desiccated Thyroid" using "2.16.840.1.113883.3.464.1003.196.12.1354"
• valueset "Desipramine" using "2.16.840.1.113883.3.464.1003.196.12.1278"
• valueset "Dexbrompheniramine / Dextromethorphan / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1426"
• valueset "Dexbrompheniramine / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1430"
• valueset "Dexbrompheniramine Maleate / Pseudoephedrine Hydrochloride" using "2.16.840.1.113883.3.464.1003.196.12.1429"
• valueset "Dexbrompheniramine" using "2.16.840.1.113883.3.464.1003.196.12.1375"
• valueset "Dexchlorpheniramine / Dextromethorphan / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1425"
• valueset "Dexchlorpheniramine / Pseudoephedrine" using "2.16.840.1.113883.3.464.1003.196.12.1425"
• valueset "Dextromethorphan / Diphenhydramine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1397"
• valueset "Dextromethorphan / Doxylamine" using "2.16.840.1.113883.3.464.1003.196.12.1452"
• valueset "Dicyclomine" using "2.16.840.1.113883.3.464.1003.196.12.1279"
• valueset "Dienogest / Estradiol Multiphasic" using "2.16.840.1.113883.3.464.1003.196.12.1398"
• valueset "Dimenhydrinate" using "2.16.840.1.113883.3.464.1003.196.12.1500"
• valueset "Diphenhydramine / Ibuprofen" using "2.16.840.1.113883.3.464.1003.196.12.1293"
• valueset "Diphenhydramine / Phenylephrine" using "2.16.840.1.113883.3.464.1003.196.12.1307"
• valueset "Diphenhydramine Hydrochloride" using "2.16.840.1.113883.3.464.1003.196.12.1371"
• valueset "Dipyridamole" using "2.16.840.1.113883.3.464.1003.196.12.1349"
• valueset "Discharge Services - Nursing Facility" using "2.16.840.1.113883.3.464.1003.101.12.1013"
• valueset "Disopyramide" using "2.16.840.1.113883.3.464.1003.196.12.1311"
• valueset "Drospirenone / Estradiol" using "2.16.840.1.113883.3.464.1003.196.12.1410"
• valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307"
• valueset "Esterified Estrogens / Methyltestosterone" using "2.16.840.1.113883.3.464.1003.196.12.1320"
• valueset "Esterified Estrogens" using "2.16.840.1.113883.3.464.1003.196.12.1419"
• valueset "Estradiol / Norethindrone" using "2.16.840.1.113883.3.464.1003.196.12.1323"
• valueset "Estradiol" using "2.16.840.1.113883.3.464.1003.196.12.1365"
• valueset "Estropipate" using "2.16.840.1.113883.3.464.1003.196.12.1319"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Glyburide / Metformin" using "2.16.840.1.113883.3.464.1003.196.12.1360"
• valueset "Glyburide" using "2.16.840.1.113883.3.464.1003.196.12.1368"
• valueset "Guanfacine" using "2.16.840.1.113883.3.464.1003.196.12.1341"
• valueset "High Risk Medications for the Elderly" using "2.16.840.1.113883.3.464.1003.196.12.1253"
• valueset "High-Risk Medications With Days Supply Criteria" using "2.16.840.1.113883.3.464.1003.196.12.1254"
• valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
• valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1.1108.15"
• valueset "Hydrochlorothiazide / Methyl dopa" using "2.16.840.1.113883.3.464.1003.196.12.1374"
• valueset "Hydroxyzine" using "2.16.840.1.113883.3.464.1003.196.12.1364"
• valueset "Hyoscyamine / Methenamine / Mblue / Phenyl Salicyl / Sodium Biphosphate" using "2.16.840.1.113883.3.464.1003.196.12.1504"
• valueset "Hyoscyamine / Methenamine / Mblue / Phenyl Salicyl" using "2.16.840.1.113883.3.464.1003.196.12.1503"
• valueset "Hyoscyamine / Methenamine / Mblue / Sodium Biphosphate" using "2.16.840.1.113883.3.464.1003.196.12.1505"
• valueset "Hyoscyamine" using "2.16.840.1.113883.3.464.1003.196.12.1501"
• valueset "Imipramine" using "2.16.840.1.113883.3.464.1003.196.12.1359"
- valueset "Indomethacin" using "2.16.840.1.113883.3.464.1003.196.12.1366"
- valueset "Isoxsuprine" using "2.16.840.1.113883.3.464.1003.196.12.1422"
- valueset "Ketorolac Tromethamine" using "2.16.840.1.113883.3.464.1003.196.12.1364"
- valueset "List of Single RxNorm Code Concepts for High Risk Drugs for the Elderly" using "2.16.840.1.113883.3.464.1003.196.12.1272"
- valueset "Meclizine" using "2.16.840.1.113883.3.464.1003.196.12.1506"
- valueset "Megestrol" using "2.16.840.1.113883.3.464.1003.196.12.1342"
- valueset "Meperidine" using "2.16.840.1.113883.3.464.1003.196.12.1351"
- valueset "Meprobamate" using "2.16.840.1.113883.3.464.1003.196.12.1284"
- valueset "Metaxalone" using "2.16.840.1.113883.3.464.1003.196.12.1358"
- valueset "Methocarbamol" using "2.16.840.1.113883.3.464.1003.196.12.1370"
- valueset "Methyldopa" using "2.16.840.1.113883.3.464.1003.196.12.1331"
- valueset "Nifedipine" using "2.16.840.1.113883.3.464.1003.196.12.1353"
- valueset "Nonbenzodiazepine hypnotics" using "2.16.840.1.113883.3.464.1003.196.12.1480"
- valueset "Nortriptyline" using "2.16.840.1.113883.3.464.1003.196.12.1507"
- valueset "Nursing Facility Visit" using "2.16.840.1.113883.3.464.1003.101.12.1012"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
- valueset "Ophthalmologic Services" using "2.16.840.1.113883.3.464.1003.101.11.1206"
- valueset "Paroxetine" using "2.16.840.1.113883.3.464.1003.196.12.1508"
- valueset "Payer" using "2.16.840.1.114222.4.11.3591"
- valueset "Phenobarbital" using "2.16.840.1.113883.3.464.1003.196.12.1348"
- valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
- valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
- valueset "Promethazine Hydrochloride" using "2.16.840.1.113883.3.464.1003.196.12.1367"
- valueset "Protriptyline" using "2.16.840.1.113883.3.464.1003.196.12.1509"
- valueset "Pseudoephedrine / Triprolidine" using "2.16.840.1.113883.3.464.1003.196.12.1345"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- valueset "Trihexyphenidyl" using "2.16.840.1.113883.3.464.1003.196.12.1334"
- valueset "Trimipramine" using "2.16.840.1.113883.3.464.1003.196.12.1285"
- valueset "Triprolidine" using "2.16.840.1.113883.3.464.1003.196.12.1408"
- "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883.3.526.3.1240)"
- "Encounter, Performed: Care Services in Long-Term Residential Facility" using (2.16.840.1.113883.3.464.1003.101.12.1014)"
- "Encounter, Performed: Discharge Services - Nursing Facility" using (2.16.840.1.113883.3.464.1003.101.12.1013)"
- "Encounter, Performed: Encounter Inpatient" using (2.16.840.1.113883.3.666.5.307)"
- "Encounter, Performed: Nursing Facility Visit" using (2.16.840.1.113883.3.464.1003.101.12.1012)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Encounter, Performed: Ophthalmologic Services" using (2.16.840.1.113883.3.464.1003.101.11.1206)"
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using2.16.840.1.113883.3.464.1003.101.12.1025)"
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1023)"
- "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Medication, Order: Acetaminophen / Butalbital / Caffeine / Codeine" using (2.16.840.1.113883.3.464.1003.196.12.1326)"
- "Medication, Order: Acetaminophen / Butalbital / Caffeine" using (2.16.840.1.113883.3.464.1003.196.12.1363)"
- "Medication, Order: Acetaminophen / Butalbital" using (2.16.840.1.113883.3.464.1003.196.12.1317)"
• "Medication, Order: Acetaminophen / Chlorpheniramine / Dextromethorphan / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1310)"

• "Medication, Order: Acetaminophen / Chlorpheniramine / Dextromethorphan / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1321)"

• "Medication, Order: Acetaminophen / Chlorpheniramine / Dextromethorphan" using (2.16.840.1.113883.3.464.1003.196.12.1421)"

• "Medication, Order: Acetaminophen / Chlorpheniramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1421)"

• "Medication, Order: Acetaminophen / Chlorpheniramine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1316)"

• "Medication, Order: Acetaminophen / Dextromethorphan" using (2.16.840.1.113883.3.464.1003.196.12.1347)"

• "Medication, Order: Acetaminophen / Dextromethorphan / Diphenhydramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1399)"

• "Medication, Order: Acetaminophen / Dextromethorphan / Doxylamine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1287)"

• "Medication, Order: Acetaminophen / Dextromethorphan / Doxylamine" using (2.16.840.1.113883.3.464.1003.196.12.1338)"

• "Medication, Order: Acetaminophen / Diphenhydramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1318)"

• "Medication, Order: Acetaminophen / Diphenhydramine" using (2.16.840.1.113883.3.464.1003.196.12.1350)"

• "Medication, Order: Amitriptyline / Chlordiazepoxide" using (2.16.840.1.113883.3.464.1003.196.12.1286)"

• "Medication, Order: Amitriptyline / Perphenazine" using (2.16.840.1.113883.3.464.1003.196.12.1428)"

• "Medication, Order: Amitriptyline Hydrochloride" using (2.16.840.1.113883.3.464.1003.196.12.1373)"

• "Medication, Order: Amoxapine" using (2.16.840.1.113883.3.464.1003.196.12.1273)"

• "Medication, Order: Anti Infectives, other" using (2.16.840.1.113883.3.464.1003.196.12.1481)"

• "Medication, Order: Aspirin / Butalbital / Caffeine" using Caffeine (2.16.840.1.113883.3.464.1003.196.12.1347)"

• "Medication, Order: Aspirin / Caffeine / Orphenadrine" using (2.16.840.1.113883.3.464.1003.196.12.1302)"
• "Medication, Order: Atropine / Diphenoxylate" using (2.16.840.1.113883.3.464.1003.196.12.1274)"

• "Medication, Order: Atropine / Hyoscyamine / Phenobarbital / Scopolamine" using (2.16.840.1.113883.3.464.1003.196.12.1355)"

• "Medication, Order: Benztropine" using (2.16.840.1.113883.3.464.1003.196.12.1361)"

• "Medication, Order: Brompheniramine / Codeine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1409)"

• "Medication, Order: Brompheniramine / Codeine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1450)"

• "Medication, Order: Brompheniramine / Dextromethorphan / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1325)"

• "Medication, Order: Brompheniramine / Dextromethorphan / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1339)"

• "Medication, Order: Brompheniramine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1332)"

• "Medication, Order: Butabarbital" using (2.16.840.1.113883.3.464.1003.196.12.1402)"

• "Medication, Order: Carbinoxamine" using (2.16.840.1.113883.3.464.1003.196.12.1306)"

• "Medication, Order: Carisoprodol" using (2.16.840.1.113883.3.464.1003.196.12.1369)"

• "Medication, Order: Chlophedianol / Chlorpheniramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1412)"

• "Medication, Order: Chlophedianol / Dextchlorpheniramine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1447)"

• "Medication, Order: Chlorpheniramine / Dextromethorphan / Phenylephrine" using Phenylephrine (2.16.840.1.113883.3.464.1003.196.12.1312)"

• "Medication, Order: Chlorpheniramine / Dextromethorphan / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1337)"

• "Medication, Order: Chlorpheniramine / Hydrocodone / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1299)"

• "Medication, Order: Chlorpheniramine / Hydrocodone" using (2.16.840.1.113883.3.464.1003.196.12.1330)"
- "Medication, Order: Chlorpheniramine / Phenylephrine" using Phenylephrine (2.16.840.1.113883.3.464.1003.196.12.1343)"
- "Medication, Order: Chlorpheniramine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1315)"
- "Medication, Order: Chlorpheniramine" using (2.16.840.1.113883.3.464.1003.196.12.1352)"
- "Medication, Order: Chlorpropamide" using (2.16.840.1.113883.3.464.1003.196.12.1303)"
- "Medication, Order: Chlorzoxazone" using (2.16.840.1.113883.3.464.1003.196.12.1362)"
- "Medication, Order: Clemastine" using (2.16.840.1.113883.3.464.1003.196.12.1308)"
- "Medication, Order: Clomipramine" using (2.16.840.1.113883.3.464.1003.196.12.1336)"
- "Medication, Order: Conjugated Estrogens / Medroxyprogesterone" using (2.16.840.1.113883.3.464.1003.196.12.1324)"
- "Medication, Order: Conjugated Estrogens" using (2.16.840.1.113883.3.464.1003.196.12.1357)"
- "Medication, Order: Cyclobenzaprine Hydrochloride" using (2.16.840.1.113883.3.464.1003.196.12.1372)"
- "Medication, Order: Cyproheptadine" using (2.16.840.1.113883.3.464.1003.196.12.1277)"
- "Medication, Order: Desiccated Thyroid" using (2.16.840.1.113883.3.464.1003.196.12.1354)"
- "Medication, Order: Desipramine" using (2.16.840.1.113883.3.464.1003.196.12.1278)"
- "Medication, Order: Dexamphetamine / Dextromethorphan / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1426)"
- "Medication, Order: Dexamphetamine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1430)"
- "Medication, Order: Dexamphetamine Maleate / Pseudoephedrine Hydrochloride" using (2.16.840.1.113883.3.464.1003.196.12.1429)"
- "Medication, Order: Dexamphetamine" using (2.16.840.1.113883.3.464.1003.196.12.1375)"
- "Medication, Order: Dechlorpheniramine / Dextromethorphan / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1300)"
• "Medication, Order: Dexchlorpheniramine / Pseudoephedrine" using (2.16.840.1.113883.3.464.1003.196.12.1425)"
• "Medication, Order: Dextromethorphan / Diphenhydramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1397)"
• "Medication, Order: Dextromethorphan / Doxylamine" using (2.16.840.1.113883.3.464.1003.196.12.1452)"
• "Medication, Order: Dicyclomine" using (2.16.840.1.113883.3.464.1003.196.12.1279)"
• "Medication, Order: Dienogest / Estradiol Multiphasic" using (2.16.840.1.113883.3.464.1003.196.12.1398)"
• "Medication, Order: Dimenhydrinate" using (2.16.840.1.113883.3.464.1003.196.12.1500)"
• "Medication, Order: Diphenhydramine / Ibuprofen" using (2.16.840.1.113883.3.464.1003.196.12.1293)"
• "Medication, Order: Diphenhydramine / Phenylephrine" using (2.16.840.1.113883.3.464.1003.196.12.1307)"
• "Medication, Order: Diphenhydramine Hydrochloride" using (2.16.840.1.113883.3.464.1003.196.12.1371)"
• "Medication, Order: Dipyridamole" using (2.16.840.1.113883.3.464.1003.196.12.1349)"
• "Medication, Order: Disopyramide" using (2.16.840.1.113883.3.464.1003.196.12.1311)"
• "Medication, Order: Drospirenone / Estradiol" using (2.16.840.1.113883.3.464.1003.196.12.1410)"
• "Medication, Order: Esterified Estrogens / Methyltestosterone" using (2.16.840.1.113883.3.464.1003.196.12.1320)"
• "Medication, Order: Esterified Estrogens" using (2.16.840.1.113883.3.464.1003.196.12.1419)"
• "Medication, Order: Estradiol / Norethindrone" using (2.16.840.1.113883.3.464.1003.196.12.1323)"
• "Medication, Order: Estradiol" using (2.16.840.1.113883.3.464.1003.196.12.1365)"
• "Medication, Order: Estropipate" using (2.16.840.1.113883.3.464.1003.196.12.1319)"
• "Medication, Order: Glyburide / Metformin" using (2.16.840.1.113883.3.464.1003.196.12.1360)"
• "Medication, Order: Glyburide" using (2.16.840.1.113883.3.464.1003.196.12.1368)"
• "Medication, Order: Guanfacine" using (2.16.840.1.113883.3.464.1003.196.12.1341)"
• "Medication, Order: High-Risk Medications With Days Supply Criteria" using (2.16.840.1.113883.3.464.1003.196.12.1254)"
• "Medication, Order: Hydrochlorothiazide / Methyldopa" using (2.16.840.1.113883.3.464.1003.196.12.1414)"
• "Medication, Order: Hydroxyzine" using (2.16.840.1.113883.3.464.1003.196.12.1374)"
• "Medication, Order: Hyoscyamine / Methenamine / Mblue / Phenyl Salicyl / Sodium Biphosphat" using (2.16.840.1.113883.3.464.1003.196.12.1504)"
• "Medication, Order: Hyoscyamine / Methenamine / Mblue / Phenyl Salicyl" using (2.16.840.1.113883.3.464.1003.196.12.1503)"
• "Medication, Order: Hyoscyamine / Methenamine / Mblue / Sodium Biphosphat" using (2.16.840.1.113883.3.464.1003.196.12.1505)"
• "Medication, Order: Hyoscyamine" using (2.16.840.1.113883.3.464.1003.196.12.1501)"
• "Medication, Order: Imipramine" using (2.16.840.1.113883.3.464.1003.196.12.1359)"
• "Medication, Order: Indomethacin" using (2.16.840.1.113883.3.464.1003.196.12.1366)"
• "Medication, Order: Isoxsuprine" using (2.16.840.1.113883.3.464.1003.196.12.1422)"
• "Medication, Order: Ketorolac Tromethamine" using (2.16.840.1.113883.3.464.1003.196.12.1364)"
• "Medication, Order: List of Single RxNorm Code Concepts for High Risk Drugs for the Elderly" using Elderly (2.16.840.1.113883.3.464.1003.196.12.1272)"
• "Medication, Order: Meclizine" using (2.16.840.1.113883.3.464.1003.196.12.1506)"
• "Medication, Order: Megestrol" using (2.16.840.1.113883.3.464.1003.196.12.1342)"
• "Medication, Order: Meperidine" using (2.16.840.1.113883.3.464.1003.196.12.1351)"
• "Medication, Order: Meprobamate" using (2.16.840.1.113883.3.464.1003.196.12.1284)"
• "Medication, Order: Metaxalone" using (2.16.840.1.113883.3.464.1003.196.12.1358)"
• "Medication, Order: Methocarbamol" using (2.16.840.1.113883.3.464.1003.196.12.1370)"
• "Medication, Order: Methyldopa" using (2.16.840.1.113883.3.464.1003.196.12.1331)"
• "Medication, Order: Nifedipine" using (2.16.840.1.113883.3.464.1003.196.12.1353)"
• "Medication, Order: Nonbenzodiazepine hypnotics" using (2.16.840.1.113883.3.464.1003.196.12.1480)"
• "Medication, Order: Nortriptyline" using (2.16.840.1.113883.3.464.1003.196.12.1507)"
• "Medication, Order: Paroxetine" using (2.16.840.1.113883.3.464.1003.196.12.1508)"
• "Medication, Order: Phenobarbital" using (2.16.840.1.113883.3.464.1003.196.12.1348)"
• "Medication, Order: Promethazine Hydrochloride" using (2.16.840.1.113883.3.464.1003.196.12.1367)"
• "Medication, Order: Protriptyline" using (2.16.840.1.113883.3.464.1003.196.12.1509)"
• "Medication, Order: Pseudoephedrine / Triprolidine" using (2.16.840.1.113883.3.464.1003.196.12.1345)"
• "Medication, Order: Trihexyphenidyl" using (2.16.840.1.113883.3.464.1003.196.12.1334)"
• "Medication, Order: Trimipramine" using (2.16.840.1.113883.3.464.1003.196.12.1285)"
• "Medication, Order: Triprolidine" using (2.16.840.1.113883.3.464.1003.196.12.1408)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
Value Sets: CMS 159V7 Depression Remission at Twelve Months

<table>
<thead>
<tr>
<th>Description</th>
<th>The percentage of adolescent patients 12 to 17 years of age and adult patients 18 years of age or older with major depression or dysthymia who reached remission 12 months (+/- 60 days) after an index event.</th>
</tr>
</thead>
</table>

- codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
- code "Patient Health Questionnaire 9 item (PHQ-9) total score [Reported]" using "LOINC version 2.63 Code (44261-6)"
- valueset "Bipolar Disorder" using "2.16.840.1.113883.3.67.1.101.1.128"
- valueset "Care Services in Long-Term Residential Facility" using "2.16.840.1.113883.3.464.1003.101.12.1014"
- valueset "Contact or Office Visit" using "2.16.840.1.113762.1.4.1080.5"
- valueset "Dysthymia" using "2.16.840.1.113883.3.67.1.101.1.254"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Major Depression Including Remission" using "2.16.840.113883.3.67.1.101.3.2444"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1.1"
- valueset "Palliative care encounter" using "2.16.840.1.113883.3.600.1.1575"
- valueset "Palliative Care" using "2.16.840.1.113883.3.600.1.1579"
- valueset "Payer" using "2.16.840.1.114222.4.11.3591"
- valueset "Personality Disorder" using "2.16.840.1.113883.3.67.1.101.1.246"
- valueset "Pervasive Developmental Disorder" using "2.16.840.1.113883.3.464.1003.105.12.1152"
- valueset "Race" using "2.16.840.1.114222.4.11.836"
- "Diagnosis: Bipolar Disorder" using (2.16.840.1.113883.3.67.1.101.1.128)"
- "Diagnosis: Dysthymia" using (2.16.840.1.113883.3.67.1.101.1.254)"
- "Diagnosis: Major Depression Including Remission" using "(2.16.840.1.113883.3.67.1.101.3.2444)"
- "Diagnosis: Personality Disorder" using (2.16.840.1.113883.3.67.1.101.1.246)"
- "Diagnosis: Pervasive Developmental Disorder" using (2.16.840.1.113883.3.464.1003.105.12.1152)"
- "Diagnosis: Schizophrenia or Psychotic Disorder" using "(2.16.840.1.113883.3.464.1003.105.12.1104)"
• "Encounter, Performed: Care Services in Long-Term Residential Facility" using (2.16.840.1.113883.3.464.1003.101.12.1014)"
• "Encounter, Performed: Contact or Office Visit" using (2.16.840.1.113762.1.4.1080.5)"
• "Encounter, Performed: Palliative care encounter" using (2.16.840.1.113883.3.600.1.1575)"
• "Intervention, Order: Palliative Care" using (2.16.840.1.113883.3.600.1.1579)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
• "Assessment, Performed: Patient Health Questionnaire 9 item (PHQ-9) total score [Reported]" using "Patient Health Questionnaire 9 item (PHQ-9) total score [Reported] (LOINC version 2.63 Code 44261-6)"
• Valueset "Schizophrenia or Psychotic Disorder" using "2.16.840.1.113883.3.464.1003.105.12.1104"

Value Sets: CMS 160V7 Depression Utilization of the PHQ-9 Tool

<table>
<thead>
<tr>
<th>Description:</th>
<th>The percentage of adolescent patients 12 to 17 years of age and adult patients age 18 and older with the diagnosis of major depression or dysthymia who have a completed PHQ-9 during each applicable 4-month period in which there was a qualifying depression encounter.</th>
</tr>
</thead>
</table>

• codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
• code "Patient Health Questionnaire 9 item (PHQ-9) total score [Reported]" using "LOINC version 2.63 Code (44261-6)"
• valueset "Bipolar Disorder" using "2.16.840.1.113883.3.67.1.101.1.128"
• valueset "Care Services in Long-Term Residential Facility" using "2.16.840.1.113883.3.464.1003.101.12.1014"
• valueset "Contact or Office Visit" using "2.16.840.1.113762.1.4.1080.5"
• valueset "Dysthymia" using "2.16.840.1.113883.3.67.1.101.1.254"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Major Depression Including Remission" using "2.16.840.113883.3.67.1.101.3.2444"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Palliative care encounter" using "2.16.840.1.113883.3.600.1.1575"
• valueset "Palliative Care" using "2.16.840.1.113883.3.600.1.1579"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Personality Disorder" using "2.16.840.1.113883.3.67.1.101.1.246"
• valueset "Pervasive Developmental Disorder" using "2.16.840.1.113883.3.464.1003.105.12.1152"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• valueset "Schizophrenia or Psychotic Disorder" using "2.16.840.1.113883.3.464.1003.105.12.1104"
• "Diagnosis: Bipolar Disorder" using (2.16.840.1.113883.3.67.1.101.1.128)"
• "Diagnosis: Dysthymia" using (2.16.840.1.113883.3.67.1.101.1.254)"
• "Diagnosis: Major Depression Including Remission" using (2.16.840.1.113883.3.67.1.101.3.2444)"
• "Diagnosis: Personality Disorder" using (2.16.840.1.113883.3.67.1.101.1.246)"
• "Diagnosis: Pervasive Developmental Disorder" using (2.16.840.1.113883.3.67.1.101.1.246)"
• "Diagnosis: Schizophrenia or Psychotic Disorder" using (2.16.840.1.113883.3.67.1.101.1.246)"
• "Encounter, Performed: Care Services in Long-Term Residential Facility" using (2.16.840.1.113883.3.464.1003.105.12.1014)"
• "Encounter, Performed: Contact or Office Visit" using (2.16.840.1.113883.3.464.1003.105.12.1014)"
• "Encounter, Performed: Palliative care encounter" using (2.16.840.1.113883.3.600.1.1575)"
• “Encounter, Performed: Telehealth Services: 2.16.840.1.113883.3.464.1003.101.12.1031
• "Intervention, Order: Palliative Care" using (2.16.840.1.113883.3.600.1.1579)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
"Assessment, Performed: Patient Health Questionnaire 9 item (PHQ-9) total score [Reported]" using "Patient Health Questionnaire 9 item (PHQ-9) total score [Reported] (LOINC version 2.63 Code 44261-6)"

**Value Sets: CMS 161V8 Adult MDD: Suicide Risk Assessment**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients aged 18 years and older with a diagnosis of MDD with a suicide risk assessment completed during the visit in which a new diagnosis or recurrent episode was identified.</th>
</tr>
</thead>
</table>

- "Encounter, Performed: ED Visit" using (2.16.840.1.113883.3.464.1003.101.12.1010)"
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
- "Encounter, Performed: Outpatient Consultation" using (2.16.840.1.113883.3.464.1003.101.12.1008)"
- "Encounter, Performed: Psych Visit - Diagnostic Evaluation" using (2.16.840.1.113883.3.526.3.1492)"
- "Encounter, Performed: Psych Visit - Psychotherapy" using (2.16.840.1.113883.3.526.3.1496)"
- "Encounter, Performed: Psychoanalysis" using (2.16.840.1.113883.3.526.3.1141)"
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
- "Intervention, Performed: Suicide risk assessment (procedure)" using "Suicide risk assessment (procedure) (SNOMEDCT version 2017-09 Code 225337009)"
- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Suicide risk assessment (procedure)" using "SNOMEDCT version 2017-09 Code (225337009)"
- valueset "ED Visit" using "2.16.840.1.113883.3.464.1003.101.12.1010"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Major Depressive Disorder-Active" using "2.16.840.1.113883.3.526.3.1491"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Outpatient Consultation" using "2.16.840.1.113883.3.464.1003.101.12.1008"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Psych Visit - Diagnostic Evaluation" using "2.16.840.1.113883.3.526.3.1492"
• valueset "Psych Visit - Psychotherapy" using "2.16.840.1.113883.3.526.3.1496"
• valueset "Psychoanalysis" using "2.16.840.1.113883.3.526.3.1141"
• valueset "Race" using "2.16.840.1.114222.4.11.836"

Value Sets: CMS 165V8 Controlling High Blood Pressure

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patients 18-85 years of age who had a diagnosis of hypertension and whose blood pressure was adequately controlled (&lt;140/90 mmHg) during the measurement period.</th>
</tr>
</thead>
</table>
• codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
• codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
• code "Diastolic blood pressure" using "LOINC version 2.63 Code (8462-4)"
• code "Discharge to healthcare facility for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428371000124100)"
• code "Discharge to home for hospice care (procedure)" using "SNOMEDCT version 2017-09 Code (428361000124107)"
• code "Systolic blood pressure" using "LOINC version 2.63 Code (8480-6)"
• valueset "Adult Outpatient Visit" using "2.16.840.1.113883.3.464.1003.101.12.1065"
• valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.3.1240"
• valueset "Chronic Kidney Disease, Stage 5" using "2.16.840.1.113883.3.526.3.1002"
• valueset "Dialysis Services" using "2.16.840.1.113883.3.464.1003.109.12.1013"
• valueset "Encounter Inpatient" using "2.16.840.1.113883.3.666.5.307"
• valueset "End Stage Renal Disease" using "2.16.840.1.113883.3.526.3.353"
• valueset "ESRD Monthly Outpatient Services" using "2.16.840.1.113883.3.464.1003.109.12.1014"
• valueset "Essential Hypertension" using "2.16.840.1.113883.3.464.1003.104.12.1011"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Home Healthcare Services" using "2.16.840.1.113883.3.464.1003.101.12.1016"
• valueset "Hospice care ambulatory" using "2.16.840.1.113762.1.4.1108.15"
• valueset "Kidney Transplant Recipient" using "2.16.840.1.113883.3.464.1003.109.12.1029"
• valueset "Kidney Transplant" using "2.16.840.1.113883.3.464.1003.109.12.1012"
• valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1.1"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Pregnancy" using "2.16.840.1.113883.3.526.3.378"
• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
• valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• "Diagnosis: Chronic Kidney Disease, Stage 5" using (2.16.840.1.113883.3.526.3.1002)
• "Diagnosis: End Stage Renal Disease" using (2.16.840.1.113883.3.526.3.353)
• "Diagnosis: Essential Hypertension" using (2.16.840.1.113883.3.464.1003.104.12.1011)
• "Diagnosis: Kidney Transplant Recipient" using (2.16.840.1.113883.3.464.1003.109.12.1029)
• "Diagnosis: Pregnancy" using (2.16.840.1.113883.3.526.3.378)
• "Encounter, Performed: Adult Outpatient Visit" using (2.16.840.1.113883.3.464.1003.101.12.1065)
• "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883.3.526.3.1240)
• "Encounter, Performed: Encounter Inpatient" using (2.16.840.1.113883.3.666.5.307)
• "Encounter, Performed: ESRD Monthly Outpatient Services" using (2.16.840.1.113883.3.464.1003.109.12.1014)
• "Encounter, Performed: Home Healthcare Services" using (2.16.840.1.113883.3.464.1003.101.12.1016)
• "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using Up (2.16.840.1.113883.3.464.1003.101.12.1025)"
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1023)"
- "Intervention, Order: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Intervention, Performed: Hospice care ambulatory" using (2.16.840.1.113762.1.4.1108.15)"
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
- "Physical Exam, Performed: Diastolic blood pressure" using (LOINC version 2.63 Code 8462-4)"
- "Physical Exam, Performed: Systolic blood pressure" using (LOINC version 2.63 Code 8480-6)"

**Value Sets: CMS 177V8 Child and Adolescent MDD: Suicide Risk Assessment**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder with an assessment for suicide risk.</th>
</tr>
</thead>
</table>

- codesystem "SNOMEDCT" using "2.16.840.1.113883.6.96 version 2017-09"
- code "Suicide risk assessment (procedure)" using "SNOMEDCT version 2017-09 Code (225337009)"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Group Psychotherapy" using "2.16.840.1.113883.3.526.3.1187"
• valueset "Major Depressive Disorder-Active" using "2.16.840.1.113883.3.526.3.1491"
• valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Outpatient Consultation" using "2.16.840.1.113883.3.464.1003.101.12.1008"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "Psych Visit - Diagnostic Evaluation" using "2.16.840.1.113883.3.526.3.1492"
• valueset "Psych Visit - Family Psychotherapy" using "2.16.840.1.113883.3.526.3.1018"
• valueset "Psych Visit - Psychotherapy" using "2.16.840.1.113883.3.526.3.1496"
• valueset "Psychoanalysis" using "2.16.840.1.113883.3.526.3.1141"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• "Encounter, Performed: Group Psychotherapy" using (2.16.840.1.113883.3.526.3.1187)"
• "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)"
• "Encounter, Performed: Outpatient Consultation" using (2.16.840.1.113883.3.464.1003.101.12.1008)"
• "Encounter, Performed: Psych Visit - Diagnostic Evaluation" using "(2.16.840.1.113883.3.526.3.1492)"
• "Encounter, Performed: Psych Visit - Family Psychotherapy" using Psychotherapy (2.16.840.1.113883.3.526.3.1018)"
• "Encounter, Performed: Psych Visit - Psychotherapy" using (2.16.840.1.113883.3.526.3.1496)"
• "Encounter, Performed: Psychoanalysis" using (2.16.840.1.113883.3.526.3.1141)"
• “Encounter, Performed: Telehealth Services: 2.16.840.1.113883.3.464.1003.101.12.1031" "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
• "Intervention,Performed:Suicide risk assessment (procedure)" using (SNOMEDCT version 2017-09 Code 225337009)"

**Value Sets: CMS 347V3 Statin Therapy for the Prevention and Treatment of Cardiovascular Disease**

<table>
<thead>
<tr>
<th>Description</th>
<th>Percentage of the following patients - all considered at high risk of cardiovascular events - who were prescribed or were on statin therapy during the measurement period:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Adults aged &gt;= 21 years who were previously diagnosed with or currently have an active diagnosis of clinical ASCVD.</td>
</tr>
<tr>
<td></td>
<td>• Adults aged &gt;= 21 years who have ever had a fasting or direct low-density lipoprotein cholesterol (LDL-C) level &gt;= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia.</td>
</tr>
<tr>
<td></td>
<td>• Adults aged 40-75 years with a diagnosis of diabetes with a fasting or direct LDL-C level of 70-189 mg/dL.</td>
</tr>
</tbody>
</table>

• valueset "Annual Wellness Visit" using "2.16.840.1.113883.3.526.2.1363"
• valueset "Atherosclerosis and Peripheral Arterial Disease" using "2.16.840.1.113762.1.4.1047.21"
• valueset "Breastfeeding" using "2.16.840.1.113762.1.4.1047.73"
• valueset "CABG Surgeries" using "2.16.840.1.113883.3.666.5.694"
• valueset "Carotid Intervention" using "2.16.840.1.113883.3.117.1.7.1.204"
• valueset "Cerebrovascular disease, Stroke, TIA" using "2.16.840.1.113883.3.117.1.7.1.204"
• valueset "Diabetes" using "2.16.840.1.113883.3.464.1003.103.12.1001"
• valueset "End Stage Renal Disease" using "2.16.840.1.113883.3.526.3.353"
• valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
• valueset "Hepatitis A" using "2.16.840.1.113883.3.464.1003.110.12.1024"
• valueset "Hepatitis B" using "2.16.840.1.113883.3.67.1.101.1.269"
• valueset "High intensity statin therapy" using "2.16.840.1.113762.1.4.1047.97"
• valueset "Hypercholesterolemia" using "2.16.840.1.113762.1.4.1047.100"
• valueset "Ischemic heart disease or coronary occlusion, rupture, or thrombosis" using "2.16.840.1.113762.1.4.1047.46"
• valueset "Liver Disease" using "2.16.840.1.113762.1.4.1047.42"
• valueset "Low intensity statin therapy" using "2.16.840.1.113762.1.4.1047.107"
• valueset "Moderate intensity statin therapy" using "2.16.840.1.113762.1.4.1047.98"
• valueset "Myocardial Infarction" using "2.16.840.1.113883.3.526.3.403"
• valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.11.1005"
• valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
• valueset "Outpatient Consultation" using "2.16.840.1.113883.3.464.1003.101.11.1040"
• valueset "Outpatient Encounters for Preventive Care" using "2.16.840.1.113762.1.4.1047.9"
• valueset "Palliative care encounter" using "2.16.840.1.113883.3.600.1.1575"
• valueset "Palliative or Hospice Care" using "2.16.840.1.113883.3.600.1.1579"
• valueset "Payer" using "2.16.840.1.114222.4.11.3591"
• valueset "PCI" using "2.16.840.1.113762.1.4.1045.67"
• valueset "Pregnancy Dx" using "2.16.840.1.113883.3.600.1.1623"
• valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.11.125"
• valueset "Preventive Care Services - Other" using "2.16.840.1.113883.3.464.1003.101.12.1030"
• valueset "Preventive Care Services-Individual Counseling" using "2.16.840.1.113883.3.464.1003.101.12.1026"
• valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.11.1115"
• valueset "Race" using "2.16.840.1.114222.4.11.836"
• valueset "Rhabdomyolysis" using "2.16.840.1.113762.1.4.1047.102"
• valueset "Stable and Unstable Angina" using "2.16.840.1.113762.1.4.1047.47"
• valueset "Statin Allergen" using "2.16.840.1.113883.3.117.1.7.1.423"
• "Adverse Event: Statin Allergen" using (2.16.840.1.113883.3.117.1.7.1.423)"
• "Allergy/Intolerance: Statin Allergen" using (2.16.840.1.113883.3.117.1.7.1.423)"
• "Diagnosis: Atherosclerosis and Peripheral Arterial Disease" using (2.16.840.1.113762.1.4.1047.21)"
• "Diagnosis: Breastfeeding" using (2.16.840.1.113762.1.4.1047.73)"
• "Diagnosis: Cerebrovascular disease, Stroke, TIA" using (2.16.840.1.113762.1.4.1047.44)"
• "Diagnosis: Diabetes" using (2.16.840.1.113883.3.464.1003.103.12.1001)"
• "Diagnosis: End Stage Renal Disease" using (2.16.840.1.113883.3.526.3.353)"
• "Diagnosis: Hepatitis A" using (2.16.840.1.113883.3.464.1003.110.12.1024)"
• "Diagnosis: Hepatitis B" using (2.16.840.1.113883.3.67.1.101.1.269)"
• "Diagnosis: Hypercholesterolemia" using (2.16.840.1.113762.1.4.1047.100)"
• "Diagnosis: Ischemic heart disease or coronary occlusion, rupture, or thrombosis" using (2.16.840.1.113762.1.4.1047.46)"
• "Diagnosis: Liver Disease" using (2.16.840.1.113762.1.4.1047.42)"
• "Diagnosis: Myocardial Infarction" using (2.16.840.1.113883.3.526.3.403)"
• "Diagnosis: Pregnancy Dx" using (2.16.840.1.113883.3.600.1.1623)"
• "Diagnosis: Rhabdomyolysis" using (2.16.840.1.113762.1.4.1047.102)"
• "Diagnosis: Stable and Unstable Angina" using (2.16.840.1.113762.1.4.1047.47)"
• "Encounter, Performed: Annual Wellness Visit" using (2.16.840.1.113883.3.526.2.1363)"
• "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.11.1005)"
• "Encounter, Performed: Outpatient Consultation" using (2.16.840.1.113883.3.464.1003.101.11.1040)"
• "Encounter, Performed: Outpatient Encounters for Preventive Care" using (2.16.840.1.113762.1.4.1047.9)"
• "Encounter, Performed: Palliative care encounter" using "(2.16.840.1.113883.3.600.1.1575)"
• "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.11.1125)"
• "Encounter, Performed: Preventive Care Services - Other" using (2.16.840.1.113883.3.464.1003.101.12.1030)"
• "Encounter, Performed: Preventive Care Services-Individual Counseling" using (2.16.840.1.113883.3.464.1003.101.12.1026)"
• "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.11.1115)"
• "Intervention, Order: Palliative or Hospice Care" using (2.16.840.1.113883.3.600.1.1579)"
• "Laboratory Test, Performed: LDL-c: 2.16.840.1.113883.3.117.1.7.1.215"
• "Medication, Active: High intensity statin therapy" using (2.16.840.1.113762.1.4.1047.97)"
• "Medication, Active: Low intensity statin therapy" using (2.16.840.1.113762.1.4.1047.107)"
• "Medication, Active: Moderate intensity statin therapy" using (2.16.840.1.113762.1.4.1047.98)"
• "Medication, Order: High intensity statin therapy" using (2.16.840.1.113762.1.4.1047.97)"
• "Medication, Order: Low intensity statin therapy" using (2.16.840.1.113762.1.4.1047.107)"
• "Medication, Order: Moderate intensity statin therapy" using (2.16.840.1.113762.1.4.1047.98)"
• "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.11.837)"
• "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.11.3591)"
• "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.11.836)"
• "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)"
• "Procedure, Performed: CABG Surgeries" using (2.16.840.1.113883.3.666.5.694)"
• "Procedure, Performed: Carotid Intervention" using (2.16.840.1.113883.3.117.1.7.1.204)"
• "Procedure, Performed: PCI" using (2.16.840.1.113762.1.4.1045.67)"

**Value Sets: CMS 349V2 HIV Screening**

| Description | Percentage of patients 15-65 years of age who have been tested for HIV within that age range. |

- codesystem "LOINC" using "2.16.840.1.113883.6.1 version 2.63"
- code "HIV 1 and 2 tests - Meaningful Use set" using "LOINC version 2.63 Code (75622-1)"
- valueset "Conditions Due To Human Immunodeficiency Virus (HIV)" using "2.16.840.1.113762.1.4.1056.54"
- valueset "Ethnicity" using "2.16.840.1.114222.4.11.837"
- valueset "Human Immunodeficiency Virus (HIV) Laboratory Test Codes (Ab and Ag)" using "2.16.840.1.113762.1.4.1056.50"
- valueset "Office Visit" using "2.16.840.1.113883.3.464.1003.101.12.1001"
- valueset "ONC Administrative Sex" using "2.16.840.1.113762.1.4.1"
- valueset "Payer" using "2.16.840.1.114222.4.1.11.3591"
- valueset "Preventive Care Services - Established Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1025"
- valueset "Preventive Care Services, Initial Office Visit, 0 to 17" using "2.16.840.1.113883.3.464.1003.101.12.1022"
- valueset "Preventive Care Services-Initial Office Visit, 18 and Up" using "2.16.840.1.113883.3.464.1003.101.12.1023"
- valueset "Preventive Care, Established Office Visit, 0 to 17" using "2.16.840.1.113883.3.464.1003.101.12.1024"
- valueset "Race" using "2.16.840.1.114222.4.1.11.836"
- "Diagnosis: Conditions Due To Human Immunodeficiency Virus (HIV)" using (2.16.840.1.113762.1.4.1056.54)
- "Encounter, Performed: Office Visit" using (2.16.840.1.113883.3.464.1003.101.12.1001)
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1025)
- "Encounter, Performed: Preventive Care Services, Initial Office Visit, 0 to 17" using (2.16.840.1.113883.3.464.1003.101.12.1022)
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using (2.16.840.1.113883.3.464.1003.101.12.1023)
- "Encounter, Performed: Preventive Care, Established Office Visit, 0 to 17" using (2.16.840.1.113883.3.464.1003.101.12.1024)
- "Laboratory Test, Performed: Human Immunodeficiency Virus (HIV) Laboratory Test Codes (Ab and Ag)" using (2.16.840.1.113762.1.4.1056.50)
- "Patient Characteristic Ethnicity: Ethnicity" using (2.16.840.1.114222.4.1.11.837)
- "Patient Characteristic Payer: Payer" using (2.16.840.1.114222.4.1.11.3591)
- "Patient Characteristic Race: Race" using (2.16.840.1.114222.4.1.11.836)
- "Patient Characteristic Sex: ONC Administrative Sex" using (2.16.840.1.113762.1.4.1)
- "Laboratory Test, Performed: HIV 1 and 2 tests - Meaningful Use set" using LOINC version 2.63 Code 75622-1), (https://r.details.loinc.org/LOINC/75622-1.html)
The Resource and Patient Management (RPMS) system is a United States Department of Health and Human Services (HHS), Indian Health Service (IHS) information system that is **FOR OFFICIAL USE ONLY**. The RPMS system is subject to monitoring; therefore, no expectation of privacy shall be assumed. Individuals found performing unauthorized activities are subject to disciplinary action including criminal prosecution.

All users (Contractors and IHS Employees) of RPMS will be provided a copy of the Rules of Behavior (ROB) and must acknowledge that they have received and read them prior to being granted access to a RPMS system, in accordance IHS policy.

- For a listing of general ROB for all users, see the most recent edition of *IHS General User Security Handbook* (SOP 06-11a).
- For a listing of system administrators/managers rules, see the most recent edition of the *IHS Technical and Managerial Handbook* (SOP 06-11b).

Both documents are available at this IHS website: [https://home.ihs.gov/security/index.cfm](https://home.ihs.gov/security/index.cfm).

**Note:** Users must be logged on to the IHS D1 Intranet to access these documents.

The ROB listed in the following sections are specific to RPMS.

### F.1 All RPMS Users

In addition to these rules, each application may include additional ROB that may be defined within the documentation of that application (e.g., Dental, Pharmacy).

#### F.1.1 Access

RPMS users shall

- Only use data for which you have been granted authorization.
- Only give information to personnel who have access authority and have a need to know.
- Always verify a caller’s identification and job purpose with your supervisor or the entity provided as employer before providing any type of information system access, sensitive information, or nonpublic agency information.
- Be aware that personal use of information resources is authorized on a limited basis within the provisions *Indian Health Manual* Part 8, “Information Resources Management,” Chapter 6, “Limited Personal Use of Information Technology Resources.”
RPMS users shall not

- Retrieve information for someone who does not have authority to access the information.
- Access, research, or change any user account, file, directory, table, or record not required to perform their official duties.
- Store sensitive files on a PC hard drive, or portable devices or media, if access to the PC or files cannot be physically or technically limited.
- Exceed their authorized access limits in RPMS by changing information or searching databases beyond the responsibilities of their jobs or by divulging information to anyone not authorized to know that information.

F.1.2 Information Accessibility

RPMS shall restrict access to information based on the type and identity of the user. However, regardless of the type of user, access shall be restricted to the minimum level necessary to perform the job.

RPMS users shall

- Access only those documents they created and those other documents to which they have a valid need-to-know and to which they have specifically granted access through an RPMS application based on their menus (job roles), keys, and FileMan access codes. Some users may be afforded additional privileges based on the functions they perform, such as system administrator or application administrator.
- Acquire a written preauthorization in accordance with IHS policies and procedures prior to interconnection to or transferring data from RPMS.

F.1.3 Accountability

RPMS users shall

- Behave in an ethical, technically proficient, informed, and trustworthy manner.
- Log out of the system whenever they leave the vicinity of their personal computers (PCs).
- Be alert to threats and vulnerabilities in the security of the system.
- Report all security incidents to their local Information System Security Officer (ISSO)
- Differentiate tasks and functions to ensure that no one person has sole access to or control over important resources.
- Protect all sensitive data entrusted to them as part of their government employment.
• Abide by all Department and Agency policies and procedures and guidelines related to ethics, conduct, behavior, and information technology (IT) information processes.

F.1.4 Confidentiality

RPMS users shall

• Be aware of the sensitivity of electronic and hard copy information, and protect it accordingly.
• Store hard copy reports/storage media containing confidential information in a locked room or cabinet.
• Erase sensitive data on storage media prior to reusing or disposing of the media.
• Protect all RPMS terminals from public viewing at all times.
• Abide by all Health Insurance Portability and Accountability Act (HIPAA) regulations to ensure patient confidentiality.

RPMS users shall not

• Allow confidential information to remain on the PC screen when someone who is not authorized to that data is in the vicinity.
• Store sensitive files on a portable device or media without encrypting.

F.1.5 Integrity

RPMS users shall

• Protect their systems against viruses and similar malicious programs.
• Observe all software license agreements.
• Follow industry standard procedures for maintaining and managing RPMS hardware, operating system software, application software, and/or database software and database tables.
• Comply with all copyright regulations and license agreements associated with RPMS software.

RPMS users shall not

• Violate federal copyright laws.
• Install or use unauthorized software within the system libraries or folders.
• Use freeware, shareware, or public domain software on/with the system without their manager’s written permission and without scanning it for viruses first.
F.1.6 System Logon
RPMS users shall
- Have a unique User Identification/Account name and password.
- Be granted access based on authenticating the account name and password entered.
- Be locked out of an account after five successive failed login attempts within a specified time period (e.g., one hour).

F.1.7 Passwords
RPMS users shall
- Change passwords a minimum of every 90 days.
- Create passwords with a minimum of eight characters.
- If the system allows, use a combination of alpha-numeric characters for passwords, with at least one uppercase letter, one lower case letter, and one number. It is recommended, if possible, that a special character also be used in the password.
- Change vendor-supplied passwords immediately.
- Protect passwords by committing them to memory or store them in a safe place (do not store passwords in login scripts or batch files).
- Change passwords immediately if password has been seen, guessed, or otherwise compromised, and report the compromise or suspected compromise to their ISSO.
- Keep user identifications (IDs) and passwords confidential.

RPMS users shall not
- Use common words found in any dictionary as a password.
- Use obvious readable passwords or passwords that incorporate personal data elements (e.g., user’s name, date of birth, address, telephone number, or social security number; names of children or spouses; favorite band, sports team, or automobile; or other personal attributes).
- Share passwords/IDs with anyone or accept the use of another’s password/ID, even if offered.
- Reuse passwords. A new password must contain no more than five characters per eight characters from the previous password.
- Post passwords.
- Keep a password list in an obvious place, such as under keyboards, in desk drawers, or in any other location where it might be disclosed.
• Give a password out over the phone.

F.1.8 Backups
RPMS users shall
• Plan for contingencies such as physical disasters, loss of processing, and disclosure of information by preparing alternate work strategies and system recovery mechanisms.
• Make backups of systems and files on a regular, defined basis.
• If possible, store backups away from the system in a secure environment.

F.1.9 Reporting
RPMS users shall
• Contact and inform their ISSO that they have identified an IT security incident and begin the reporting process by providing an IT Incident Reporting Form regarding this incident.
• Report security incidents as detailed in the IHS Incident Handling Guide (SOP 05-03).

RPMS users shall not
• Assume that someone else has already reported an incident. The risk of an incident going unreported far outweighs the possibility that an incident gets reported more than once.

F.1.10 Session Timeouts
RPMS system implements system-based timeouts that back users out of a prompt after no more than 5 minutes of inactivity.

RPMS users shall
• Utilize a screen saver with password protection set to suspend operations at no greater than 10 minutes of inactivity. This will prevent inappropriate access and viewing of any material displayed on the screen after some period of inactivity.

F.1.11 Hardware
RPMS users shall
• Avoid placing system equipment near obvious environmental hazards (e.g., water pipes).
• Keep an inventory of all system equipment.
• Keep records of maintenance/repairs performed on system equipment.
RPMS users shall not
• Eat or drink near system equipment.

F.1.12 Awareness
RPMS users shall
• Participate in organization-wide security training as required.
• Read and adhere to security information pertaining to system hardware and software.
• Take the annual information security awareness.
• Read all applicable RPMS manuals for the applications used in their jobs.

F.1.13 Remote Access
Each subscriber organization establishes its own policies for determining which employees may work at home or in other remote workplace locations. Any remote work arrangement should include policies that
• Are in writing.
• Provide authentication of the remote user through the use of ID and password or other acceptable technical means.
• Outline the work requirements and the security safeguards and procedures the employee is expected to follow.
• Ensure adequate storage of files, removal, and nonrecovery of temporary files created in processing sensitive data, virus protection, and intrusion detection, and provide physical security for government equipment and sensitive data.
• Establish mechanisms to back up data created and/or stored at alternate work locations.
Remote RPMS users shall
• Remotely access RPMS through a virtual private network (VPN) whenever possible. Use of direct dial in access must be justified and approved in writing and its use secured in accordance with industry best practices or government procedures.
Remote RPMS users shall not
• Disable any encryption established for network, internet, and Web browser communications.
F.2  RPMS Developers

RPMS developers shall

• Always be mindful of protecting the confidentiality, availability, and integrity of RPMS when writing or revising code.

• Always follow the IHS RPMS Programming Standards and Conventions (SAC) when developing for RPMS.

• Only access information or code within the namespaces for which they have been assigned as part of their duties.

• Remember that all RPMS code is the property of the U.S. Government, not the developer.

• Not access live production systems without obtaining appropriate written access, and shall only retain that access for the shortest period possible to accomplish the task that requires the access.

• Observe separation of duties policies and procedures to the fullest extent possible.

• Document or comment all changes to any RPMS software at the time the change or update is made. Documentation shall include the programmer’s initials, date of change, and reason for the change.

• Use checksums or other integrity mechanism when releasing their certified applications to assure the integrity of the routines within their RPMS applications.

• Follow industry best standards for systems they are assigned to develop or maintain, and abide by all Department and Agency policies and procedures.

• Document and implement security processes whenever available.

RPMS developers shall not

• Write any code that adversely impacts RPMS, such as backdoor access, “Easter eggs,” time bombs, or any other malicious code or make inappropriate comments within the code, manuals, or help frames.

• Grant any user or system administrator access to RPMS unless proper documentation is provided.

• Release any sensitive agency or patient information.

F.3  Privileged Users

Personnel who have significant access to processes and data in RPMS, such as, system security administrators, systems administrators, and database administrators, have added responsibilities to ensure the secure operation of RPMS.
Privileged RPMS users shall

- Verify that any user requesting access to any RPMS system has completed the appropriate access request forms.
- Ensure that government personnel and contractor personnel understand and comply with license requirements. End users, supervisors, and functional managers are ultimately responsible for this compliance.
- Advise the system owner on matters concerning information technology security.
- Assist the system owner in developing security plans, risk assessments, and supporting documentation for the certification and accreditation process.
- Ensure that any changes to RPMS that affect contingency and disaster recovery plans are conveyed to the person responsible for maintaining continuity of operations plans.
- Ensure that adequate physical and administrative safeguards are operational within their areas of responsibility and that access to information and data is restricted to authorized personnel on a need-to-know basis.
- Verify that users have received appropriate security training before allowing access to RPMS.
- Implement applicable security access procedures and mechanisms, incorporate appropriate levels of system auditing, and review audit logs.
- Document and investigate known or suspected security incidents or violations and report them to the ISSO, Chief Information Security Officer (CISO), and systems owner.
- Protect the supervisor, superuser, or system administrator passwords.
- Avoid instances where the same individual has responsibility for several functions (i.e., transaction entry and transaction approval).
- Watch for unscheduled, unusual, and unauthorized programs.
- Help train system users on the appropriate use and security of the system.
- Establish protective controls to ensure the accountability, integrity, confidentiality, and availability of the system.
- Replace passwords when a compromise is suspected. Delete user accounts as quickly as possible from the time that the user is no longer authorized system. Passwords forgotten by their owner should be replaced, not reissued.
- Terminate user accounts when a user transfers or has been terminated. If the user has authority to grant authorizations to others, review these other authorizations. Retrieve any devices used to gain access to the system or equipment. Cancel logon IDs and passwords, and delete or reassign related active and backup files.
• Use a suspend program to prevent an unauthorized user from logging on with the current user's ID if the system is left on and unattended.

• Verify the identity of the user when resetting passwords. This can be done either in person or having the user answer a question that can be compared to one in the administrator’s database.

• Shall follow industry best standards for systems they are assigned to, and abide by all Department and Agency policies and procedures.

Privileged RPMS users shall not

• Access any files, records, systems, etc., that are not explicitly needed to perform their duties

• Grant any user or system administrator access to RPMS unless proper documentation is provided.

• Release any sensitive agency or patient information.
# Acronym List

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Term Meaning</th>
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<tbody>
<tr>
<td>AACE/ACE</td>
<td>American Association of Clinical Endocrinologists/American College of Endocrinology</td>
</tr>
<tr>
<td>AAP</td>
<td>American Academy of Pediatrics</td>
</tr>
<tr>
<td>ACC</td>
<td>American College of Cardiology</td>
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<tr>
<td>ACCF</td>
<td>American College of Cardiology Foundation</td>
</tr>
<tr>
<td>ACE</td>
<td>Angiotensin-Converting Enzyme</td>
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<tr>
<td>ACIP</td>
<td>Advisory Committee on Immunization Practices</td>
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<tr>
<td>ACOG</td>
<td>American College of Obstetricians and Gynecologists</td>
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<tr>
<td>ADA</td>
<td>American Diabetes Association</td>
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<tr>
<td>ADE</td>
<td>Adverse Drug Event</td>
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<tr>
<td>AHA</td>
<td>American Heart Association</td>
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<tr>
<td>AHRQ</td>
<td>Agency for Healthcare Research and Quality</td>
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<tr>
<td>AIDS</td>
<td>Acquired Immunodeficiency Syndrome</td>
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<tr>
<td>AMA</td>
<td>Against Medical Advice</td>
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<td>AMI</td>
<td>Acute Myocardial Infarction</td>
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<tr>
<td>AOD</td>
<td>Alcohol or Other Drug abuse</td>
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<tr>
<td>ARB</td>
<td>Angiotensin Receptor Blocker</td>
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<td>ART</td>
<td>Antiretroviral therapy</td>
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<td>ASCVD</td>
<td>Atherosclerotic Cardiovascular Disease</td>
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<td>BDI</td>
<td>Beck Depression Inventory</td>
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<td>BMI</td>
<td>Body Mass Index</td>
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<tr>
<td>CAH</td>
<td>Critical Access Hospital</td>
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<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention</td>
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<td>CEHR</td>
<td>Certified Electronic Health Record</td>
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<td>CES-D</td>
<td>Center for Epidemiologic Studies</td>
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<td>CMS</td>
<td>Centers for Medicare and Medicaid Services</td>
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<tr>
<td>CPT</td>
<td>Current Procedural Terminology</td>
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<td>CVD</td>
<td>Cardiovascular Disease</td>
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<tr>
<td>CY</td>
<td>Calendar Year</td>
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<tr>
<td>DBT</td>
<td>Digital Breast Tomosynthesis</td>
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<tr>
<td>DTaP</td>
<td>Diphtheria, Tetanus and Acellular Pertussis</td>
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<td>DVT</td>
<td>Deep Vein Thrombosis</td>
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<td>ECG</td>
<td>Electrocardiogram</td>
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<tr>
<td>eCQI</td>
<td>Electronic Clinical Quality Improvement</td>
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<tr>
<td>Acronym</td>
<td>Term Meaning</td>
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<tr>
<td>eCQM</td>
<td>Electronic Clinical Quality Measures</td>
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<tr>
<td>ED</td>
<td>Emergency Department</td>
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<tr>
<td>EGA</td>
<td>Estimated Gestational Age</td>
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<td>EH</td>
<td>Eligible Hospital</td>
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<tr>
<td>EP</td>
<td>Eligible Professional</td>
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<td>ER</td>
<td>Emergency Room</td>
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<td>ESRD</td>
<td>End Stage Renal Disease</td>
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<td>FDA</td>
<td>Food and Drug Administration</td>
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<tr>
<td>FOBT</td>
<td>Fecal Occult Blood Test</td>
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<tr>
<td>GCS</td>
<td>Graduated Compression Stockings</td>
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<td>GDMT</td>
<td>Guideline-Directed Medical Therapy</td>
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<td>GLAD-PC</td>
<td>Guidelines for Adolescent Depression in Primary Care</td>
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<tr>
<td>HbA1c</td>
<td>Hemoglobin A1c</td>
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<td>HBIG</td>
<td>Hepatitis B Immune Globulin</td>
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<td>HF</td>
<td>Heart Failure</td>
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<td>HFrEF</td>
<td>Heart Failure with reduced Ejection Fraction</td>
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<tr>
<td>HHS</td>
<td>Health and Human Services</td>
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<tr>
<td>HIV</td>
<td>Human Immunodeficiency Virus</td>
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<td>HPV</td>
<td>Human Papillomavirus</td>
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<tr>
<td>ICD</td>
<td>International Classification of Diseases</td>
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<tr>
<td>ICSI</td>
<td>Institute for Clinical Systems Improvement</td>
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<tr>
<td>ICU</td>
<td>Intensive Care Unit</td>
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<td>IHS</td>
<td>Indian Health Service</td>
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<tr>
<td>IIV</td>
<td>Inactivated Influenza Vaccine</td>
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<tr>
<td>LDL-C</td>
<td>Low-Density Lipoprotein Cholesterol</td>
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<tr>
<td>LOINC</td>
<td>Logical Observation Identifiers Names and Codes</td>
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<tr>
<td>LVEF</td>
<td>Left Ventricular Ejection Fraction</td>
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<tr>
<td>MDD</td>
<td>Major Depressive Disorder</td>
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<tr>
<td>MDE</td>
<td>Major Depressive Episode</td>
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<tr>
<td>MMR</td>
<td>Measles, Mumps, and Rubella</td>
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<tr>
<td>NHANES</td>
<td>National Health and Nutrition Examination Survey</td>
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<tr>
<td>NHLBI</td>
<td>National Heart Lung and Blood Institute</td>
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<tr>
<td>NQF</td>
<td>National Quality Forum</td>
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<tr>
<td>NVAF</td>
<td>Nonvalvular Atrial Fibrillation</td>
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<tr>
<td>OB/GYN</td>
<td>Obstetrician/Gynecologist</td>
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<tr>
<td>PCI</td>
<td>Percutaneous Coronary Intervention</td>
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<tr>
<td>Acronym</td>
<td>Term Meaning</td>
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<tr>
<td>PCP</td>
<td>Primary Care Provider</td>
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<td>PE</td>
<td>Pulmonary Embolism</td>
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<td>POV</td>
<td>Purpose of Visit</td>
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<td>QRDA</td>
<td>Quality Reporting Data Architecture</td>
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<td>RPMS</td>
<td>Resource and Patient Management System</td>
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<td>RV</td>
<td>Rotavirus</td>
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<td>SCIP</td>
<td>Surgical Care Improvement Project</td>
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<tr>
<td>SNO-CT</td>
<td>Systematized Nomenclature of Medicine – Clinical Terms</td>
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<tr>
<td>SSRI</td>
<td>Selective Serotonin Reuptake Inhibitor</td>
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<tr>
<td>STEMI</td>
<td>ST-Segment Myocardial Infarction</td>
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<tr>
<td>TIA</td>
<td>Transient Ischemic Attack</td>
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<tr>
<td>UACR</td>
<td>Urinary Albumin-to-Creatinine Ratio</td>
</tr>
<tr>
<td>USHIK</td>
<td>United States Health Information Knowledgebase</td>
</tr>
<tr>
<td>USPSTF</td>
<td>U.S. Preventive Services Task Force</td>
</tr>
<tr>
<td>VSAC</td>
<td>Value Set Authority Center</td>
</tr>
<tr>
<td>VTE</td>
<td>Venous Thromboembolism</td>
</tr>
</tbody>
</table>
Contact Information
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If you have any questions or comments regarding this distribution, please contact the IHS IT Service Desk.

**Phone:** (888) 830-7280 (toll free)

**Web:** [https://www.ihs.gov/itsupport/](https://www.ihs.gov/itsupport/)

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