TO: NQF Members and Public

FR: NQF Staff

RE: Pre-voting review for Patient Safety - Complications Endorsement Maintenance: Phase I

DA: February 17, 2012

NQF has previously endorsed consensus standards to evaluate patient safety across conditions and care settings. This project seeks to identify and endorse measures for public reporting and quality improvement that specifically address complications of health care. An evaluation of previously NQF-endorsed complications measures and consideration of new measures will ensure the currency of NQF's portfolio of voluntary consensus standards. Given the volume and variety of measures being considered under this project, evaluation of the measures will be conducted in two phases. The attached report represents the results of Phase I.

A 26-member Steering Committee representing a range of stakeholder perspectives was appointed to evaluate 22 previously-endorsed measures for maintenance review in Phase I.

The draft document, *National Voluntary Consensus Standards: Patient Safety – Complications Endorsement Maintenance: Phase I* is posted on the NQF website along with the measure submission forms. This report recommends continued endorsement of 15 measures, with an additional two measures recommended for continued endorsement under 'reserve status.'

Pursuant to section II.A of the Consensus Development Process v. 1.8, this draft document, along with the accompanying material, is being provided to you at this time for purposes of review and comment only and is not intended to be used for voting purposes. You may post your comments and view the comments of others on the NQF website.

Please note that the organization of this report has been modified, similar to the recent Renal and Cardiovascular Endorsement Maintenance reports. The intention is to begin with high-level information (e.g., overarching evaluation issues and lists of measures) followed by more detail about the evaluation ratings and rationale in the measure evaluation summary tables. The detailed measure specifications for the recommended measures are in Appendix A and all submitted measure information is posted on the project web page.

All comments must be submitted no later than 6:00 pm ET, March 19, 2012.

Thank you for your interest in NQF's work. We look forward to your review and comments.

NATIONAL VOLUNTARY CONSENSUS STANDARDS: PATIENT SAFETY – COMPLICATIONS ENDORSEMENT MAINTENANCE: PHASE I

DRAFT TECHNICAL REPORT FOR COMMENT February 17, 2011

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PATIENT SAFETY MEASURES – COMPLICATIONS ENDORSEMENT MAINTENANCE: PHASE I Draft Technical Report

INTRODUCTION

Americans are exposed to more preventable medical errors than patients in other industrialized nations. It's estimated that preventable errors cost the United States \$17-\$29 billion per year in healthcare expenses, lost worker productivity, and disability. The costs are passed on in a number of ways—premiums, taxes, lost work time and wages, and health threats, to name a few. Proactively addressing medical errors and unsafe care will protect patients from harm and lead to more affordable, effective, and equitable care.

The Patient Safety Measures - Complications Endorsement Maintenance project will be executed in two phases, each addressing a number of specific complication-related domains. The first phase will focus on medication safety, venous thromboembolism, surgery, and care coordination, while the second phase will focus on falls, pressure ulcers, healthcare associated infections, and mortality. The Complications project builds on the work an earlier Patient Safety Measures project launched in 2009, which focused on healthcare-associated infections and radiation safety, among other issues. Endorsement maintenance provides the opportunity to harmonize specifications and to ensure that an endorsed measure represents the best in class. Composite and outcome measures and measures sensitive to the needs of vulnerable populations, including racial/ethnic minorities and Medicaid populations, were a priority.

MEASURE EVALUATION

On December 15-16, 2011, the Patient Safety - Complications Steering Committee evaluated one new measure and twenty four measures undergoing maintenance review against NQF's standard evaluation criteria. To facilitate the evaluation, the committee and candidate standards were divided into four workgroups for preliminary review of the measures against the evaluation sub-criteria prior to consideration by the entire Steering Committee. The Committee's discussion and ratings of the criteria are summarized in the evaluation tables beginning on page 5.

	MAINTENANCE	NEW	TOTAL
Measures under consideration	22	0	22
Withdrawn from consideration	5	0	5
Recommended	17	0	17
Not recommended	5	0	5
Reasons for Not	Importance – 4		
Recommending	Scientific Acceptability – 1		
	Overall – 0		
	Competing measure – 0		

PATIENT SAFETY - COMPLICATIONS

Overarching Issues

During the Steering Committee's discussion of the measures, several overarching issues emerged that were factored into the Committee's ratings and recommendations for multiple measures and are not repeated in detail with each individual measure:

Current Evidence and Relationship to Outcomes

The Committee expressed its preference for measures that provide clear and direct evidence of the measure's proximity to an improved outcome. Ensuring that the evidence provided to support the measure is current was highlighted, particularly for measures undergoing maintenance. In addition, several of the measures were focused on processes of care and discussions centered on whether what was measured truly was proximal to outcomes. This concern is reflected in the evaluation and underlying rationale for supporting a measure or not.

Impact on Quality

The Committee suggested measure developers provide detail on how their measure(s) impact quality. The Committee considered such information important when deciding whether a measure should be endorsed.

Measure with a Limited Performance Gap

The Committee suggested that the two Transfusion Reaction measures, which are similar but focus on different patient populations, have been performing at such high levels that continued efforts to publicly report on performance may not be warranted. They agreed that these measures should be maintained in the NQF portfolio with 'Reserve Status' designation, as they continue to address a critical aspect of patient safety and quality that should be sustained. The measures also fully meet all endorsement criteria with the exception of importance (as long as failure to meet this criterion was due to a high level of performance). The Committee acknowledged that placing these measures in 'reserve' could lead to the unintended consequence of inattention to the relevant processes or outcomes and consequently to potentially reduced levels of performance and poor patient outcomes. However, Committee members agreed that the 'Reserve Status' measures should be reviewed and reassessed in subsequent endorsement cycles to ensure that the performance remains at consistently high levels.

Continuum of Care

The Committee noted gaps in care that could be improved by addressing the patient's treatment across multiple settings of care. Committee members noted that aspects of a patient's condition should be reassessed when they are admitted to other departments within a healthcare facility and upon each interaction with a provider in an outpatient setting. Several of the measures under review were limited in their scope to a specific setting. This limitation was due to the focus of the developer and data source but when viewed from a patient-centered approach they should be broadened. The Committee suggested that when measures undergo the next maintenance cycle, to the extent possible, developers should focus on expanding the measure's scope.

Counts versus Rates

The Committee debated the usefulness of reporting rare but serious events, specifically related to several measures submitted by the Agency for Healthcare Research and Quality (AHRQ), in the form of counts or rates. While counts provide a more detailed breakdown of the data, rates may be more applicable for comparisons across settings and more useful to consumers. The Committee recognized that in these circumstances healthcare facilities may have no safety events captured by the measures, but stressed that continued monitoring of performance is necessary to improve quality and encourage transparency.

Discussion on Patient Safety Indicators (PSIs) and Pediatric Safety Indicators (PDIs)

Because most of the corresponding PSIs and PDIs are identical except for the population covered, the Committee often discussed both together. The rationales and information provided only vary when there was a separate concern given the patient population.

Discussion of Related and Competing Measures

The Steering Committee reviewed a number of previously-endorsed measures (0097, 0554, and 0646) that had been identified as related to and potentially competing with measure 0419. In general, the Committee saw the measures as related but not competing, and agreed that in the future they would like to see a single medication reconciliation measure that applies across populations, settings, and care transitions.

RECOMMENDATIONS FOR FUTURE MEASURE DEVELOPMENT

- 1. Wound care measures:
 - Vascular screening for patients with existing leg ulcers
 - Adequate venous compression for patients with existing venous leg ulcers
 - Adequate offloading patients with diabetic foot ulcers
 - Adequate support surface for patients with stage III-IV pressure ulcers
- 2. Obstetric measures:
 - Induction and augmentation of labor
 - Outcomes of neonatal birth injury
- 3. Infection measures:
 - Clostridium difficile colitis is epidemic in US and should be measured.
 - Vascular catheter infections in other settings including--dialysis catheters, home infusion, peripherally inserted central catheter lines; nursing home catheters
- 4. Equipment related injury:
 - Monitoring of product related events
- 5. Information technology:
 - EHR programming related errors
- 6. The expectation for physical mobility among hospitalized adults:
 - The severity of the inactivity among people who are hospitalized was described in this article: Brown CJ, Redden DT, Flood KL, Allman RM. The under recognized epidemic of low mobility during hospitalization of older adults. *J Am Geriatr* Soc. 2009;57(9): 1660-1665 (see http://onlinelibrary.wiley.com/doi/10.1111/j.1532-5415.2009.02393.x/full). There are extensive adverse effects associated with prolonged bed rest and much of these adverse effects are preventable with daily activity.

MEASURES RECOMMENDED

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0419 Documentation of Current Medications in the Medical Record	7
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0372 Intensive Care Unit Venous Thromboembolism Prophylaxis	
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0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)	14
0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant	17
0344 Accidental Puncture or Laceration Rate (PDI 1)	18
0345 Accidental Puncture or Laceration Rate (PSI 15)	20
0362 Foreign Body left after procedure (PDI 3)	22
0363 Foreign Body Left During Procedure (PSI 5)	24
0263 Patient Burn	25
0346 Iatrogenic Pneumothorax Rate (PSI 6)	25
0348 Iatrogenic Pneumothorax Rate (PDI 5)	27

MEASURES RECOMMENDED FOR RESERVE STATUS

0349	Transfusion Reaction	(PSI 16)	3
0350	Transfusion Reaction	(PDI 13))
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MEASURES NOT RECOMMENDED

0374 Venous Thromboembolism Patients Recieving Unfractionated Heparin with Dosages/Platelet	
Count Monitoring by Protocol or Nomogram	.30
0375 Venous Thromoboembolism Warfarin Therapy Discharge Instructions	.32
0501 Confirmation of Endotracheal Tube Placement	.33
0523 Pain Assessment Conducted	.34
0524 Pain Interventions Implemented During Short Term Episodes Of Care	35

0022 Use of High Risk Medications in the Elderly

Measure Submission Form Description: a: Percentage of Medicare members 65 years of age and older who received at least one high-risk medication. b: Percentage of Medicare members 65 years of age and older who received at least two different high-risk medications. For both rates, a lower rate represents better performance. Numerator Statement: a: At least one prescription dispensed for any high-risk medication during the measurement year. b: At least two prescriptions dispensed for different high-risk medications during the measurement year. Denominator Statement: All patients ages 65 years and older as of December 31 of the measurement year. Exclusions: N/A Adjustment/Stratification: No risk adjustment or risk stratification N/A N/A Level of Analysis: Health Plan Type of Measure: Process Data Source: Administrative claims, Electronic Clinical Data, Electronic Clinical Data : Pharmacy Healthcare Effectiveness Data and Information Set (HEDIS) Measure Steward: National Committee for Quality Assurance STEERING COMMITTEE MEETING 12/15-16/2011 1. Importance to Measure and Report: Y-19; N-3 (1a. High Impact: 1b. Performance Gap, 1c. Evidence) 1a. Impact: H-6; M-1; L1-; I-0; 1b. Performance Gap: H-6; M-1; L-0; I-1 1c. Evidence Quantity: H-5; M-1; L-1; I-1; Quality: H-4; M-2; L-1; I-1; Consistency: H-5; M-1; L-1; I-1 Rationale: The measure focuses on medications that are known to cause harm or lead to adverse events in the elderly. The literature cited, including the 2003 Beers criteria, provides further evidence for the measure's focus. The committee and developer acknowledged

that the American Geriatrics Society is currently reviewing and updating the list of medications and the measure will be updated to reflect those changes when they are released.

2. Scientific Acceptability of Measure Properties: Y-22; N-0

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NATIONAL QUALITY FORUM
0022 Use of High Risk Medications in the Elderly
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity) 2a. Reliability: H-5; M-3; L-0; I-0; 2b. Validity: H-4; M-4; L0-; I-0
Rationale: The measure is well specified and the denominator is clear.
 Usability: H-9; M-12; L-1; I-0 (<i>Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvemen</i> Public Reporting: H-4; M-4; L-0; I-0 Ql: H-4; M-4; L-0; I-0
Rationale: The measure will be useful for patient safety and provide valuable information to consumers.
 4. Feasibility: H-8; M-13; L-1; I-0 (4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented) 4a. Byproduct of Care Processes: H-5; M-1; L-1; I-0 4b. Electronic data sources: H-5; M-1; L-1; I-0
 4c. Suscep inaccuracies, consequences: H-5; M-1; L-1; I-0 4d. Data collection strategy: H-5; M-1; L-1; I-0
<u>Rationale</u> : The measure may need to be updated in the future to accommodate any changes in medication monitoring or remove any medications that are no longer available. The developer indicated that it would be reviewed frequently.
Steering Committee Recommendation for Endorsement: Y-20; N-2
Rationale: The measure would inform patient safety efforts and the consumer. It represents a major patient safety initiative.
0419 Documentation of Current Medications in the Medical Record
Measure Submission Form
Description: Percentage of patients aged 18 years and older with a list of current medications (includes prescription, over-the-counter, herbals, vitamin/mineral/dietary [nutritional] supplements) documented by the provider, including drug name, dosage, frequency and route
Numerator Statement: Current medications including name, dosage, frequency and route documented by the provider Denominator Statement: All patients aged 18 years and older on date of patient encounter
Exclusions: Not Eligible – A patient is not eligible if one or more of the following condition(s) exist: Patient refuses to participate
Patient is in an urgent or emergent medical situation where time is of the essence and to delay treatment would jeopardize the patient's health status
Patient cognitively impaired and no authorized representative available
Adjustment/Stratification: No risk adjustment or risk stratification. N/A No stratification. All eligible patients are subject to the same numerator criteria.
Level of Analysis: Clinician : Individual, Population : National
Type of Measure: Process
Data Source: Administrative claims, Electronic Clinical Data : Registry Medicare Part B claims data
Measure Steward: Centers for Medicare & Medicaid STEERING COMMITTEE MEETING 12/15-16/2011
1. Importance to Measure and Report: Y-19; N-2
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-6; M-0; L2-; I-0; 1b. Performance Gap: H-6; M-0; L-2; I-0
1c. Evidence Quantity: H-1; M-3; L-3; I-1; Quality: H-1; M-3; L-3; I-1; Consistency: H-1; M-4; L-2; I-1
<u>Rationale</u> : The Committee affirmed the importance of the measure's goals: to prompt discussions between physicians and patients, to increase knowledge of patients' medical histories, and to reduce adverse drug events. The Committee also discussed the importance of medication reconciliation in general. Since reporting on this measure is voluntary, the Committee noted that it is not possible to clearly define the performance gap but current rates demonstrate a gap for just documentation of current medications in the medical record.
 Scientific Acceptability of Measure Properties: Y-15; N-5 (2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-0; M-4; L-4; I-0; 2b. Validity: H-0; M-5; L-3; I-0
NQF REVIEW DRAFT—DO NOT CITE OR QUOTE
Comments due by March 19, 2012 by 6:00pm ET

0419 Documentation of Current Medications in the Medical Record

<u>Rationale</u>: The Committee had several concerns related to whether the specifications were precise and understandable and whether the results would be valid. The Committee was concerned that it would be difficult to effectively document a patient's vitamin and over-thecounter medication use. The Committee requested that the developer clarify language in the measure to focus on whether a medical history was taken and a patient's medications were documented rather than the creation of a current and complete medication list. Committee members suggested that the measure should be rewritten to more clearly reflect that providers are being measured on whether patients were asked about their medications on each visit. Concerns regarding the validity of the data were discussed. The measure currently asks the provider to report on whether they have current medications documented in the medical record but it is not known whether what is documented actually is what the patient is taking and if any were missed.

3. Usability: H-7; M-7; L-5; I-1

(*Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement*) **3a.** Public Reporting: H-1; M-4; L-3; I-0

3b. QI: H-1; M-4; L-2; I-0

<u>Rationale</u>: Recognizing that the measure is currently being used in both public reporting and quality improvement programs, the Steering Committee agreed that the measure meets the usability criterion.

4. Feasibility: H-2; M-11; L-6; I-1

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-3; M-3; L-2; I-0

4b. Electronic data sources: H-1; M-3; L-4; I-0

4c. Suscep inaccuracies, consequences: H-0; M-5; L-2; I-1

4d. Data collection strategy: H-1; M-4; L-2; I-0

Rationale: The measure is currently being collected and no concerns with feasibility were raised.

Steering Committee Recommendation for Endorsement: Y-14; N-6

Rationale: The Steering Committee agreed that documentation of patients' current medications is an area where there is a great need and opportunity for improvement. Many Committee members stated that they would prefer an outcome measure in this area but acknowledged that no such measure existed, and agreed that in the absence of an outcome measure that correlates with reconciliation, this measure was a good starting point. The Steering Committee also reviewed a number of medication reconciliation measures (0097, 0554, and 0646) that had been identified as related and potentially competing with measure 0419. In general, the Committee saw the measures as related but not competing, and agreed that in the future they would like to see a single medication reconciliation measure that applies across populations, settings, and care transitions.

0371 Venous Thromboembolism Prophylaxis

Measure Submission Form

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. This measure is part of a set of six nationally implemented prevention and treatment measures that address VTE (VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: Incidence of Potentially-Preventable VTE) that are used in The Joint Commission's accreditation process.

Numerator Statement: Patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given: •the day of or the day after hospital admission

•the day of or the day after surgery end date for surgeries that start the day of or the day after hospital admission **Denominator Statement:** All discharged hospital inpatients

Exclusions:

•Patients less than 18 years of age

•Patients who have a length of stay (LOS) less than two days and greater than 120 days

•Patients with Comfort Measures Only documented on day of or day after hospital arrival

Patients enrolled in clinical trials

•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 LOS greater than or equal to one day

•Patients with ICD-9-CM Principal Diagnosis Code of Mental Disorders or Stroke as defined in Appendix A, Table 7.01, 8.1 or 8.2 •Patients with ICD-9-CM Principal or Other Diagnosis Codes of Obstetrics or VTE as defined in Appendix A, Table 7.02, 7.03 or 7.04 •Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected surgeries as defined in

0371 Venous Thromboembolism Prophylaxis Appendix A, Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, 5.24 Adjustment/Stratification: No risk adjustment or risk stratification; Not applicable; Not Applicable, the measure is not stratified. Level of Analysis: Facility, Population : National Type of Measure: Process Data Source: Administrative claims, Electronic Clinical Data : Electronic Health Record, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. The vendor may not offer the measure set to hospitals until verification has been passed. Measure Steward: The Joint Commission STEERING COMMITTEE MEETING 12/15-16/2011 1. Importance to Measure and Report: Y-20; N-1 (1a. High Impact: 1b. Performance Gap, 1c. Evidence) 1a. Impact: H-8; M-1; L-0; I-0; 1b. Performance Gap: H-8; M-1; L-0; I-0 1c. Evidence Quantity: H-8; M-1; L-0; I-0; Quality: H-8; M-1; L-0; I-0; Consistency: H-8; M-1; L-0; I-0 Rationale: There is strong evidence to support the measure and recent reporting indicated a performance gap of 17%. However, the Committee stated that the measure has a limited ability to impact outcomes since it lacks a validated risk assessment model and remains vague. Yet, the Committee agreed that it would encourage hospitals to have standardized policies for VTE prophylaxis among inpatients. 2. Scientific Acceptability of Measure Properties: Y-17: N-4 (2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity) 2a. Reliability: H-5; M-4; L-0; I-0; 2b. Validity: H-7; M-2; L-0; I-0 Rationale: The measure was considered a good starting point for examining whether patients were given prophylaxis. However, the Committee expressed concern about the measure's lack of specificity on how risk is assessed. Patients at low risk as well as certain other populations, such as renal patients and older patients, should not be given prophylaxis since it increases the risk of bleeding. The American College of Chest Physicians is expected to release new guidelines (ACCP-9) around VTE prophylaxis in February 2012; these guidelines may offer additional direction to providers. Additionally, the Committee expressed reservations regarding the lack of a definition for "effective prophylaxis". Hospitals may vary in their interpretation of this language; therefore, the measure may not be limited in its usefulness for comparison. It was suggested that in the future the measure could be further specified and improved. While there were concerns with the specificity of the measure, the measure as specified demonstrated reliable results and face validity was provided. 3. Usability: H-3; M-14; L-4; I-0 (Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement) 3a. Public Reporting: H-8; M-1; L-0; I-0 3b. QI: H-8; M-1; L-0; I-0 Rationale: This measure is part of a VTE measure set that will be implemented nationally in January 2013. While the Committee guestioned whether the measure alone will provide useful information to consumers, members agreed that measuring VTE prophylaxis will lead to quality improvement. 4. Feasibility: H-8; M-10; L-3; I-0 (4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented) 4a. Byproduct of Care Processes: H-8; M-1; L-0; I-0 4b. Electronic data sources: H-3; M-2; L-3; I-0 4c. Suscep inaccuracies, consequences: H-5; M-4; L-0; I-0 4d. Data collection strategy: H-7; M-1; L-0; I-1 Rationale: Creating a risk assessment model would have made data collection more complicated, which would further limit feasibility. The measure will be reevaluated and updated every six months by the developer. Steering Committee Recommendation for Endorsement: Y-17; N-4 Rationale: The measure recognizes that VTE prophylaxis is an important part of the process of care for a variety of diagnoses and

0371 Venous Thromboembolism Prophylaxis

treatment plans. While there are concerns about the implementation and usefulness of the measure, the Committee agreed that it continues to be a good starting point in the assessment of hospital performance related to VTE prophylaxis.

0372 Intensive Care Unit Venous Thromboembolism Prophylaxis

Measure Submission Form

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 the initial admission (or transfer) to the Intensive Care Unit (ICU) or surgery end date for surgeries that start the day of or the day after ICU admission (or transfer). This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring by Protocol, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: VTE Incidence of Potentially-Preventable VTE).

Numerator Statement: Patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given: •the day of or the day after ICU admission (or transfer)

•the day of or the day after surgery end date for surgeries that start the day of or the day after ICU admission (or transfer) **Denominator Statement:** Patients directly admitted or transferred to ICU

Exclusions:

•Patients less than 18 years of age

•Patients who have a hospital length of stay (LOS) less than two days and greater than 120 days

•Patients with Comfort Measures Only documented on day of or day after hospital arrival

•Patients enrolled in clinical trials

•Patients with ICU LOS less than one day without VTE prophylaxis administered and documentation for no VTE prophylaxis •Patients with ICD-9-CM Principal or Other Diagnosis Code of Obstetrics or VTE as defined in Appendix A, Table 7.02, 7.03, or 7.04 •Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected surgeries as defined in Appendix A, Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, 5.24 that start the day of or the day after ICU admission or transfer **Adjustment/Stratification:** No risk adjustment or risk stratification. Not applicable Not Applicable, the measure is not stratified.

Level of Analysis: Facility, Population : National Type of Measure: Process

Data Source: Administrative claims, Electronic Clinical Data, Paper Records Each data element in the data dictionary includes suggested data sources.

The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. The vendor may not offer the measure set to hospitals until verification has been passed. **Measure Steward:** The Joint Commission

STEERING COMMITTEE MEETING 12/15-16/2011

1. Importance to Measure and Report: Y-21; N-0

(1a. High Impact: 1b. Performance Gap, 1c. Evidence)

1a. Impact: H-8; M-1; L-0; I-0; 1b. Performance Gap: H-7; M-2; L-0; I-0

1c. Evidence Quantity: H-7; M-2; L-0; I-0; Quality: H-7; M-2; L-0; I-0; Consistency: H-7; M-2; L0-; I-0

<u>Rationale</u>: There is strong evidence for the measure given the population – patients in intensive care units (ICU) and the measure noted an aggregate performance rate of 87.9 %, indicating a potential performance gap of 12.1 %...

2. Scientific Acceptability of Measure Properties: Y-21; N-0

(2a. Reliability - precise specifications, testing; 2b. Validity - testing, threats to validity)

2a. Reliability: H-6; M-3; L-0; I-0; 2b. Validity: H-7; M-2; L-0; I-0

<u>Rationale</u>: The specifications for this measure are similar to Measure #0371 but as was discussed with that measure, while there are limitations to the measure the measure addresses an important population and aspect of care. The populations for this measure and Measure #0371 differ since this measure looks at patients who are admitted to the ICU at any point during the hospitalization, ensuring that patients are assessed when they are at highest risk regardless of whether they were initially assessed at the time of admission (the focus for Measure #0371). In the future, the measure could be improved by also including patients who are transferred out of the ICU since that point in time is not currently captured in the measures under consideration. The measure as specified demonstrated reliable results and face validity was provided.

3. Usability: H-10; M-11; L-0; I-0

(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)

0372 Intensive Care Unit Venous Thromboembolism Prophylaxis

3a. Public Reporting: H-8; M-1; L-0; I-0 3b. QI: H-8; M-1; L-0; I-0

<u>Rationale</u>: This measure is part of a VTE measure set that will be implemented nationally in January 2013. While the Committee questioned whether the measure alone will provide useful information to consumers, members agreed that measuring VTE prophylaxis will lead to quality improvement.

4. Feasibility: H-12; M-8; L-1; I-0

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-8; M-1; L-0; I-0

4b. Electronic data sources: H-2; M-6; L-1; I-0

4c. Suscep inaccuracies, consequences: H-5; M-4; L-0; I-0

4d. Data collection strategy: H-7; M-1; L-1; I-0

<u>Rationale</u>: Creating a risk assessment model would have made data collection more complicated, which would further limit feasibility. The measure will be reevaluated and updated every six months by the developer.

Steering Committee Recommendation for Endorsement: Y-21; N-0

Rationale: The measure recognizes that VTE prophylaxis is an important part of the process of care for a variety of diagnoses and treatment plans. While there are concerns about the implementation and usefulness of the measure, the Committee agreed that this measure addresses a very high risk population and room for improvement exists.

0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy

Measure Submission Form

Description: This measure assesses the number of patients diagnosed with confirmed VTE who received an overlap of Parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulation and warfarin therapy. For patients who received less than five days of overlap therapy, they should be discharged on both medications and have a Reason for Discontinuation of Overlap Therapy. Overlap therapy should be administered for at least five days with an international normalized ratio (INR) greater than or equal to 2 prior to discontinuation of the parenteral anticoagulation therapy, or INR less than 2 but discharged on both medications or have a Reason for Discontinuation of Overlap Therapy. This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: Incidence of Potentially-Preventable VTE). Numerator Statement: Patients who received overlap therapy: Included Populations: Patients who received warfarin and parenteral anticoagulation: • Five or more days, with an INR greater than or equal to 2 prior to discontinuation of parenteral therapy OR •Five or more days, with an INR less than 2 and discharged on overlap therapy OR ·Less than five days and discharged on overlap therapy OR •With documentation of reason for discontinuation of overlap therapy OR •With documentation of a reason for no overlap therapy Denominator Statement: Patients with confirmed VTE who received warfarin. The target population includes patients discharged with an ICD-9-CM Principal or Other Diagnosis Codes for VTE as defined in Table 7.03 or Table 7.04. Exclusions: •Patients less than 18 years of age •Patients who have a length of stay greater than 120 days Patients with Comfort Measures Only documented Patients enrolled in clinical trials Patients discharged to a health care facility for hospice care •Patients discharged to home for hospice care ·Patients who expired ·Patients who left against medical advice •Patients discharged to another hospital Patients without warfarin therapy during hospitalization •Patients without VTE confirmed by diagnostic testing Adjustment/Stratification: No risk adjustment or risk stratification; Not Applicable; Not Applicable, the measure is not stratified.

0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy

Level of Analysis: Facility, Population : National

Type of Measure: Process

Data Source: Administrative claims, Electronic Clinical Data, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. Verification must be completed and passed before the vendor can offer the data collection tool to hospitals.

Measure Steward: The Joint Commission

STEERING COMMITTEE MEETING 12/15-16/2011

1. Importance to Measure and Report: Y-20; N-1

(1a. High Impact: 1b. Performance Gap, 1c. Evidence)

1a. Impact: H-9; M-0; L-0; I-0; 1b. Performance Gap: H-9; M-0; L-0; I-0

1c. Evidence Quantity: H-4; M-4; L-1; I-0; Quality: H-5; M-4; L-0; I-0; Consistency: H-7; M-2; L-0; I-0

<u>Rationale</u>: The measure is based on multiple guidelines, primarily from the American College of Chest Physicians (ACCP), that indicate overlap therapy of heparin and warfarin should be used to reduce a patient's risk of increased hypercoagulability. The body of evidence supports the measure's focus and a clear performance gap remains.

2. Scientific Acceptability of Measure Properties: Y-18; N-3

(2a. Reliability - precise specifications, testing; 2b. Validity - testing, threats to validity)

2a. Reliability: H-7; M-2; L-0; I-0; 2b. Validity: H-7; M-1; L-1; I-0

<u>Rationale</u>: The Committee was concerned that the timeframes specified in the measure were complex but the testing demonstrated that the measure as specified was reliable and valid. The Committee agreed with the exclusion allowing a clinician to document an explicit reasoning for not discharging with overlap therapy. It was suggested that in the future the settings be expanded to include patients in the emergency room, since a number of patients are not admitted to the hospital.

3. Usability: H-7; M-9; L-5; I-0

(*Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement*) **3a.** Public Reporting: **H-8**; **M-1**; **L-0**; **I-0**

3b. QI: H-7; M-2; L-0; I-0

Rationale: The measure is part of a VTE measure set that will be nationally implemented in January 2013.

4. Feasibility: H-6; M-9; L-6; I-0

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-8; M-1; L-0; I-0

4b. Electronic data sources: H-3; M-4; L-2; I-0

4c. Suscep inaccuracies, consequences: H-5; M-4; L0-; I-0

4d. Data collection strategy: H-6; M-2; L-1; I-0

<u>Rationale</u>: Some members of the Committee expressed concern about how timeframes were defined within the measure and whether the data was feasible to collect. The developer explained that the measure scope and timeframe were specified to ensure that the data would be feasible to capture.

Steering Committee Recommendation for Endorsement: Y-18; N-3

Rationale: The measure will address a lack of knowledge regarding the importance of overlap therapy.

0376 Incidence of Potentially Preventable Venous Thromboembolism

Measure Submission Form

Description: This measure assesses the number of patients with confirmed venous thromboembolism (VTE) during hospitalization (not present at admission) who did not receive VTE prophylaxis between hospital admission and the day before the VTE diagnostic testing order date. This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring by Protocol, and VTE-5: VTE Warfarin Therapy Discharge Instructions). **Numerator Statement:** Patients who received no VTE prophylaxis prior to the VTE diagnostic

NATIONAL QUALITY FORUM	
0376 Incidence of Potentially Preventable Venous Thromboembolism	
test order date	
Denominator Statement: Patients who developed confirmed VTE during hospitalization. The target population includes patients	
discharged with an ICD-9-CM Secondary Diagnosis Codes for VTE as defined in Table 7.03 or Table 7.04.	
Exclusions: Patients less than 18 years of age	
Patients who have a length of stay greater than 120 days	
Patients with Comfort Measures Only documented	
Patients enrolled in clinical trials	
 Patients with ICD-9-CM Principal Diagnosis Code of VTE as defined in Appendix A, Table 7.03 or 7.04 	
Patients with VTE Present at Admission	
 Patients with reasons for not administering mechanical and pharmacologic prophylaxis 	
Patients without VTE confirmed by diagnostic testing	
Adjustment/Stratification: No risk adjustment or risk stratification; No risk adjustment or risk stratification as intermediate outcome	me;
Not Applicable	
Level of Analysis: Facility, Population : National	
Type of Measure: Outcome	
Data Source: Administrative claims, Paper Records Each data element in the data dictionary includes suggested data sources.	
The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based or	n the
measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the	
collection tool with the measure specifications. Verification must be completed and passed before the vendor can offer the data	
collection tool to hospitals.	
Measure Steward: The Joint Commission	
STEERING COMMITTEE MEETING 12/15-16/2011	
1. Importance to Measure and Report: Y-20; N-2	
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)	
1a . Impact: H-8; M-0; L-0; I-1; 1b. Performance Gap: H-6; M-2; L-0; I-1	
1c. Evidence Quantity: H-6; M-2; L-0; I-0; Quality: H-6; M-1; L-1; I-0; Consistency: H-6; M-1; L-1; I-0	
Rationale: The measure is important because it indicates the adequacy of the hospital's risk assessment profile by reporting the ra	ate at
which patients acquired VTE and did not receive prophylaxis. The measure presented an aggregate performance gap of 13.2% a	
stated that the gap would ideally be reduced to 0%. However, the Committee expressed concern that the measure did not gauge t	
adequacy of the prophylaxis. They also recognized that patients receiving adequate prophylaxis could still develop adverse event	
regardless of the quality of the provider's care.	0
2. Scientific Acceptability of Measure Properties: Y-20; N-1	
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)	
2a. Reliability: H-7; H-1; L-1; I-0; 2b. Validity: H-6; H-3; L-0; I-0	
Rationale: The measure as specified includes the rate of patients who had a confirmed VTE that was not present on admission – t	he
Committee was interested with the idea that while the measure focused on those patients who had a treatment failure (i.e., were n	
assessed and treated resulting in a VTE), the denominator itself also provided valuable information. Reliability and validity were	01
demonstrated.	
3. Usability: H-7; M-14; L-1; I-0	
(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improve	omont)
3a. Public Reporting: H-7; M-2; L-0; I-0	ennenny
3b. QI: H-7; M-2; L-0; I-0	
ЗВ. ЦІ. П-7, №-2, L-0, I-0	
Rationale: The measure will assist hospitals with quality improvement by reporting patients not risk-assessed for VTE.	
4. Feasibility: H-7; M-13; L-1; I-0	
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequen	Ces
identified 4d. Data collection strategy can be implemented)	
4a. Byproduct of Care Processes: H-8; M-1; L-0; I-0	
4b. Electronic data sources: H-4; M-5; L-0; I-0	
4c. Suscep inaccuracies, consequences: H-5; M-3; L-1; I-0	
4d. Data collection strategy: H-6; M-2; L-1; I-0	
Detionals. The measure is surroutly being collected and an energy with face it if the surroutly d	
Rationale: The measure is currently being collected and no concerns with feasibility were raised.	
NQF REVIEW DRAFT—DO NOT CITE OR QUOTE	
Comments due by March 19, 2012 by 6:00pm ET	

0376 Incidence of Potentially Preventable Venous Thromboembolism

Steering Committee Recommendation for Endorsement: Y-20; N-2

Rationale: The measure indicates whether facilities are adequately assessing patients for their risk of developing VTE and provide important information on the incidence of VTEs that could have potentially been avoided through appropriate assessment and treatment.

0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)

Measure Submission Form

Description: Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM codes for deep vein thrombosis or pulmonary embolism in any secondary diagnosis field.

Numerator Statement: Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM codes for deep vein thrombosis or pulmonary embolism in any secondary diagnosis field.

Denominator Statement: All surgical discharges age 18 and older defined by specific DRGs or MS-DRGs and an ICD-9-CM code for an operating room procedure

Exclusions: Exclude cases:

- with principal diagnosis of deep vein thrombosis or pulmonary embolism or secondary diagnosis present on admission

- where a procedure for interruption of vena cava is the only operating room procedure

- where a procedure for interruption of vena cava occurs before or on the same day as the first operating room procedure

- MDC 14 (pregnancy, childbirth, and puerperium)

- with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)

Adjustment/Stratification: Statistical risk model. The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate.

Tale, muit	ipileu by li
Age	18 to 24
Age	25 to 29
Age	30 to 34
Age	35 to 39
Age	40 to 44
Age	45 to 49
Age	50 to 59
Age	65 to 74
Age	75 to 79
Age	80 to 84
Age	85+
MDRG	101
MDRG	102
MDRG	103
MDRG	104
MDRG	105
MDRG	107
MDRG	108
MDRG	401
MDRG	402
MDRG	502
MDRG	503
MDRG	505
MDRG	507
MDRG	508
MDRG	509
MDRG	511

	stoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)
MDRG	514
MDRG	519
MDRG	601
MDRG	602
MDRG	603
MDRG	604
MDRG	611
MDRG	701
MDRG	705
MDRG	801
MDRG	802
MDRG	804
MDRG	805
MDRG	806
MDRG	807
MDRG	808
MDRG	811
MDRG	815
MDRG	1001
MDRG	1003
MDRG	1006
MDRG	1101
MDRG	1102
MDRG	1103
MDRG	1104
MDRG	1107
MDRG	1109
MDRG	1201
MDRG	1301
MDRG	1302
MDRG	1303
MDRG	1304
MDRG	1707
MDRG	1708
MDRG	1709
MDRG	1801
MDRG	1802
MDRG	2104
MDRG MDRG	2406 2407
MDRG	2407
MDRG	2501
MDRG	7701
MDRG	7702
MDRG	1
MDC	4
MDC	5
MDC	7
MDC	, 11
MDC	12
MDC	16
MDC	17
MDC	18
MDC	21
NDC	21

0/E0 Do	stonarati	ue Dulmonary Embalism or Doop Voin Thrombosis Data (DSI 12)
		ve Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)
MDC MDC	22 24	
MDC	24 25	
TRNSFE		Transfer in
COMOR		Transfer-in CHF
COMOR COMOR		VALVE PULMCIRC
COMOR		PERIVASC
COMOR		
COMOR		HTN_C PARA
COMOR		NEURO
COMOR		CHRNLUNG
COMOR		DM
COMOR		НҮРОТНҮ
COMOR		RENLFAIL
COMOR		AIDS
COMOR		AIDS LYMPH
COMOR		METS
COMOR		TUMOR
COMOR		OBESE
COMOR		WGHTLOSS
COMOR		BLDLOSS
COMOR		ANEMDEF
COMOR		ALCOHOL
COMOR		DRUG
COMOR		PSYCH
COMOR		DEPRESS Not applicable
	f Analysis	
	Measure:	
		ninistrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
		rch and Quality, Rockville, MD.
		: Agency for Healthcare Research and Quality
		AITTEE MEETING 12/15-16/2011
-		Measure and Report: Y-20; N-2
		1b. Performance Gap, 1c. Evidence)
		I-1; L-0; I-0; 1b. Performance Gap: H-6; M-3; L-0; I-0
		ntity: H-3; M-2; L-1; I-1; Quality: H-3; M-2; L-1; I-1; Consistency: H-3; M-2; L-1; I-1
		asure was considered important due to its high impact and opportunity for improvement.
		ptability of Measure Properties: Y-20; N-1
		recise specifications, testing; 2b. Validity – testing, threats to validity)
		; M-2; L-1; I-0; 2b. Validity: H-4; M-2; L-3; I-0
		mmittee discussed the measure in light of new studies provided by AHRQ representatives further indicating the
		ility of the measure. The evidence demonstrated that changes in the ICD-9 codes and present-on-admission
		the number of false positives captured by the measure. The Committee accepted that the data provided from the
		decrease in false positives that would be indicative of a larger body of evidence.
		M-12; L-1; I-0
		rstandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
		ng: H-6; M-2; L-1; I-0
	H-6; M-2; I	5
Rational		
		3; M-7; L-1; I-0
		generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences
		collection strategy can be implemented)
		care Processes: H-9; M-0; L-0; I-0
		NOF REVIEW DRAFT-DO NOT CITE OR OLIOTE

0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12) 4b. Electronic data sources: H-8; M-1; L-0; I-0 4c. Suscep inaccuracies, consequences: H-7; M-1; L-1; I-0 4d. Data collection strategy: H-8; M-1; L-0; I-0 Rationale: Steering Committee Recommendation for Endorsement: Y-20; N-1 Rationale: The measure indicated an opportunity for improvement and was proven to be scientifically acceptable through new studies, which demonstrated a reduction in the number of false positives captured by the measure. 0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant Measure Submission Form Description: Percentage of ASC admissions experiencing a wrong site, wrong side, wrong patient, wrong procedure, or wrong implant event. Numerator Statement: ASC admissions experiencing a wrong site, wrong side, wrong patient, wrong procedure, or wrong implant Denominator Statement: All ASC admissions Exclusions: None Adjustment/Stratification: No risk adjustment or risk stratification Not applicable The measure is not stratified Level of Analysis: Facility Type of Measure: Outcome Data Source: Paper Records ASC medical records, as well as incident/occurrence reports, and variance reports may serve as data sources. No specific collection instrument is required although the ASC Quality Collaboration has developed a sample data collection instrument that may be used as desired. Facilities may use any collection instrument that allows tracking of all wrong site, wrong side, wrong patient, wrong procedure, and wrong implant events. Measure Steward: Ambulatory Surgical Center Quality Collaboration STEERING COMMITTEE MEETING 12/15-16/2011 1. Importance to Measure and Report: Y-22; N-0 (1a. High Impact: 1b. Performance Gap, 1c. Evidence) 1a. Impact: H-6; M-1; L-0; I-0; 1b. Performance Gap: H-5; M-2; L-0; I-0 1c. Evidence Quantity: H-4; M-2; L-1; I-0; Quality: H-3; M-3; L-1; I-0; Consistency: H-3; M-4; L-0; I-0 Rationale: The measure provides a way to collect information on a serious reportable event and will improve ambulatory surgical care. The rate for surgeries involving the wrong site, side, patient, procedure or implant ranged from a minimum of 0.00% to a maximum of 0.31%. 2. Scientific Acceptability of Measure Properties: Y-21; N-1 (2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity) 2a. Reliability: H-5; M-1; L-1; I-0; 2b. Validity: H-5; M-2; L-0; I-0 Rationale: The measure is reported in the ambulatory care setting, increasing the monitoring of wrong site, wrong side procedures beyond the in-patient setting. The Committee suggested that in the future the measure be stratified by procedure and reported as a count to keep it consistent with hospitals' current monitoring practices. Reliability and validity of the measure as specified was demonstrated. 3. Usability: H-15; M-6; L-1; I-0 (Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement) 3a. Public Reporting: H-6; M-1; L-0; I-0 3b. QI: H-6; M-1; L-0; I-0 Rationale: The measure is currently being collected on a voluntary basis and will be included in CMS' mandatory reporting program beginning October 1, 2012. It is reported on a publicly available website, and in the future the developer will be able to report statistics based on demographics, procedure and state. 4. Feasibility: H-12; M-9; L-0; I-0 (4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented) 4a. Byproduct of Care Processes: H-5; M-1; L-1; I-0 4b. Electronic data sources: H-2; M-3; L-2; I-0 NQF REVIEW DRAFT-DO NOT CITE OR QUOTE Comments due by March 19, 2012 by 6:00pm ET

NATIONAL QUALITY FORUM
0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
4c. Suscep inaccuracies, consequences: H-4; M-3; L-0; I-0 4d. Data collection strategy: H-6; M-1; L-0; I-0
Rationale: The measure is effectively collected from manual reviews of paper records. Steering Committee Recommendation for Endorsement: Y-21; N-1
Rationale : The measure is used to track wrong site, wrong side surgeries in the ambulatory surgery setting b for mandatory reporting on a serious reportable event. The gap in care demonstrates an opportunity for improvement with a maximum rate for surgeries involving the wrong site, side, patient, procedure or implant of 0.31%.
0244 Assidental Duncture or Lassration Data (DDL1)
0344 Accidental Puncture or Laceration Rate (PDI 1)
Measure Submission Form Description: Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field. Numerator Statement: Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field. Denominator Statement: All surgical and medical discharges under age 18 defined by specific DRGs or MS-DRGs. Exclusions: Exclude cases:
 with principal diagnosis denoting accidental cut, puncture, perforation, or laceration or secondary diagnosis present on admission normal newborn
 neonate with birth weight less than 500 grams (Birth Weight Category 1) MDC 14 (pregnancy, childbirth, and puerperium) with ICD-9-CM code for spine surgery
- with missing discharge gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing) See Pediatric Quality Indicators Appendices:
 Appendix I – Definitions of, Neonate, Newborn, Normal Newborn, and Outborn Appendix L – Low Birth Weight Categories
Adjustment/Stratification: Statistical risk model The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, birthweight (500g groups), age in days (29-60, 61-90, 91+), age in years (in 5-year age groups), modified CMS DRG and AHRQ CCS comorbities. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 43 states and approximately 6 million pediatric discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Covariates used in this measures:
MDC 5 MDC 6
MDC 8 MDC 11
MDC 15 MDC OTHER
Procedure Type 2 Procedure Type 3 Procedure Type 4 to 5
Procedure Type 4 to 5 Procedure Type 6 Procedure Type 7
*** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures
2. Only minor therapeutic procedure with any or no diagnostic procedures
 One major therapeutic without diagnostic procedure One major therapeutic with only minor diagnostic procedure(s)
5. One major therapeutic with major diagnostic procedure(s)
6. Two major therapeutic procedures with any or no diagnostic procedures NOE REVIEW DRAFT_DO NOT CITE OR OUTOTE

0344 Accidental Puncture or Laceration Rate (PDI 1)
Three or more major therapeutic procedures with any or no diagnostic procedures; Clinical categories for PDI 1 are based on Major Diagnostic Categories (MDC).
Stratum 1. Eye, ear, nose, mouth, throat, skin, breast, and other low-risk procedures (MDC 2, 3, 9, 19, 22, 23)
Stratum 2. Thoracic, cardiovascular, and specified neoplastic procedures (MDC 4, 5, 17)
Stratum 3. Kidney, and male/female reproductive procedures
MDC 11, 12, 13) Stratum 4. Infectious, immunological, hematological, and ungroupable procedures
(MDC 0/99, 16, 18, 25) Stratum 5. Trauma, orthopedic, and neurologic procedures
(MDC 1, 8, 21, 24) Stratum 6. Gastrointestinal, hepatobiliary, and endocrine procedures
(MDC 6, 7, 10) Level of Analysis: Facility
Type of Measure: Outcome Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD.
Measure Steward: Agency for Healthcare Research and Quality
STEERING COMMITTEE MEETING 12/15-16/2011
1. Importance to Measure and Report: Y-18; N-3
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-3; M-2; L-1; I-0; 1b. Performance Gap: H-2; M-1; L-3; I-0
1c. Evidence Quantity: H-0; M-3; L-3; I-0; Quality: H-1; M-4; L-1; I-0; Consistency: H-1; M-2; L-3; I-0
<u>Rationale</u> : The Committee recognized that the measure affects small numbers of patients. They agreed that the key problem with accidental lacerations is those that occur without detection, resulting in a complication.
2. Scientific Acceptability of Measure Properties: Y-18; N- 3
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-2; M-3; L-1; I-0; 2b. Validity: H-;1 M-3; L-2; I-0
<u>Rationale</u> : The Committee noted that the risk adjustment could include additional factors, such as specialty or body part and that the measure's validity was impacted by the reliance on administrative data. The developer stated that work continues to determine if data from laboratories, electronic health records, and other sources could be incorporated into the measure to increase its validity. Coding updates and refinements are continuously made to address the issue and it has improved since the measure was initially developed.
 3. Usability: H-3; M-13; L-4; I-0 (<i>Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)</i> 3a. Public Reporting: H-0; M-4; L-1; I-0 3b. QI: H-1; M-4; L-0; I-0
Rationale: The Committee stated that the measure has been publicly reported for several years.
 4. Feasibility: H-8; M-10; L-2; I-0 (4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented) 4a. Byproduct of Care Processes: H-5; M-1; L-0; I-0
4b. Electronic data sources: H-4; M-1; L-1; I-0
 4c. Suscep inaccuracies, consequences: H-3; M-3; L-0; I-0 4d. Data collection strategy: H-5; M-1; L-0; I-0
Rationale: Given its reliance on administrative data, the measure is feasible as specified.
Steering Committee Recommendation for Endorsement: Y-19; N-2
Rationale: The measure is a useful indicator of quality by monitoring rates of accidental cuts, punctures, perforations, or lacerations among pediatric patients.
v. 1

0345 Accidental Puncture or Laceration Rate (PSI 15)
Measure Submission Form
Description: Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code
denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
Numerator Statement: Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code
denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
Denominator Statement: All surgical and medical discharges age 18 years and older defined by specific DRGs or MS-DRGs.
Exclusions: Exclude cases:
- with principal diagnosis denoting accidental cut, puncture, perforation, or laceration or secondary diagnosis present on admission
- MDC 14 (pregnancy, childbirth, and puerperium)
- with ICD-9-CM code for spine surgery
- with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis
(DX1=missing)
Adjustment/Stratification: Statistical risk model. The predicted value for each case is computed using a hierarchical model (logistic
regression with hospital random effect) and covariates for gender, age, modified CMS DRG, transfer status, procedure day availability,
and the AHRQ Comorbidity category. The reference population used in the regression is the universe of discharges for states that
participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 42 states and approximately 30 million
adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the
unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided
by the expected rate, multiplied by the reference population rate.
Covariates used in this measures:
Sex Female
Age 18 to 24 Age 25 to 29
5
Age 30 to 59
MDRG 101 MDRG 103
MDRG 103 MDRG 107
MDRG 302
MDRG 401
MDRG 402
MDRG 416
MDRG 502
MDRG 503
MDRG 504
MDRG 505
MDRG 506
MDRG 507
MDRG 508
MDRG 510
MDRG 511
MDRG 513
MDRG 514
MDRG 519
MDRG 520
MDRG 522
MDRG 601
MDRG 602
MDRG 603
MDRG 604
MDRG 606
MDRG 609 MDRG 610
MDRG 611

	cidental Puncture or Laceration Rate (PSI 15)
MDRG	621
MDRG	701
MDRG	702
MDRG	703
MDRG	704
MDRG	705
MDRG	712
MDRG	806
MDRG	807
MDRG	815
MDRG	816
MDRG	1001
MDRG	1003
MDRG	1005
MDRG	1006
MDRG	1101
MDRG	1102
MDRG	1103
MDRG	1104
MDRG	1105
MDRG	1107
MDRG	1109 1201
MDRG	1201
MDRG MDRG	1301
MDRG	1302
MDRG	1303
MDRG	1304
MDRG	1305
MDRG	1306
MDRG	1307
MDRG	1308
MDRG	1707
MDRG	1709
MDRG	1801
MDRG	1802
MDRG	2104
MDRG	2108
MDRG	2408
MDRG	7702
MDC	3
MDC	4
MDC	5
MDC	6
MDC	7
MDC	8
MDC	9
MDC	11
MDC	12
MDC	13
MDC	16
MDC	17
MDC	18
MDC	21

0345 Accidental	Puncture or Laceration Rate (PSI 15)
MDC 24	
MDC 24 MDC Other	
TRNSFER	Transfer-in
NOPRDAY	Procedure Days Data Not Available
COMORB	PERIVASC
COMORB	DM
COMORB	DMCX
COMORB	RENLFAIL
COMORB	OBESE
COMORB	WGHTLOSS
COMORB	BLDLOSS
COMORB	
Level of Analysis	ANEMDEF Not applicable
Type of Measure:	
	ninistrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
	rch and Quality, Rockville, MD.
	I: Agency for Healthcare Research and Quality MITTEE MEETING 12/15-16/2011
	Masure and Report: Y-20; N-2
	1b. Performance Gap, 1c. Evidence)
	1-2; L-0; I-0; Ib. Performance Gap: H-4; M-1; L-1; I-0
	ntity: H-;2 M-3; L-1; I-0; Quality: H-3; M-3; L-0; I-0; Consistency: H-2; M-3; L-1; I-0
IC. EVIDENCE QUA	Tuity. H-, 2 M-3, L-1, H0, Quality. H-3, M-3, L-0, H0, CONSISTENCY. H-2, M-3, L-1, H0
Rationale: The Co	ommittee recognized that the measure affects small numbers of patients. They agreed that the key problem with
	ons is those that occur without detection, resulting in a complication.
	eptability of Measure Properties: Y-20; N-2
	recise specifications, testing; 2b. Validity – testing, threats to validity)
	2 M-3; L-1; I-0; 2b. Validity: H-2; M-3; L-1; I-0
	2 10-3, $1-1$, $1-0$, 20 , 20 , $10-2$, $10-3$, $1-7$, $1-0$
	mmittee noted that the risk adjustment could include additional factors, such as specialty or body part and that the
	was impacted by the reliance on administrative data. The developer stated that work continues to determine if data
	electronic health records, and other sources could be incorporated into the measure to increase its validity. Coding
updates and refine	ements are continuously made to address the issue and it has improved since the measure was initially developed.
3. Usability: H-3;	M-16; L-3; I-0
(Meaningful, unde	rstandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
3a. Public Reportin	ng: H-3; M-2; L-1; I-0
3b. QI: H-4; M-0;	L-2; I-0
	ommittee stated that the measure has been publicly reported for several years.
4. Feasibility: H-9	
	generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences
	collection strategy can be implemented)
	Care Processes: H-5; M-1; L-0; I-0
	a sources: H-5; M-0; L-1; I-0
	iracies, consequences: H-4; M-2; L-0; I-0
4d. Data collection	n strategy: H-5; M-1; L-0; I-0
Rationale: Givon	its reliance on administrative data, the measure is feasible as specified.
	tee Recommendation for Endorsement: Y-20 ; N-2
Steering Commit	נכב הכנטוווווכוועמנוטון וטו בוועטו זכוווכוונ. ד-20 , וו-2
Rationale. The m	neasure is a useful indicator of quality by monitoring rates of accidental cuts, punctures, perforations, or lacerations
among adult patie	
among addit patie	

0362 Foreign Body left after procedure (PDI 3)

NATIONAL QUALITY FORUM
0362 Foreign Body left after procedure (PDI 3)
Measure Submission Form
Description: Count of discharges with foreign body left in during procedure in medical and surgical discharges among patients less than
18 years and not MDC 14 (pregnancy, childbirth, and puerperium)
Numerator Statement: Discharges under age 18 with ICD-9-CM codes for foreign body left in during procedure in any secondary
diagnosis field of medical and surgical discharges defined by specific DRGs or MS-DRGs where several exclusions are applied to the
numerator. (Details of the numerator, medical and surgical discharges DRGs and MS-DRGs, and exclusions appear in 2a1.3).
Denominator Statement: Not applicable
Exclusions: Not applicable
Adjustment/Stratification: No risk adjustment or risk stratification Not applicable Not applicable
Level of Analysis: Facility
Type of Measure: Outcome
Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
Healthcare Research and Quality, Rockville, MD.
Measure Steward: Agency for Healthcare Research and Quality
STEERING COMMITTEE MEETING 12/15-16/2011
1. Importance to Measure and Report: Y-22; N-0
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-4; M-1; L-0; I-0; 1b. Performance Gap: H-4; M-1; L-0; I-0
1c. Evidence Quantity: H-3; M-1; L-0; I-0; Quality: H-3; M-1; L-0; I-0; Consistency: H-3; M-1; L-0; I-0
Rationale: All agreed that this measure continues to address an important patient safety area. The Committee discussed the incidence
of foreign bodies being retained after a procedure and noted that once the statistics were further broken down to exclude foreign bodies
left behind intentionally, there appeared to be a much lower rate of occurrence. They also suggested that the measure name could be
changed to reduce confusion based on objects that were intentionally retained.
2. Scientific Acceptability of Measure Properties: Y-15; N- 7
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-3; M-1; L-1; I-0; 2b. Validity: H-;2 M-2; L-1; I-0
<u>Rationale</u> : The Committee debated at what point a foreign body would be considered "left after procedure" – i.e., at what point the
surgical procedure officially ends – and noted the differences between a foreign body being left intentionally after surgery versus a
foreign body left accidentally. Foreign bodies that affect the care management of a patient are counted in the measure and AHRQ confirmed that the definitions and time windows are consistent with the definitions for the similar serious reportable event (SRE). The
Committee noted that device fragments may be left intentionally to reduce the potential for further injury inflicted by retrieval and stated
that this would be coded as a "foreign body" within the measure as currently specified. The Committee requested that future versions of
the measure be stratified by intended retained bodies, unintended retained bodies, and device malfunctions. The developer indicated it
would be possible to capture this information through ICD-10 codes in the future.
3. Usability: H-2; M-12; L-8; I-0
(<i>Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement</i>)
3a. Public Reporting: H-1; M-3; L-1; I-0
3b. Ql: H-3; M-1; L-1; I-0
Rationale: The Committee questioned how the measure would improve quality and whether capturing the data would lead to a decrease
in foreign bodies left after a procedure but agreed that it continued to be useful for both consumers and providers.
4. Feasibility: H-6; M-11; L-5; I-0
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences
identified 4d. Data collection strategy can be implemented)
4a. Byproduct of Care Processes: H-5; M-0; L-0; I-0
4b. Electronic data sources: H-5; M-0; L-0; I-0
4c. Suscep inaccuracies, consequences: H-3; M-1; L-1; I-0
4d. Data collection strategy: H-3; M-1; L-1; I-0
Rationale: They encouraged the developer to utilize codes in the future that would reflect irretrievable device fragments, to differentiate
between types of foreign bodies left after a procedure, which will hopefully be achieved when ICD10 is implemented. Because this
measure is collected using administrative data, it was considered feasible.
Steering Committee Recommendation for Endorsement: Y- 17; N-4
Rationale: The Committee agreed the measure was important and encouraged the developer to further differentiate between types of
foreign bodies left after procedure in future iterations.

363 Foreign Body Left During Procedure (PSI 5)
Aeasure Submission Form
Description: Count of discharges with foreign body left in during procedure in medical and surgical discharges among patients 18 years nd older or MDC 14 (pregnancy, childbirth, and puerperium)
Iumerator Statement: Discharges, 18 years and older or MDC 14 (pregnancy, childbirth, and puerperium), with ICD-9-CM codes for
preign body left in during procedure in any secondary diagnosis field of medical and surgical discharges defined by specific DRGs or
IS-DRGs. (Details of medical and surgical discharges defined by specific DRGs or MS-DRGs and exclusions noted in 2a1.3).
Denominator Statement: Not applicable
xclusions: Not applicable
djustment/Stratification: No risk adjustment or risk stratification Not applicable Not applicable
evel of Analysis: Facility
ype of Measure: Outcome
Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
lealthcare Research and Quality, Rockville, MD.
leasure Steward: Agency for Healthcare Research and Quality
TEERING COMMITTEE MEETING 12/15-16/2011
. Importance to Measure and Report: Y-22; N-0
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
a. Impact: H-4; M-1; L-0; I-0; 1b. Performance Gap: H-4; M-1; L-0; I-0
c. Evidence Quantity: H-3; M-1; L-0; I-0; Quality: H-3; M-1; L-0; I-0; Consistency: H-3; M-1; L-0; I-0
Rationale: All agreed that this measure continues to address an important patient safety area. The Committee discussed the incidence
f foreign bodies being retained after a procedure and noted that once the statistics were further broken down to exclude foreign bodies
eft behind intentionally, there appeared to be a much lower rate of occurrence. They also suggested that the measure name could be
hanged to reduce confusion based on objects that were intentionally retained.
. Scientific Acceptability of Measure Properties: Y-15; N- 7
2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
a. Reliability: H-3; M-1; L-1; I-0; 2b. Validity: H-;2 M-2; L-1; I-0
Rationale: Rationale: The Committee debated at what point a foreign body would be considered "left after procedure" – i.e., at what point
ne surgical procedure officially ends – and noted the differences between a foreign body being left intentionally after surgery versus a
preign body left accidentally. Foreign bodies that affect the care management of a patient are counted in the measure and AHRQ
onfirmed that the definitions and time windows are consistent with the definitions for the similar serious reportable event (SRE). The
Committee noted that device fragments may be left intentionally to reduce the potential for further injury inflicted by retrieval and stated
nat this would be coded as a "foreign body" within the measure as currently specified. The Committee requested that future versions of
ne measure be stratified by intended retained bodies, unintended retained bodies, and device malfunctions. The developer indicated it
rould be possible to capture this information through ICD-10 codes in the future.
. Usability: H-2; M-12; L-8; I-0
Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
a. Public Reporting: H-1; M-3; L-1; I-0
b. QI: H-3; M-1; L-1; I-0
Rationale: The Committee questioned how the measure would improve quality and whether capturing the data would lead to a decrease
n foreign bodies left after a procedure but agreed that it continued to be useful for both consumers and providers.
. Feasibility: H-6; M-11; L-5; I-0
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences
dentified 4d. Data collection strategy can be implemented)
a. Byproduct of Care Processes: H-5; M-0; L-0; I-0
b. Electronic data sources: H-5; M-0; L-0; I-0
c. Suscep inaccuracies, consequences: H-3; M-1; L-1; I-0
d. Data collection strategy: H-3; M-1; L-1; I-0
Ationale: They encouraged the developer to utilize codes in the future that would reflect irretrievable device fragments, to differentiate
etween types of foreign bodies left after a procedure, which will hopefully be achieved when ICD10 is implemented. Because this
neasure is collected using administrative data, it was considered feasible.
Steering Committee Recommendation for Endorsement: Y- 17; N-4
Detionaley. The Committee agreed the measure was important and ensured and the developer to further differentiate between twee of
Rationale: The Committee agreed the measure was important and encouraged the developer to further differentiate between types of

NATIONAL QUALITY FORUM
0363 Foreign Body Left During Procedure (PSI 5)
foreign bodies left after procedure in future iterations.
0263 Patient Burn
Measure Submission Form
Description: Percentage of ASC admissions experiencing a burn prior to discharge
Numerator Statement: Ambulatory surgical center (ASC) admissions experiencing a burn prior to discharge.
Denominator Statement: All ASC admissions.
Exclusions: None
Adjustment/Stratification: No risk adjustment or risk stratification. None. This measure is not stratified.
Level of Analysis: Facility
Type of Measure: Outcome
Data Source: Paper Records ASC medical records, as well as incident/occurrence reports, and variance reports may serve as data
sources. No specific collection instrument is required although the ASC Quality Collaboration has developed a sample data collection
instrument that may be used as desired. Facilities may use any collection instrument that allows tracking of all burns prior to discharge.
Measure Steward: Ambulatory Surgical Center Quality Collaboration
STEERING COMMITTEE MEETING 12/15-16/2011
Importance to Measure and Report: Y-22; N-0
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-4; M-1; L-1; I-0; 1b. Performance Gap: H-2; M-3; L-1; I-0
1c. Evidence Quantity: H-2; M-1; L-2; I-0; Quality: H-1; M-2; L-2; I-0; Consistency: H-2; M-2; L-1-; I-0
Rationale: The Committee agreed that while a patient burn is a rare event, it could lead to serious consequences for both the patient and
hospital staff. This measure would provide an avenue for ambulatory surgical centers to collect data on a serious reportable event.
2. Scientific Acceptability of Measure Properties: Y-21; N-1
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-4; M-2; L-0; I-0; 2b. Validity: H-3; M-3; L-0; I-0
Rationale: The measure provides a precise definition of burns, which is designed to capture the variety of ways a patient could be
injured.
3. Usability: H-15; M-5; L-0; I-0
(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
3a. Public Reporting: H-5; M-0; L-0; I-1
3b. QI: H-5; M-0; L-0; I-1
Rationale: The measure will lead to public reporting and quality improvement of a serious reportable event in the ambulatory setting.
4. Feasibility: H-18; M-5; L-0; I-0
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences
identified 4d. Data collection strategy can be implemented)
4a. Byproduct of Care Processes: H-2; M-2; L-0; I-1
4b. Electronic data sources: H-1; M-1; L-2; I-1
4c. Suscep inaccuracies, consequences: H-2; M-1; L-1; I-1
4d. Data collection strategy: H-3; M-0; L-1; I-1
Rationale: The Committee expressed concern that some burns may not be captured due to a patient's short length of stay, but
acknowledged that these events were already being voluntarily reported
Steering Committee Recommendation for Endorsement: Y- 22; N-0
Detionals. While notions have a core quant they can lead to parisus accore manage. The management will relate a surgery when the
Rationale: While patient burns are a rare event, they can lead to serious consequences. The measure will raise awareness about the
varying types of burns that may result in patient injury.
024/ Introgenia Droumetherov Data (DSI ()
0346 latrogenic Pneumothorax Rate (PSI 6)

Measure Submission Form

Description: Percent of discharges with ICD-9-CM code for iatrogenic pneumothorax in any secondary diagnosis field among cases meeting the inclusion and exclusion rules for the denominator

Numerator Statement: Discharges with ICD-9-CM code for iatrogenic pneumothorax in any secondary diagnosis field among cases meeting the inclusion and exclusion rules for the denominator.

Denominator Statement: All surgical and medical discharges age 18 years and older defined by specific DRGs or MS-DRGs.

	neumothorax Rate (PSI 6)
Exclusions: Exclu	
	gnosis of iatrogenic pneumothorax or secondary diagnosis present on admission
	ncy, childbirth, and puerperium)
	s code of chest trauma or pleural effusion
	phragmatic surgery repair in any procedure field
- with any code ind	licating thoracic procedure, lung or pleural biopsy, or cardiac procedure
- with missing gene	der (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis
(DX1=missing)	
	fication: Statistical risk model. The predicted value for each case is computed using a hierarchical model (logistic
	spital random effect) and covariates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRO
	bry. The reference population used in the regression is the universe of discharges for states that participate in the
	ent Data (SID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges.
	is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of
	tal). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected
	the reference population rate.
Sex Female	
Age 65 to 85-	+
MDRG 416	
MDRG 504	
MDRG 510	
MDRG 601	
MDRG 602	
MDRG 1103	
MDRG 1801	
MDRG 1807	
MDC 1	
MDC 6	
MDC 8	
MDC 25	
NOPRDAY	Procedure Days Data Not Available
COMORB	HTN_C
COMORB	NEURO
COMORB	CHRNLUNG
COMORB	
	DM
COMORB	DMCX
COMORB	METS
COMORB	OBESE
COMORB	WGHTLOSS
COMORB	DRUG Not applicable
Level of Analysis	
Type of Measure:	
	ninistrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
	ch and Quality, Rockville, MD.
	: Agency for Healthcare Research and Quality
	AITTEE MEETING 12/15-16/2011
	Measure and Report: Y-18; N-1
	1b. Performance Gap, 1c. Evidence)
	-2; L-0; I-0; Ib. Performance Gap: H-3; M-1; L-0; I-0
	ntity: H-1; M-2; L-1; I-0; Quality: H-2; M-1; L-1; I-0; Consistency: H-1; M-2; L-1; I-0
	easure indicates a small performance gap, but focuses on an event which is relatively common. Additionally, it may be
	fferences in performance between hospitals based on low volumes of iatrogenic pneumothoraxes. However, the
	t it was important to capture these serious adverse events, many of which are preventable.
	ptability of Measure Properties: Y-17; N-2
(2a. Reliability – pr	ecise specifications, testing; 2b. Validity – testing, threats to validity)

0346 latrogenic Pneumothorax Rate (PSI 6)

2a. Reliability: H-3; M-1; L-0; I-0; 2b. Validity: H-4; M-0; L-0; I-0

<u>Rationale</u>: The measure is derived from administrative claims data, which has been shown to be consistent and reliable. The Committee noted that the measure had a number of exclusions but agreed that they were necessary and reasonable. The Committee encouraged the developer to continue work on appropriate validation studies. The positive predictive values both for the adult and pediatric measures were low but determined to be acceptable.

3. Usability: H-6; M-12; L-1; I-0

(*Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement*) **3a.** Public Reporting: **H-3**; **M-1**; **L-0**; **I-0**

3b. QI: H-3; M-1; L-0; I-0

Rationale: The Committee stated that the measure has been reported in the public domain and has led to quality improvement.

4. Feasibility: H-9; M-9; L-1; I-0

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-3; M-1; L-0; I-0

4b. Electronic data sources: H-3; M-1; L-0; I-0

4c. Suscep inaccuracies, consequences: H-3; M-1; L-0; I-0

4d. Data collection strategy: H-3; M-1; L-0; I-0

<u>Rationale</u>: The measure has been updated to reduce the likelihood of inaccuracies and appropriately capture the iatrogenic pneumothorax rate. Because this measure is collected using administrative data, it was considered feasible..

Steering Committee Recommendation for Endorsement: Y- 18; N-1

Rationale: The measure continues to provide information on an event that may be preventable and facilitates quality improvement.

0348 latrogenic Pneumothorax Rate (PDI 5)

Measure Submission Form

Description: Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code of iatrogenic pneumothorax in any secondary diagnosis field

Numerator Statement: Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code of iatrogenic pneumothorax in any secondary diagnosis field

Denominator Statement: Discharges, age under 18 years, defined by specific surgical and medical DRGs

Exclusions: Exclude cases:

- neonates with birth weight less than 2500 grams (Birth Weight Category 1-8)

- with principal diagnosis of iatrogenic pneumothorax or secondary diagnosis present on admission
- with any diagnosis code of chest trauma or pleural effusion
- with an ICD-9-CM procedure code of thoracic surgery, lung or pleural biopsy, diaphragmatic surgery repair, OR cardiac surgery
- normal newborn
- MDC 14 (pregnancy, childbirth, and puerperium)

- with missing discharge gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)

Adjustment/Stratification: Statistical risk model The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, birthweight (500g groups), age in days (29-60, 61-90, 91+), age in years (in 5-year age groups), modified CMS DRG and AHRQ CCS comorbities. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 43 states and approximately 6 million pediatric discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate.

Age in Years 13 to 18

Age in Years 1 to 13 Not applicable

Level of Analysis: Facility

Type of Measure: Outcome

Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD.

Measure Steward: Agency for Healthcare Research and Quality

STEERING COMMITTEE MEETING 12/15-16/2011

0240 Jahrs mania Draumath arou Data (DDI E)
0348 latrogenic Pneumothorax Rate (PDI 5)
1. Importance to Measure and Report: Y-18; N-1
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-2; M-2; L-0; I-0; 1b. Performance Gap: H-3; M-1; L-0; I-0
1c. Evidence Quantity: H-1; M-2; L-1; I-0; Quality: H-2; M-1; L-1; I-0; Consistency: H-1; M-2; L-1; I-0
Rationale: The Committee noted that the performance gap had decreased on the measure over time. However, a continued reduction in
the prevalence of these events shows that the performance gap can still be improved.
2. Scientific Acceptability of Measure Properties: Y-17; N-2
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-3; M-1; L-0; I-0; 2b. Validity: H-4; M-0; L-0; I-0
Rationale: The measure is derived from administrative claims data, which has been shown to be consistent and reliable. The
Committee noted that the measure had a number of exclusions but agreed that they were necessary and reasonable. The Committee
noted that the developer should continue to work on appropriate validation studies. The positive predictive values both for the adult and
pediatric measures were low but determined to be acceptable.
3. Usability: H-6; M-12; L-1; I-0
(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
3a. Public Reporting: H-3; M-1; L-0; I-0
3b. QI: H-3; M-1; L-0; I-0
Rationale: The Committee stated that the measure has been reported in the public domain and led to quality improvement.
4. Feasibility: H-9; M-9; L-1; I-0
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences
identified 4d. Data collection strategy can be implemented)
4a. Byproduct of Care Processes: H-3; M-1; L-0; I-0
4b. Electronic data sources: H-3; M-1; L-0; I-0
4c. Suscep inaccuracies, consequences: H-3; M-1; L-0; I-0
4d. Data collection strategy: H-3; M-1; L-0; I-0
Rationale: The measure has been updated to reduce the likelihood of inaccuracies and appropriately capture the iatrogenic
pneumothorax rate. Because this measure is collected using administrative data, it was considered feasible
Steering Committee Recommendation for Endorsement: Y- 18; N-1
Rationale: The measure continues to provide information on an event that may be preventable and facilitates quality improvement.
MEASURES RECOMMENDED FOR RESERVE STATUS

0349 Transfusion Reaction (PSI 16)

Measure Submission Form

Description: The count of medical and surgical discharges for patients age greater than or equal to 18 or in MDC 14 with ICD-9-CM code for transfusion reaction in any secondary diagnosis field.

Numerator Statement: Discharges 18 years and older or in MDC 14 with ICD-9-CM codes for transfusion reaction in any secondary diagnosis field of all medical and surgical discharges defined by specific DRGs or MS-DRGs

See Patient Safety Indicators Appendices:

- Appendix B – Medical Discharge DRGs

- Appendix C – Medical Discharge MS-DRGs

- Appendix D – Surgical Discharge DRGs

- Appendix E – Surgical Discharge MS-DRGs

Link to PSI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf Exclude cases:

-with principal diagnosis of transfusion reaction or secondary diagnosis present on admission

-with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)

Denominator Statement: Not applicable

Exclusions: Not applicable

Adjustment/Stratification: No risk adjustment or risk stratification Not applicable Not applicable

Level of Analysis: Facility

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0349 Transfusion Reaction (PSI 16)
Type of Measure: Outcome
Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
Healthcare Research and Quality, Rockville, MD.
Measure Steward: Agency for Healthcare Research and Quality
STEERING COMMITTEE MEETING 12/15-16/2011
1. Importance to Measure and Report: Y-6; N-15
(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
1a. Impact: H-2; H-2; L-0; I-1; 1b. Performance Gap: H-2; H-1; L-2; I-0
1c. Evidence Quantity: H-2; M-2; L-1; I-0; Quality: H-2; M-2; L-1; I-0; Consistency:
Rationale: The Committee questioned whether the measure would reduce transfusion reactions as the performance rate is currently low.
However, the Committee agreed that collecting data on transfusion reactions may be used to reduce events in the future. The
Committee also suggested that the measure's title may be wrongly interpreted to indicate a patient being given the wrong blood, when it
collects data on a variety of transfusion reactions, such as reactions to antigens. The Committee affirmed that the low number of events
provides evidence of industry success at managing transfusions and still meets two of the three criteria – high impact and evidence.
2. Scientific Acceptability of Measure Properties: Y-19; N- 2
(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity)
2a. Reliability: H-3; M-1; L-0; I-1; 2b. Validity: H-3; M-1; L-0; I-1
Rationale: The Committee stated that the measure provides precise specifications to count a variety of transfusion events and is used to
monitor a serious reportable event.
3. Usability: H-5; M-10; L-6; I-
(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)
3a. Public Reporting: H-2; M-2; L-1; I-0 3b. QI: H-2; M-1; L-2; I-0
Rationale: The Committee expressed concern about the value of publicly reporting transfusion reactions and whether it would affect
quality improvement. However, they also noted the measure could be easily interpreted by the public.
4. Feasibility: H-14; M-;5 L-2; I-
(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences
identified 4d. Data collection strategy can be implemented)
4a. Byproduct of Care Processes: H-4; M-1; L-0; I-0
4b. Electronic data sources: H-4; M-1; L-0; I-0
4c. Suscep inaccuracies, consequences: H-3; M-1; L-1; I-0
4d. Data collection strategy: H-4; M-1; L-0; I-0
Rationale: The measure has been implemented for a number of years, using administrative data, indicating that it is feasible to collect.
Steering Committee Recommendation for Endorsement: Reserve Status Y- 19; N-1
Rationale: The measure provides important information to the industry and consumers, highlighting a small but important performance
gap. The Committee was concerned of the implications if endorsement was removed and no longer reported because of the low
performance gap. Based on the Committee's discussion and votes, the measure continues to meet all of the criteria with the exception
of an opportunity for improvement. The Committee stressed that this achievement should be celebrated but also it should be monitored
to ensure that this event continues to be low; thus, they recommend that the measure be endorsed with reserve status.
0350 Transfusion Reaction (PDI 13)

Measure Submission Form

Description: The count of medical and surgical discharges for patients age less than 18 and not in MDC 14 with ICD-9-CM code for transfusion reaction in any secondary diagnosis field.

Numerator Statement: Discharges under age 18 with ICD-9-CM codes for transfusion reaction in any secondary diagnosis field of all medical and surgical discharges defined by specific DRGs or MS-DRGs with the exclusion of neonates, cases in MDC 14 and instances with the outcome of interest was present on admission.

See Pediatric Quality Indicators Appendices:

- Appendix B Surgical DRGs
- Appendix C Surgical MS-DRGs
- Appendix D Medical DRGs
- Appendix E Medical MS-DRGs
- Appendix I Definitions of, Neonate, Newborn, Normal Newborn, and Outborn

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	0350 Transfusion Reaction (PDI 13)
	Link to PDI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf
	Cases excluded with missing gender (SEX=missig, age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal
	diagnosis (DX1=missing)
	Denominator Statement: Not applicable
	Exclusions: Not applicable
	Adjustment/Stratification: No risk adjustment or risk stratification Not applicable Not applicable
	Level of Analysis: Facility
	Type of Measure: Outcome
	Data Source: Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for
	Healthcare Research and Quality, Rockville, MD.
ļ	Measure Steward: Agency for Healthcare Research and Quality
	STEERING COMMITTEE MEETING 12/15-16/2011
	1. Importance to Measure and Report: Y-6; N-15
	(1a. High Impact: 1b. Performance Gap, 1c. Evidence)
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	quality improvement. However, they also noted the measure could be easily interpreted by the public.
ľ	4. Feasibility: H-14; M-;5 L-2; I-
	(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences
	identified 4d. Data collection strategy can be implemented)
	4a. Byproduct of Care Processes: H-4; M-1; L-0; I-0
	4b. Electronic data sources: H-4; M-1; L-0; I-0
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	Steering Committee Recommendation for Endorsement: Reserve Status Y- 19; N-1
	Rationale. The measure provides important information to the industry and consumers, highlighting a small but important performance
	performance gap. Based on the Committee's discussion and votes, the measure continues to meet all of the criteria with the exception
-	 4. Feasibility: H-14; M-;5 L-2; I- (<i>4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c. Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented</i>) 4a. Byproduct of Care Processes: H-4; M-1; L-0; I-0 4b. Electronic data sources: H-4; M-1; L-0; I-0 4c. Suscep inaccuracies, consequences: H-3; M-1; L-1; I-0 4d. Data collection strategy: H-4; M-1; L-0; I-0 4d. Data collection strategy: H-4; M-1; H-0;

of an opportunity for improvement. The Committee stressed that this achievement should be celebrated but also it should be monitored to ensure that this event continues to be low; thus, they recommend that the measure be endorsed with reserve status.

MEASURES NOT RECOMMENDED

0374 Venous Thromboembolism Patients Recieving Unfractionated Heparin with Dosages/Platelet Count Monitoring by Protocol or Nomogram

Measure Submission Form

	ents Recieving Unfractionated Heparin with Dosages/Platelet Count Monitoring by
Protocol or Nomogram	
intravenous (IV) unfractionated heparin (as a nomogram or protocol. This measu Prophylaxis, VTE-2: ICU VTE Prophylax Discharge Instructions and VTE-6: Incide	we their IV UFH therapy dosages AND platelet counts monitored according to defined
Denominator Statement: Patients with with an ICD-9-CM Principal or Other Dia	confirmed VTE receiving IV UFH therapy. The target population includes patients discharged gnosis Codes for VTE as defined in Table 7.03 or Table 7.04.
Exclusions:	
 Patients less than 18 years of age 	
Patients who have a length of stay grea Patients with Comfort Measures Only de Detients arralled in aligned trials	
Patients enrolled in clinical trials	194. for bouries core
Patients discharged to a health care fac	
Patients discharged to home for hospice	ecare
Patients who expired	
•Patients who left against medical advice	
Patients discharged to another hospital	
Patients without UFH Therapy Administ	
Patients without VTE confirmed by diag	
	ustment or risk stratification Not applicable Not applicable
evel of Analysis: Facility, Population :	National
ype of Measure: Process	
	ctronic Clinical Data, Paper Records. Each element in the data dictionary includes suggested
based on the measure specifications. Th	ng contracted Performance Measurement Systems (vendors) that develop data collection tools tools are verified and tested by Joint Commission staff to confirm the accuracy and vith the specifications. The vendor may not offer the measure set to hospitals until verification
has been passed.	
Measure Steward: The Joint Commission	on
STEERING COMMITTEE MEETING 12/	
1. Importance to Measure and Report:	: Y-10; N-11
(1a. High Impact: 1b. Performance Gap	
1a. Impact: H-8; M-1; L-0; I-0; 1b. Perfo	
	0; Quality: H-5; M-4; L-0; I-0; Consistency: H-7; M-2; L-0; I-0
Rationale: The Committee expressed co range was achieved. There was evidence	ncern that the measure focused only on the use of a nomogram, and not whether therapeutic ce to support the measure focus and a gap exists. Because the vote on whether the measure ort, the Committee continued discussions on the remaining criteria.
2. Scientific Acceptability of Measure	
	esting; 2b. Validity – testing, threats to validity)
Rationale: The Committee was concerne	ed that the measure only applied to a small number of patients. Additionally, the Committee ogram was not a direct indication of improvement in patient care. There was concern related to
he validity of the measure as it is not me	easuring what is of most interest – whether therapeutic range was achieved. Also, it was nould be its own measure rather than included here.
3. Usability: H-; M-; L-; I- (<i>Meaningful, understandable, and usefu</i> 3a. Public Reporting: H-9; M-0; L-0; I-0	I to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvemer
3b. QI: H-9; M-0; L-0; I-0 Rationale:	
4. Feasibility: H-; M-; L-; I- (4a. Clinical data generated during care identified 4d. Data collection strategy ca	e delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences n be implemented)

0374 Venous Thromboembolism Patients Recieving Unfractionated Heparin with Dosages/Platelet Count Monitoring by Protocol or Nomogram

4a. Byproduct of Care Processes: H-7; M-2; L-0; I-0

4b. Electronic data sources: H-6; M-3; L-0; I-0

4c. Suscep inaccuracies, consequences: H-6; M-3; L-1; I-0

4d. Data collection strategy: H-5; M-3; L-1; I-0

Rationale:

Steering Committee Recommendation for Endorsement: Did not pass the Scientific Acceptability of Measure Properties criteria, which is required for endorsement.

Rationale: The goal of the measure is for a patient to be within the therapeutic range; however, measuring the use of a nomogram alone does not necessarily lead to an improvement in patient outcomes.

0375 Venous Thromoboembolism Warfarin Therapy Discharge Instructions

Measure Submission Form

Description: This measure assesses the number of patients diagnosed with confirmed VTE that are discharged on warfarin to home, home with home health or home hospice with written discharge instructions that address all four criteria: compliance issues, dietary advice, follow-up monitoring, and information about the potential for adverse drug reactions/interactions. This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring by Protocol and VTE-6: Incidence of Potentially-Preventable VTE). Numerator Statement: Patients with documentation that they or their caregivers were given written discharge instructions or other educational material about warfarin that addressed all of the following: 1. compliance issues 2. dietary advice 3. follow-up monitoring 4. potential for adverse drug reactions and interactions **Denominator Statement:** Patients with confirmed VTE discharged on warfarin therapy. The target population includes patients discharged with an ICD-9-CM Principal or Other Diagnosis Codes for VTE as defined in Table 7.03 or Table 7.04 that are discharged to home, homecare or court/law enforcement or home for hospice care. Please note: The allowable values of the data element Discharge Disposition are used to designate which locations are included. Exclusions: • Patients less than 18 years of age • Patients who have a length of stay greater than 120 days · Patients enrolled in clinical trials Patients without Warfarin Prescribed at Discharge Patients without VTE confirmed by diagnostic testing Adjustment/Stratification: No risk adjustment or risk stratification. Not applicable Not applicable, the measure is not stratified. Level of Analysis: Facility, Population : National Type of Measure: Process Data Source: Administrative claims, Electronic Clinical Data, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. The vendor may not offer the measure set to hospitals until verification has been passed. Measure Steward: The Joint Commission STEERING COMMITTEE MEETING 12/15-16/2011 1. Importance to Measure and Report: Y-10; N-11 (1a. High Impact: 1b. Performance Gap, 1c. Evidence) 1a. Impact: H-7; M-2; L0-; I-0; 1b. Performance Gap: H-6; M-3; L-0; I-0 1c. Evidence Quantity: H-1; M-7; L-1; I-0; Quality: H-0; M-4; L-4; I-0; Consistency: H-4; M-3; L-1; I-1 Rationale: The measure documents whether patients were provided with written instructions for the use of warfarin therapy at discharge. However, the measure is limited in that it does not assess a patient's understanding of the discharge instructions nor the effectiveness of the education (i.e., improved compliance post discharge). An opportunity for improvement does continue to exist but it was not clear whether the measure's continued use would lead to further improvement in patient outcomes. Because the vote on whether the measure passed importance to measure and report, the Committee continued discussions on the remaining criteria.

0375 Venous Thromoboembolism Warfarin Therapy Discharge Instructions

2. Scientific Acceptability of Measure Properties: Y-4; N-17

(2a. Reliability - precise specifications, testing; 2b. Validity - testing, threats to validity)

2a. Reliability: H-4; M-5; L-0; I-0; 2b. Validity: H-2; M-;6 L-1; I-0

<u>Rationale</u>: The Committee underscored the importance of patient education but noted that the measure instructions should be offered in a patient's native language to ensure understanding. They encouraged the developer to continue working on measures focused on patient education but to ensure that the measure uses validated educational materials. A Committee member suggested a 24 hour postdischarge follow-up phone call could be used to clarify how well instructions were adhered to by the patient. Because the measure did not pass scientific acceptability, the Committee did not discuss the remaining criteria.

3. Usability: H-; M-; L-; I-

(*Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement*) **3a.** Public Reporting: H-4; M-5; L-0; I-0

3b. QI: H-6; M-2; L-1; I-0

Rationale:

4. Feasibility: H-; M-; L-; I-

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-6; M-2; L-1; I-0

4b. Electronic data sources: H-4; M-4; L-1; I-0

4c. Suscep inaccuracies, consequences: H-5; M-3; L-1; I-0

4d. Data collection strategy: H-6; M-2; L-1; I-0

Rationale:

Steering Committee Recommendation for Endorsement: Did not pass the Importance to Measure and Report and the Scientific Acceptability of Measure Properties criteria, which are required for endorsement.

Rationale: The measure may not directly lead to an improvement in patient outcomes and lacks validated educational materials.

0501 Confirmation of Endotracheal Tube Placement

Measure Submission Form

Description: Any time an endotracheal tube is placed into a patients airway in the Emergency Department (ED)or a patient arrives to the ED with an endotracheal tube already in place (via EMS or hospital transfer) there should be appropriate confirmation of ETT placement and documentation of its performance in the medical record.

Numerator Statement: Number of ED patients with an endotracheal tube(ETT) placed or assessed with an endotracheal already in place who had the ETT confirmation performed

Denominator Statement: Total number of endotracheal tubes evaluated including those patients who had an ETT's placed in the ED and those patients who arruived to the ED with an ETT already in palce.

Exclusions: No exclusions

Adjustment/Stratification: No risk adjustment or risk stratification

Level of Analysis: Clinician : Group/Practice, Clinician : Individual, Facility, Integrated Delivery System, Population : Community Type of Measure: Process

Data Source: Administrative claims, Electronic Clinical Data, Electronic Clinical Data : Electronic Health Record, Electronic Clinical Data : Registry, Paper Records The data will need to be collected from each patient's medical record. For those patients that are intubated in the Emergency Department, there will likely be a billed procedure for an endotracheal tube intubation. Other charts like patients who expired or patients who admitted to an ICU may be another source of identification of patients who either arrived intubated or are intubated in surveillance mechanism is in place (i.e., airway registry) is in place to capture all patients who either arrived intubated or are intubated in the Emergency Department then the data can be collected from there.

Measure Steward: Cleveland Clinic

STEERING COMMITTEE MEETING 12/15-16/2011

1. Importance to Measure and Report: Y-17; N-3

(1a. High Impact: 1b. Performance Gap, 1c. Evidence)

1a. Impact: H-6; M-3; L-0; I-0; 1b. Performance Gap: H-6; M-3; L-0; I-0

1c. Evidence Quantity: H-3; M-5; L-1; I-0; Quality: H-3; M-4; L-2; I-0; Consistency: H-3; M-4; L-2; I-0

<u>Rationale</u>: The Committee acknowledged the performance gap and that was a strong correlation between an incorrect endotracheal tube placement and morbidity and mortality. They stated that providers may be unfamiliar with best practices and the appropriate methods for assessing endotracheal tube placement but there is evidence in support of specific methods.

0501 Confirmation of Endotracheal Tube Placement 2. Scientific Acceptability of Measure Properties: Y-6; N-14 (2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity) 2a. Reliability: H-3; M-3; L-2; I-1; 2b. Validity: H-4; M-4; L-1; I-0 Rationale: The Committee noted that the specifications should include further definitions of what was considered appropriate confirmation, and were concerned about the variability of ET tube placement confirmation. In the future, the Committee encouraged the developer to expand the measure to additional care settings beyond the emergency department. 3. Usability: H-: M-: L-: I-(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement) 3a. Public Reporting: H-5; M-4; L-0; I-0 3b. QI: H-5; M-4; L-0; I-0 Rationale: 4. Feasibility: H-; M-; L-; I-(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented) 4a. Byproduct of Care Processes: H-3; M-4; L-2; I-0 4b. Electronic data sources: H-2; M-3; L-4; I-0 4c. Suscep inaccuracies, consequences: H-3; M-4; L-2; I-0 4d. Data collection strategy: H-3; M-4; L-2; I-0 Rationale: Steering Committee Recommendation for Endorsement: Did not pass the Importance to Measure criteria, which is required for endorsement. Rationale: The Committee agreed that the measure should be further specified in the future. 0523 Pain Assessment Conducted Measure Submission Form Description: Percentage of home health episodes of care in which the patient was assessed for pain, using a standardized pain assessment tool, at start/resumption of care. Numerator Statement: Number of home health episodes of care in which the patient was assessed for pain, using a standardized pain assessment tool, at start/resumption of care. Denominator Statement: Number of home health episodes of care ending during the reporting period, other than those covered by generic exclusions. Exclusions: No measure specific exclusions. See details of generic exclusions in 2a1.9. Adjustment/Stratification: No risk adjustment or risk stratification. N/A - process measure. N/A - measure not stratified Level of Analysis: Facility Type of Measure: Process Data Source: Electronic Clinical Data OASIS-C Measure Steward: Centers for Medicare & Medicaid STEERING COMMITTEE MEETING 12/15-16/2011 1. Importance to Measure and Report: Y-8; N-11 (1a. High Impact: 1b. Performance Gap, 1c. Evidence) 1a. Impact: H-1; M-4; L-; I-0; 1b. Performance Gap: H-0; M-5; L-; I-0 1c. Evidence Quantity: H-3; M-1; L-1; I-0; Quality: H-2; M-1; L-2; I-0; Consistency: H-3; M-0; L-2; I-0 Rationale: The Committee agreed that pain should be assessed across the continuum of care and during each visit for patients who are receiving home care but noted that there is little evidence that pain assessment alone does not improve outcomes. The Committee encouraged the developer to link the measure to an appropriate pain treatment plan in the future. Because the measure did not pass importance to measure and report, the Committee did not discuss the remaining criteria. 2. Scientific Acceptability of Measure Properties: Y-; N-(2a. Reliability – precise specifications, testing; 2b. Validity – testing, threats to validity) 2a. Reliability: H-3; M-0; L-1; I-1; 2b. Validity: H-2; M-1; L-1; I-1 Rationale:

3. Usability: H-; M-; L-; I-

(Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement)

0523 Pain Assessment Conducted

3a. Public Reporting: H-2; M-1; L-1; I-1 3b. QI: H-2; M-1; L-1; I-1

Rationale:

4. Feasibility: H-; M-; L-; I-

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-3; M-0; L-1; I-1

4b. Electronic data sources: H-3; M-0; L-1; I-1

4c. Suscep inaccuracies, consequences: H-3; M-0; L-1; I-1

4d. Data collection strategy: H-3; M-0; L-1; I-1

Rationale:

Steering Committee Recommendation for Endorsement: Did not pass the Importance to Measure criteria, which is required for endorsement.

Rationale: The evidence does not indicate that the assessment of pain alone will lead to an improvement in patient outcomes.

0524 Pain Interventions Implemented During Short Term Episodes Of Care

Measure Submission Form

Description: Percentage of short term home health episodes of care during which pain interventions were included in the physicianordered plan of care and implemented.

Numerator Statement: Number of home health episodes of care during which pain interventions were included in the physician-ordered plan of care and implemented.

Denominator Statement: Number of home health episodes of care ending during the reporting period, other than those covered by generic or measure-specific exclusions.

Exclusions: Episodes in which the patient did not have pain since the last OASIS assessment, as evidenced by a formal assessment that indicated no pain. Long-term episodes (as indicated by the presence of a follow-up assessment between admission and transfer or discharge). Episodes ending in patient death.

Adjustment/Stratification: No risk adjustment or risk stratification. N/A - process measure. N/A measure not stratified.

Level of Analysis: Facility

Type of Measure: Process

Data Source: Electronic Clinical Data OASIS-C

Measure Steward: Centers for Medicare & Medicaid

STEERING COMMITTEE MEETING 12/15-16/2011

1. Importance to Measure and Report: Y-7; N-12

(1a. High Impact: 1b. Performance Gap, 1c. Evidence)

1a. Impact: H-2; M-1; L-2; I-0; 1b. Performance Gap: H-2; M-2; L-1; I-0

1c. Evidence Quantity: H-3; M-2; L-0; I-0; Quality: H-2; M-3; L-0-; I-0; Consistency: H-4; M-0; L-1; I-0

<u>Rationale</u>: The Committee agreed that pain should be assessed across the continuum of care but noted that implementing a pain intervention does not necessarily improve a patient's outcome. The measure did not look at the quality of the intervention and was even more limited due to the fact that it only included patients who reported pain at the time of the OASIS assessment and not every visit. Because the measure did not pass importance to measure and report, the Committee did not discuss the remaining criteria.

2. Scientific Acceptability of Measure Properties: Y-; N-

(2a. Reliability - precise specifications, testing; 2b. Validity - testing, threats to validity)

2a. Reliability: H-2; M-1; L-2; I-0; 2b. Validity: H-2; M-1; L-2; I-0

Rationale:

3. Usability: H-; M-; L-; I-

(*Meaningful, understandable, and useful to the intended audiences for 3a. Public Reporting/Accountability and 3b. Quality Improvement*) **3a.** Public Reporting: H-1; M-2; L-2; I-0

3b. QI: H-1; M-2; L-2; I-0

Rationale:

4. Feasibility: H-; M-; L-; I-

(4a. Clinical data generated during care delivery; 4b. Electronic sources; 4c.Susceptibility to inaccuracies/ unintended consequences identified 4d. Data collection strategy can be implemented)

4a. Byproduct of Care Processes: H-4; M-0; L-1; I-0

0524 Pain Interventions Implemented During Short Term Episodes Of Care

4b. Electronic data sources: H-3; M-1; L-1; I-0

4c. Suscep inaccuracies, consequences: H-2; M-2; L-1; I-0

4d. Data collection strategy: H-4; M-0; L-1; I-0

Rationale:

Steering Committee Recommendation for Endorsement: Did not pass the Importance to Measure criteria, which is required for endorsement.

Rationale: The evidence does not assess pain on every visit or consider the quality of the pain intervention.

MEASURES WITHDRAWN FROM CONSIDERATION

Four measures previously endorsed by NQF have not been re-submitted or withdrawn from maintenance of endorsement. Two of these measures were withdrawn after initial submission. The following measures are being retired from endorsement:

Measure	Reason for withdrawal
0019: Documentation of medication list in the	Developer elected not to pursue maintenance
outpatient record	of endorsement.
0020: Documentation of allergies and adverse	Developer elected not to pursue maintenance
reactions in the outpatient record	of endorsement.
0021: Annual monitoring for patients on	Withdrawn related to Steering Committee
persistent medications	discussion.
0503: Anticoagulation for acute pulmonary	This measure was moved to Phase II to provide
embolus	the developer additional time to complete
	testing.
1729: Polytherapy with Oral Antipsychotics	Withdrawn related to Steering Committee
	discussion.

NATIONAL VOLUNTARY CONSENSUS STANDARDS FOR PATIENT SAFETY: COMPLICATIONS ENDORSEMENT MAINTENANCE

APPENDIX A: MEASURE SPECIFICATIONS

The following tables present the detailed specifications for the National Quality Forum (NQF)endorsed® *National Voluntary Consensus Standards Patient Safety: Complications Endorsement Maintenance*. All information presented has been derived directly from measure sources/developers without modification or alteration (except when the measure developed agreed to such modification during the NQF Consensus Development Process) and is current as of February 17, 2012. All NQF-endorsed voluntary consensus standards are open source, meaning they are fully accessible and disclosed. Measures stewards include the Agency for Healthcare Research and Quality, Ambulatory Surgical Center Quality Collaboration, Centers for Medicare & Medicaid Services, National Committee for Quality Assurance, and The Joint Commission.

0022 Use of High Risk Medications in the Elderly
0263 Patient Burn
0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
0344 Accidental Puncture or Laceration Rate (PDI 1)A-5
0345 Accidental Puncture or Laceration Rate (PSI 15)
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0371 Venous Thromboembolism Prophylaxis
0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
0376 Incidence of Potentially Preventable Venous Thromboembolism
0419 Documentation of Current Medications in the Medical Record
0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12) A-71

	0022 Use of High Risk Medications in the Elderly
Steward	National Committee for Quality Assurance
-	 a: Percentage of Medicare members 65 years of age and older who received at least one high-risk medication. b: Percentage of Medicare members 65 years of age and older who received at least two different high-risk medications. For both rates, a lower rate represents better performance.
Туре	Process

	0022 Use of High Risk Medications in the Elderly
Data Source	Administrative claims, Electronic Clinical Data, Electronic Clinical Data : Pharmacy Healthcare Effectiveness Data and Information Set (HEDIS)
Level	Clinician : Group/Practice, Clinician : Individual, Health Plan, Integrated Delivery System
Setting	Ambulatory Care : Clinician Office, Pharmacy
Numerator Statement	a: At least one prescription dispensed for any high-risk medication during the measurement year. b: At least two prescriptions dispensed for different high-risk medications during the measurement year.
Numerator Details	Time Window: The measurement year.
	Antianviety (includes combination drugs) asplirin-meprobamate and meprobamate Analgesis: (includes combination drugs) acetaminophen-diphenhydramine, diphenhydramine-magnesium salicylate, and ketorolac Antihistamines (includes combination drugs) APAP/dxtromethorphan/diphenhydramine, APAP/diphenhydramine/phenylephrine, APAP/dixtromethorphan/diphenhydramine, acetaminophen-diphenhydramine/phenylephrine, APAP/dixtromethorphan/diphenhydramine, caretaminophen-diphenhydramine/phenylephrine, atorpine/CPM/poscyamine/EP/PA/scopolamine, caretaepane/diphenhydramine/phenylephrine , codeine/phenylephrine/promethazine, codeine-promethazine, cyproheptadine, dexchlorpheniramine/ghenylephrine, dexchlorpheniramine/dixtomethorphan/PSE, dexchlorpheniramine/guaifnesin/PSE, dexchlorpheniramine/ dexchlorpheniramine/maynesium salicylate, diphenhydramine, diphenhydramine/PSE, dexchlorpheniramine- pseudoephedrine, dextromethorphan-promethazine, diphenhydramine, diphenhydramine/Pydrocodone/phenylephrine, diphenhydramine-gseudoephedrine, acetaminopheniramine/guaifnesin/PSE, dexchlorpheniramine- mesordiazine and thioridazine Antipszychotic, typical mesordiazine and thioridazine Amphetamines amphetamine-dextroamphetamine, benzphetamine, dexmethylphenidate, dextroamphetamine, diethylpropion, methamphetamine, methylphenidate, pemoline, phendimetrazine, phentermine Barbitrates ambarbital, butabarbital, mephobarbital, pentobarbital, Phenobarbital, and secobarbital Long-acting benzodiazepoixide, chlordiazepoxide, chlordiazepoxide-clidinium, diazepam Calatum channel blockers nifedipine—short-acting only Gastrominestinal anti-spasmodics dicyclomine and propaniheline Beladonna alkaloids (includes combination drugs) atripine, dirophenzini, beladonna, beladonna/caffinel/ergotamine/PB/scopolamine , atropine-difenoxin, atropine- diphenoxylate, atropine-dirophonium, beladonna, caffinel/ergotamine/PB/scopolamine , atropine-difenoxin, atropine- diphenoxylate, atropine-dorophonium, beladonna, caffinel/ergotamine/PD/scopolamine , atropine-difenoxin, atropine-

	0022 Use of High Risk Medications in the Elderly
	Vasodilators cyclandelate, dipyridamole—short-acting only, ergot mesyloid, isoxsuprine Others (including androgens and anabolic steroids, thyroid drugs, urinary anti-infectives) methyltestosterone, nitrofurantoin, nitrofurantoin macrocrystals, nitrofurantoin macrocrystals-monohydrate, thyroid desiccated
Denominator Statement	All patients ages 65 years and older as of December 31 of the measurement year.
Denominator Details	Time Window: December 31 of the measurement year Use administrative data for eligible population
Exclusions	N/A
Exclusion Details	N/A
	No risk adjustment or risk stratification N/A
Stratification	N/A
Type Score	Rate/proportion better quality = lower score
Algorithm	 Step 1. Determine the eligible population. The eligible population is all members who satisfy all specified criteria, including any age, continuous enrollment, benefit, event, or anchor date enrollment requirement. Step 2. Search administrative systems to identify numerator events for all members in the eligible population. Step 3. If applicable, for members for whom administrative data do not show a positive numerator event, search administrative data for an exclusion to the service/procedure being measured. Note: This step applies only to measures for which optional exclusions are specified and for which the organization has chosen to search for exclusions. The organization is not required to search for optional exclusions. Step 4. Exclude from the eligible population members from step 3 for whom administrative system data identified an exclusion to the service/procedure being measured. Step 5. Calculate the rate.
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	0263 Patient Burn
Steward	Ambulatory Surgical Center Quality Collaboration
Description	Percentage of ASC admissions experiencing a burn prior to discharge
Туре	Outcome
Data Source	Paper Records ASC medical records, as well as incident/occurrence reports, and variance reports may serve as data sources. No specific collection instrument is required although the ASC Quality Collaboration has developed a sample data collection instrument that may be used as desired. Facilities may use any collection instrument that allows tracking of all burns prior to discharge. URL http://ascquality.org/documents/ASCQualityCollaborationImplementationGuide.pdf Not needed URL http://ascquality.org/documents/ASCQualityCollaborationImplementationGuide.pdf Not needed
Level	Facility
Setting	Ambulatory Care : Ambulatory Surgery Center (ASC)
Numerator Statement	Ambulatory surgical center (ASC) admissions experiencing a burn prior to discharge.
Numerator Details	Time Window: In-facility, prior to discharge DEFINITIONS: Admission: Completion of registration upon entry into the facility. Burn: Unintended tissue injury caused by any of the six recognized mechanisms: scalds, contact, fire, chemical, electrical, or

	0263 Patient Burn
	radiation (e.g. warming devices, prep solutions, electrosurgical unit, or laser). Discharge: Occurs when the patient leaves the confines of the ASC.
Denominator Statement	All ASC admissions.
Denominator Details	Time Window: In-facility, prior to discharge
	DEFNITIONS: Admission: Completion of registration upon entry into the facility.
Exclusions	None
Exclusion Details	No denominator exclusions
Risk Adjustment	No risk adjustment or risk stratification None.
Stratification	This measure is not stratified
Type Score	Rate/proportion better quality = lower score
Algorithm	The number of admissions experiencing a burn prior to discharge is divided by the number of ASC admissions during the reporting period, yielding the rate of burns prior to discharge for the reporting period.
Copyright	None

	0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
Steward	Ambulatory Surgical Center Quality Collaboration
Description	Percentage of ASC admissions experiencing a wrong site, wrong side, wrong patient, wrong procedure, or wrong implant event.
Туре	Outcome
Data Source	Paper Records ASC medical records, as well as incident/occurrence reports, and variance reports may serve as data sources. No specific collection instrument is required although the ASC Quality Collaboration has developed a sample data collection instrument that may be used as desired. Facilities may use any collection instrument that allows tracking of all wrong site, wrong side, wrong patient, wrong procedure, and wrong implant events. URL http://ascquality.org/documents/ASCQualityCollaborationImplementationGuide.pdf Not needed URL http://ascquality.org/documents/ASCQualityCollaborationImplementationGuide.pdf Not needed
Level	Facility
Setting	Ambulatory Care : Ambulatory Surgery Center (ASC)
Numerator Statement	ASC admissions experiencing a wrong site, wrong side, wrong patient, wrong procedure, or wrong implant
Numerator Details	Time Window: In-facility, prior to discharge DEFINITIONS: Admission: completion of registration upon entry into the facility Wrong: not in accordance with intended site, side, patient, procedure or implant
Denominator Statement	All ASC admissions
Denominator Details	Time Window: In-facility, prior to discharge DEFINITION: Admission: completion of registration upon entry into the facility
Exclusions	None
Exclusion Details	Not applicable

	0267 Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
	No risk adjustment or risk stratification
Adjustment	Not applicable
Stratification	The measure is not stratified
Type Score	Rate/proportion better quality = lower score
_	The number of admissions experiencing a wrong site, wrong side, wrong patient, wrong procedure, or wrong implant event is divided by the number of ASC admissions during the reporting period, yielding the rate of wrong site, wrong side, wrong patient, wrong procedure, and wrong implant events for the reporting period.
Copyright	None

	0344 Accidental Puncture or Laceration Rate (PDI 1)
Steward	Agency for Healthcare Research and Quality
Description	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
Туре	Outcome
	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Ũ	Hospital/Acute Care Facility
	Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
	Time Window: User may specify the time window; generally one calendar year ICD-9-CM Accidental puncture or laceration diagnosis codes: E8700 SURGICAL OPERATION E8701 INFUSION OR TRANSFUSION E8702 KIDNEY DIALYSIS OR OTHER PERFUSION E8703 INJECTION OR VACCINATION E8704 ENDOSCOPIC EXAMINATION E8705 ASPIRATION OF FLUID OR TISSUE, PUNCTURE, AND CATHETERIZATION E8706 HEART CATHETERIZATION E8707 ADMINISTRATION OF ENEMA
	E8708 OTHER SPECIFIED MEDICAL CARE E8709 UNSPECIFIED MEDICAL CARE 9982 ACCIDENTAL PUNCTURE OR LACERATION DURING A PROCEDURE
Denominator Statement	All surgical and medical discharges under age 18 defined by specific DRGs or MS-DRGs.

	0344 Accidental Puncture or Laceration Rate (PDI 1)
	Time Window: User may specify the time window; generally one calendar year
Details	
	See Pediatric Quality Indicators Appendices:
	- Appendix B – Surgical Discharge DRGs - Appendix C – Surgical Discharge MS-DRGs
	- Appendix C – Surgical Discharge DRGs
	- Appendix E – Medical Discharge MS-DRGs
	Link to PDI appendices:
	http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf
Exclusions	Exclude cases: - with principal diagnosis denoting accidental cut, puncture, perforation, or laceration or secondary diagnosis present on
	admission
	- normal newborn
	- neonate with birth weight less than 500 grams (Birth Weight Category 1)
	- MDC 14 (pregnancy, childbirth, and puerperium)
	- with ICD-9-CM code for spine surgery
	 with missing discharge gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)
	See Pediatric Quality Indicators Appendices:
	- Appendix I – Definitions of, Neonate, Newborn, Normal Newborn, and Outborn
	- Appendix L – Low Birth Weight Categories
Exclusion	ICD-9-CM Spine surgery procedure codes:
Details	0301 DEMOVAL OF FOREICN RODY FROM SPINAL CANAL
	REMOVAL OF FOREIGN BODY FROM SPINAL CANAL 0302
	REOPENING OF LAMINECTOMY SITE
	0309
	OTHER EXPLORATION AND DECOMPRESSION OF SPINAL CANAL
	0353 REPAIR OF VERTEBRAL FRACTURE
	036
	LYSIS OF ADHESIONS OF SPINAL CORD AND NERVE ROOTS
	8053
	REPAIR OF THE ANULUS FIBROSUS WITH GRAFT OR PROSTHESIS (OCT08) 8054
	OTHER AND UNSPECIFIED REPAIR OF THE ANULUS FIBROSUS (OCT08)
	8100
	SPINAL FUSION, NOT OTHERWISE SPECIFIED
	ATLAS-AXIS SPINAL FUSION 8102
	OTHER CERVICAL FUSION, ANTERIOR TECHNIQUE
	8103
	OTHER CERVICAL FUSION, POSTERIOR TECHNIQUE
	DORSAL AND DORSOLUMBAR FUSION, ANTERIOR TECHNIQUE 8105
	DORSAL AND DORSOLUMBAR FUSION, POSTERIOR TECHNIQUE
	8106
	LUMBAR AND LUMBOSACRAL FUSION, ANTERIOR TECHNIQUE
	8107
	LUMBAR AND LUMBOSACRAL FUSION, LATERAL TRANSVERSE PROCESS TECHNIQUE
	8108

	0344 Accidental Puncture or Laceration Rate (PDI 1)
	LUMBAR AND LUMBOSACRAL FUSION, POSTERIOR TECHNIQUE
	8130
	REFUSION OF SPINE, NOT OTHERWISE SPECIFIED
	8131 REFUSION OF ATLAS-AXIS SPINE
	8132
	REFUSION OF OTHER CERVICAL SPINE, ANTERIOR TECHNIQUE
	8133
	REFUSION OF OTHER CERVICAL SPINE, POSTERIOR TECHNIQUE 8134
	REFUSION OF DORSAL AND DORSOLUMBAR SPINE, ANTERIOR TECHNIQUE
	8135
	REFUSION OF DORSAL AND DORSOLUMBAR SPINE, POSTERIOR TECHNIQUE
	8136 REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, ANTERIOR TECHNIQUE
	8137
	REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, LATERAL TRANSVERSE PROCESS TECHNIQUE
	8138 REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, POSTERIOR TECHNIQUE
	8139
	REFUSION OF SPINE, NOT ELSEWHERE CLASSIFIED
	FUSION OR REFUSION OF 2-3 VERTEBRAE* 8163
	FUSION OR REFUSION OF 4-8 VERTEBRAE*
	8164
	FUSION OR REFUSION OF 9 OR MORE VERTEBRAE* 8165
	VERTEBROPLASTY
	8166
	KYPHOPLASTY
	8451 INSERTION OF INTERBODY SPINAL FUSION DEVICE*
	8452
	INSERTION OF RECOMBINANT BONE MORPHOGENETIC PROTEIN*
	IMPLANTATION OF INTERSPINOUS PROCESS DECOMPRESSION DEVICE (PRIOR TO OCT 1, 2007) 8459
	INSERTION OF OTHER SPINAL DEVICES
	INSERTION OF SPINAL DISC PROSTHESIS, NOT OTHERWISE SPECIFIED 8461
	INSERTION OF PARTIAL SPINAL DISC PROSTHESIS, CERVICAL
	8462
	INSERTION OF TOTAL SPINAL DISC PROSTHESIS, CERVICAL
	8463 INSERTION OF SPINAL DISC PROSTHESIS, THORACIC
	8464
	INSERTION OF PARTIAL SPINAL DISC PROSTHESIS, LUMBOSACRAL
	8465 INSERTION OF TOTAL SPINAL DISC PROSTHESIS, LUMBOSACRAL
	8466
	REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, CERVICAL
-	

	0344 Accidental Puncture or Laceration Rate (PDI 1)
	8467 REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, THORACIC
	8468
	REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, LUMBOSACRAL
	8469 REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, NOT OTHERWISE SPECIFIED
	INSERTION OR REPLACEMENT OF INTERSPINOUS PROCESS DEVICE(S) 8481
	REVISION OF INTERSPINOUS PROCESS DEVICE(S)
	8482 INSERTION OR REPLACEMENT OF PEDICLE-BASED DYNAMIC STABILIZATION DEVICE(S)
	8483
	REVISION OF PEDICLE-BASED DYNAMIC STABILIZATION DEVICE(S) 8485
	REVISION OF FACET REPLACEMENT DEVICE(S)
	* code has "code also" instructions
	See Pediatric Quality Indicators Appendices: - Appendix I – Definitions of, Neonate, Newborn, Normal Newborn, and Outborn
	- Appendix L – Low Birth Weight Categories
	Link to PDI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf
Risk	Statistical risk model
Adjustment	The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) an
	covariates for gender, birthweight (500g groups), age in days (29-60, 61-90, 91+), age in years (in 5-year age groups),
	modified CMS DRG and AHRQ CCS comorbities. The reference population used in the regression is the universe of
	discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 43 states and approximately 6 million pediatric discharges. The expected rate is computed as the sum of the predicted value for
	each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed
	using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate.
	Covariates used in this measures:
	MDC 5
	MDC 6
	MDC 8
	MDC 11 MDC 15
	MDC 0THER
	Procedure Type 2
	Procedure Type 3
	Procedure Type 4 to 5
	Procedure Type 6
	Procedure Type 6 Procedure Type 7
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type)
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure 4. One major therapeutic with only minor diagnostic procedure(s)
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure 4. One major therapeutic with only minor diagnostic procedure(s) 5. One major therapeutic with major diagnostic procedure(s) 6. Two major therapeutic procedures with any or no diagnostic procedures
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure 4. One major therapeutic with only minor diagnostic procedure(s) 5. One major therapeutic with major diagnostic procedure(s) 6. Two major therapeutic procedures with any or no diagnostic procedures 7. Three or more major therapeutic procedures with any or no diagnostic procedures;
	Procedure Type 6 Procedure Type 7 *** Risk adjust by risk category (Procedure Type) 1. No therapeutic procedure with any or no diagnostic procedures 2. Only minor therapeutic procedure with any or no diagnostic procedures 3. One major therapeutic without diagnostic procedure 4. One major therapeutic with only minor diagnostic procedure(s) 5. One major therapeutic with major diagnostic procedure(s) 6. Two major therapeutic procedures with any or no diagnostic procedures

	0344 Accidental Puncture or Laceration Rate (PDI 1)
	Stratum 1. Eye, ear, nose, mouth, throat, skin, breast, and other low-risk procedures (MDC 2, 3, 9, 19, 22, 23)
	Stratum 2. Thoracic, cardiovascular, and specified neoplastic procedures
	(MDC 4, 5, 17) Stratum 3. Kidney, and male/female reproductive procedures
	MDC 11, 12, 13)
	Stratum 4. Infectious, immunological, hematological, and ungroupable procedures (MDC 0/99, 16, 18, 25)
	Stratum 5. Trauma, orthopedic, and neurologic procedures (MDC 1, 8, 21, 24)
	Stratum 6. Gastrointestinal, hepatobiliary, and endocrine procedures (MDC 6, 7, 10)
Type Score	Rate/proportion better quality = lower score
Algorithm	Each indicator is expressed as a rate, is defined as outcome of interest / population at risk or numerator / denominator. The AHRQ Quality Indicators (AHRQ QI) software performs six steps to produce the rates. 1) Discharge-level data is used to mark inpatient records containing the outcome of interest and 2) the population at risk. For provider indicators, the population at risk is also derived from hospital discharge records; for area indicators, the population at risk is derived from U.S. Census data. 3) Calculate observed rates. Using output from steps 1 and 2, rates are calculated for user-specified combinations of stratifiers. 4) Calculate expected rates. Regression coefficients from a reference population database are applied to the discharge records and aggregated to the provider or area level. For indicators that are not risk-adjusted, this is the reference population rate. 5) Calculate risk-adjusted rate. Use the indirect standardization to account for case-mix. For indicators that are not risk-adjusted, this is the same as the observed rate. 6) Calculate smoothed rate. A Univariate shrinkage factor is applied to the risk-adjusted rates. The shrinkage estimate reflects a reliability adjustment unique to each indicator URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable

	0345 Accidental Puncture or Laceration Rate (PSI 15)
Steward	Agency for Healthcare Research and Quality
Description	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary diagnosis field.
Numerator Details	Time Window: User may specify the time window; generally one calendar year ICD-9-CM Accidental puncture or laceration diagnosis codes: E8700 SURGICAL OPERATION E8701 INFUSION OR TRANSFUSION E8702 KIDNEY DIALYSIS OR OTHER PERFUSION E8703

	0345 Accidental Puncture or Laceration Rate (PSI 15)
	INJECTION OR VACCINATION
	E8704
	ENDOSCOPIC EXAMINATION
	E8705 ASPIRATION OF FLUID OR TISSUE, PUNCTURE, AND CATHETERIZATION
	E8706
	HEART CATHETERIZATION
	E8707
	ADMINISTRATION OF ENEMA E8708
	OTHER SPECIFIED MEDICAL CARE E8709
	UNSPECIFIED MEDICAL CARE
	9982
	ACCIDENTAL PUNCTURE OR LACERATION DURING A PROCEDURE
Denominator Statement	All surgical and medical discharges age 18 years and older defined by specific DRGs or MS-DRGs.
	Time Window: User may specify the time window; generally one calendar year
Dotans	See Patient Safety Indicators Appendices:
	- Appendix B – Medical Discharge DRGs
	- Appendix C – Medical Discharge MS-DRGs
	- Appendix D – Surgical Discharge DRGs
	- Appendix E – Surgical Discharge MS-DRGs Link to PSI appendices:
	http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf
Exclusions	Exclude cases:
	- with principal diagnosis denoting accidental cut, puncture, perforation, or laceration or secondary diagnosis present on
	admission
	- MDC 14 (pregnancy, childbirth, and puerperium) - with ICD-9-CM code for spine surgery
	- with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal
	diagnosis (DX1=missing)
Exclusion	ICD-9-CM Spine surgery procedure codes:
Details	
	REMOVAL OF FOREIGN BODY FROM SPINAL CANAL 0302
	REOPENING OF LAMINECTOMY SITE
	0309
	OTHER EXPLORATION AND DECOMPRESSION OF SPINAL CANAL
	0353 REPAIR OF VERTEBRAL FRACTURE
	036
	LYSIS OF ADHESIONS OF SPINAL CORD AND NERVE ROOTS
	8053
	REPAIR OF THE ANULUS FIBROSUS WITH GRAFT OR PROSTHESIS (OCT08) 8054
	OTHER AND UNSPECIFIED REPAIR OF THE ANULUS FIBROSUS (OCT08) 8100
	SPINAL FUSION, NOT OTHERWISE SPECIFIED
	8101 ATLAS-AXIS SPINAL FUSION
L	l

0345 Accidental Puncture or Laceration Rate (PSI 15)
8102
OTHER CERVICAL FUSION, ANTERIOR TECHNIQUE 8103
OTHER CERVICAL FUSION, POSTERIOR TECHNIQUE
8104
DORSAL AND DORSOLUMBAR FUSION, ANTERIOR TECHNIQUE
8105 DORSAL AND DORSOLUMBAR FUSION, POSTERIOR TECHNIQUE
8106
LUMBAR AND LUMBOSACRAL FUSION, ANTERIOR TECHNIQUE 8107
LUMBAR AND LUMBOSACRAL FUSION, LATERAL TRANSVERSE PROCESS TECHNIQUE
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LUMBAR AND LUMBOSACRAL FUSION, POSTERIOR TECHNIQUE 8130
REFUSION OF SPINE, NOT OTHERWISE SPECIFIED
8131
REFUSION OF ATLAS-AXIS SPINE 8132
REFUSION OF OTHER CERVICAL SPINE, ANTERIOR TECHNIQUE
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REFUSION OF OTHER CERVICAL SPINE, POSTERIOR TECHNIQUE 8134
REFUSION OF DORSAL AND DORSOLUMBAR SPINE, ANTERIOR TECHNIQUE
8135
REFUSION OF DORSAL AND DORSOLUMBAR SPINE, POSTERIOR TECHNIQUE 8136
REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, ANTERIOR TECHNIQUE
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REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, LATERAL TRANSVERSE PROCESS TECHNIQUE 8138
REFUSION OF LUMBAR AND LUMBOSACRAL SPINE, POSTERIOR TECHNIQUE
8139
REFUSION OF SPINE, NOT ELSEWHERE CLASSIFIED 8162
FUSION OR REFUSION OF 2-3 VERTEBRAE*
8163
FUSION OR REFUSION OF 4-8 VERTEBRAE* 8164
FUSION OR REFUSION OF 9 OR MORE VERTEBRAE*
8165
VERTEBROPLASTY 8166
KYPHOPLASTY
8451
INSERTION OF INTERBODY SPINAL FUSION DEVICE* 8452
INSERTION OF RECOMBINANT BONE MORPHOGENETIC PROTEIN*
8458
IMPLANTATION OF INTERSPINOUS PROCESS DECOMPRESSION DEVICE (ONLY BEFORE OCT 1, 2007) 8459
INSERTION OF OTHER SPINAL DEVICES
8460

	0345 Accidental Puncture or Laceration Rate (PSI 15)
	INSERTION OF SPINAL DISC PROSTHESIS, NOT OTHERWISE SPECIFIED
	8461 INSERTION OF PARTIAL SPINAL DISC PROSTHESIS, CERVICAL
	8462
	INSERTION OF TOTAL SPINAL DISC PROSTHESIS, CERVICAL
	8463 INSERTION OF SPINAL DISC PROSTHESIS, THORACIC
	8464
	INSERTION OF PARTIAL SPINAL DISC PROSTHESIS, LUMBOSACRAL 8465
	INSERTION OF TOTAL SPINAL DISC PROSTHESIS, LUMBOSACRAL
	8466
	REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, CERVICAL 8467
	REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, THORACIC
	8468 REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, LUMBOSACRAL
	8469 REVISION OR REPLACEMENT OF ARTIFICIAL SPINAL DISC PROSTHESIS, NOT OTHERWISE SPECIFIED
	8480 INSERTION OR REPLACEMENT OF INTERSPINOUS PROCESS DEVICE(S)
	REVISION OF INTERSPINOUS PROCESS DEVICE(S) 8482
	INSERTION OR REPLACEMENT OF PEDICLE-BASED DYNAMIC STABILIZATION DEVICE(S)
	8483 REVISION OF PEDICLE-BASED DYNAMIC STABILIZATION DEVICE(S)
	8485
	REVISION OF FACET REPLACEMENT DEVICE(S)
Risk Adjustment	Statistical risk model The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, age, modified CMS DRG, transfer status, procedure day availability, and the AHRQ Comorbidity category. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Covariates used in this measures: Sex Female Age 18 to 24 Age 25 to 29 Age 30 to 59 MDRG 101 MDRG 103 MDRG 107 MDRG 302 MDRG 401 MDRG 401 MDRG 401 MDRG 402 MDRG 416 MDRG 502

0345 A	ccidental Puncture or Laceration Rate (PSI 15)
MDRG	506
MDRG	507
MDRG	508
MDRG	510
MDRG	511
MDRG	513
MDRG	514
MDRG	519
MDRG	520
MDRG	522
MDRG	601
MDRG	602
MDRG	603
MDRG	604
MDRG	606
MDRG	609
MDRG	610
MDRG	611
MDRG	621
MDRG	701
MDRG	702
MDRG	703
MDRG	704
MDRG	705
MDRG	712
MDRG	806
MDRG	807
MDRG	815
MDRG	816
MDRG	1001
MDRG	1003
MDRG	1005
MDRG	1006
MDRG	1101
MDRG	1102
MDRG	1103
MDRG	1104
MDRG	1105
MDRG	1107
MDRG	1109
MDRG	1201
MDRG	1204
MDRG	1301
MDRG	1302
MDRG	1303
MDRG	1304
MDRG	1305
MDRG	1306
MDRG	1307
MDRG	1308
MDRG	1707
MDRG	1709
MDRG	1801

	0345 Accidental Puncture or Laceration Rate (PSI 15)
	MDRG 1802
	MDRG 2104
	MDRG 2108
	MDRG 2408
	MDRG 7702
	MDC 3
	MDC 4
	MDC 5
	MDC 6 MDC 7
	MDC 7 MDC 8
	MDC 9
	MDC 9 MDC 11
	MDC 12
	MDC 13
	MDC 16
	MDC 17
	MDC 18
	MDC 21
	MDC 24
	MDC Other
	TRNSFER Transfer-in
	NOPRDAY Procedure Days Data Not Available
	COMORB PERIVASC
	COMORB DM
	COMORB DMCX
	COMORB RENLFAIL
	COMORB OBESE
	COMORB WGHTLOSS
	COMORB BLDLOSS COMORB ANEMDEF
	URL http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/Risk%20Adjustment%20Tables%20PSI%204.3.pdf Not
	applicable
	Not applicable Rate/proportion better quality = lower score
51	Each indicator is expressed as a rate, is defined as outcome of interest / population at risk or numerator / denominator. The
Algorithm	AHRQ Quality Indicators (AHRQ QI) software performs six steps to produce the rates. 1) Discharge-level data is used to mark
	inpatient records containing the outcome of interest and 2) the population at risk. For provider indicators, the population at risk
	is also derived from hospital discharge records; for area indicators, the population at risk is derived from U.S. Census data. 3)
	Calculate observed rates. Using output from steps 1 and 2, rates are calculated for user-specified combinations of stratifiers.
	4) Calculate expected rates. Regression coefficients from a reference population database are applied to the discharge
	records and aggregated to the provider or area level. For indicators that are not risk-adjusted, this is the reference population
	rate. 5) Calculate risk-adjusted rate. Use the indirect standardization to account for case-mix. For indicators that are not risk-
	adjusted, this is the same as the observed rate. 6) Calculate smoothed rate. A Univariate shrinkage factor is applied to the
	risk-adjusted rates. The shrinkage estimate reflects a reliability adjustment unique to each indicator URL Not applicable
	http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
	Not applicable
15 5	

	0346 latrogenic Pneumothorax Rate (PSI 6)
Steward	Agency for Healthcare Research and Quality
Description	Percent of discharges with ICD-9-CM code for iatrogenic pneumothorax in any secondary diagnosis field among cases

	0346 latrogenic Pneumothorax Rate (PSI 6)
	meeting the inclusion and exclusion rules for the denominator
Туре	Outcome
-	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges with ICD-9-CM code for iatrogenic pneumothorax in any secondary diagnosis field among cases meeting the inclusion and exclusion rules for the denominator.
Numerator Details	Time Window: User may specify the time window; generally one calendar year
	ICD-9-CM latrogenic Pneumothorax diagnosis code: 5121 IATROGENIC PNEUMOTHORAX
Denominator Statement	All surgical and medical discharges age 18 years and older defined by specific DRGs or MS-DRGs.
Denominator Details	Time Window: User may specify the time window; generally one calendar year
	See Patient Safety Indicators Appendices: - Appendix B – Medical Discharge DRGs - Appendix C – Medical Discharge MS-DRGs - Appendix D – Surgical Discharge DRGs - Appendix E – Surgical Discharge MS-DRGs Link to PSI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf
Exclusions	Exclude cases: - with principal diagnosis of iatrogenic pneumothorax or secondary diagnosis present on admission - MDC 14 (pregnancy, childbirth, and puerperium) - with any diagnosis code of chest trauma or pleural effusion - with a code of diaphragmatic surgery repair in any procedure field - with any code indicating thoracic procedure, lung or pleural biopsy, or cardiac procedure - with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)
Details	ICD-9-CM Chest trauma diagnosis codes: 80700 FRACTURE RIB NOS-CLOSED 80701 FRACTURE ONE RIB-CLOSED 80702 FRACTURE TWO RIBS-CLOSED 80703 FRACTURE THREE RIBS-CLOS 80704 FRACTURE FOUR RIBS-CLOSE 80705 FRACTURE FIVE RIBS-CLOSE 80706 FRACTURE SIX RIBS-CLOSED 80707

	0346 latrogenic Pneumothorax Rate (PSI 6)
	FRACTURE SEVEN RIBS-CLOS
	80708
	FX EIGHT/MORE RIB-CLOSED
	80709
	FX MULT RIBS NOS-CLOSED
	80710
	FRACTURE RIB NOS-OPEN
	FRACTURE ONE RIB-OPEN 80712
	FRACTURE TWO RIBS-OPEN
	80713
	FRACTURE THREE RIBS-OPEN
	80714
	FRACTURE FOUR RIBS-OPEN
	80715
	FRACTURE FIVE RIBS-OPEN
	80716
	FRACTURE SIX RIBS-OPEN
	80717
	FRACTURE SEVEN RIBS-OPEN
	FX EIGHT/MORE RIBS-OPEN 80719
	FX MULT RIBS NOS-OPEN
	8072
	FRACTURE OF STERNUM-CLOS
	8073
	FRACTURE OF STERNUM-OPEN
	8074
	FLAIL CHEST
	8075
	FX LARYNX/TRACHEA-CLOSED
	8076
	FX LARYNX/TRACHEA-OPEN
	8090 FRACTURE TRUNK BONE-CLOS
	8091
	FRACTURE TRUNK BONE-OPEN
	8600
	TRAUM PNEUMOTHORAX-CLOSE
	8601
	TRAUM PNEUMOTHORAX-OPEN
	8602
	TRAUM HEMOTHORAX-CLOSED
	8603
	TRAUM HEMOTHORAX-OPEN
	8604
	TRAUM PNEUMOHEMOTHOR-CL 8605
	TRAUM PNEUMOHEMOTHOR-OPN
	86100
	HEART INJURY NOS-CLOSED
L	

0346 latrogenic Pneumothorax Rate (PSI 6)
86101
HEART CONTUSION-CLOSED
86102
HEART LACERATION-CLOSED
86103
HEART CHAMBER LACERAT-CL
86110
HEART INJURY NOS-OPEN
86111
HEART CONTUSION-OPEN
86112
HEART LACERATION-OPEN
86113
HEART CHAMBER LACER-OPN
86120
LUNG INJURY NOS-CLOSED
86121
LUNG CONTUSION-CLOSED
86122
LUNG LACERATION-CLOSED
86130
LUNG INJURY NOS-OPEN
86131
LUNG CONTUSION-OPEN
86132
LUNG LACERATION-OPEN
8620
DIAPHRAGM INJURY-CLOSED
8621
DIAPHRAGM INJURY-OPEN
86221
BRONCHUS INJURY-CLOSED
ESOPHAGUS INJURY-CLOSED
INTRATHORACIC INJ NEC-CL
86231 BRONCHUS INJURY-OPEN
86232
ESOPHAGUS INJURY-OPEN
86239
INTRATHORAC INJ NEC-OPEN
8628
INTRATHORACIC INJ NOS-CL
8629
INTRATHORAC INJ NOS-OPEN
8750
OPEN WOUND OF CHEST
8751
OPEN WOUND CHEST-COMPL
8760
OPEN WOUND OF BACK
8761

0346 latrogenic Pneumothorax Rate (PSI 6)
OPEN WOUND BACK-COMPL
9010
INJURY THORACIC AORTA
9011
INJ INNOMIN/SUBCLAV ART
9012
INJ SUPERIOR VENA CAVA
9013
INJ INNOMIN/SUBCLAV VEIN
90140
INJ PULMONARY VESSEL NOS
INJURY PULMONARY ARTERY 90142
INJURY PULMONARY VEIN
90181
INJ INTERCOSTAL ART/VEIN
90182
INJ INT MAMMARY ART/VEIN
90183
INJ MULT THORACIC VESSEL
90189
INJ THORACIC VESSEL NEC
9019
INJ THORACIC VESSEL NOS
9110
ABRASION TRUNK
9111
ABRASION TRUNK-INFECTED
9118
SUPERFIC INJU TRUNK NEC
SUPERFIC INJU TRUNK NEC-INF
9220 CONTUSION OF BREAST
9221
CONTUSION OF CHEST WALL
9223
BACK CONTUSION
92231
BACK CONTUSION
92233
INTERSCPLR REG CONTUSION
9228
MULIPLE CONTUSION TRUNK
9229
CONTUSION OF TRUNK
92611
CRUSHING INJURY BACK
CRUSHING INJ TRUNK NEC
9268 MULT CRUSHING INJ TRUNK

0346 latrogenic Pneumothorax Rate (PSI 6)
9269
CRUSHING INJ TRUNK NOS
9290 CRUSH INJ MULT SITE NEC
9299
CRUSHING INJURY NOS
9541
INJ SYMPA NERVE NEC 9548
INJURY TRUNK NERVE NEC
9549
INJURY TRUNK NERVE NOS
95911
INJURY OF CHEST WALL NEC 95919
TRUNK INJURY-SITES NEC
9599
INJURY-SITE NOS
ICD-9-CM Pleural effusion diagnosis codes: 0101
TUBERCULOUS PLEURISY IN PRIMARY PROGRESSIVE TUBERCULOSIS
01010
TUBERCULOUS PLEURISY IN PRIMARY PROGRESSIVE TUBERCULOSIS, UNSPECIFIED
01011
TPIPPT, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE 01012
TPIPPT, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN
01013
TPIPPT, TUBERCLE BACILI FOUND BY MICROSCOPY
01014 TPIPPT, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE
01015
TPIPPT, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY
01016
TPIPPT, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER
METHODS 0117
TUBRCULOUS PNEUMOTHORAX
01170
TUBRCULOUS PNEUMOTHORAX, UNSPECIFIED
01171 TRNELL RACTERIAL OR LUSTOLOGICAL EVANANOT RONE
TPNEU, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE 01172
TPNEU, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN
01173
TPNEU, TUBERCLE BACILI FOUND BY MICROSCOPY
01174 TPNEU, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE
01175
TPNEU, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY
01176
TPENU, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER
METHODS

0346 latrogenic Pneumothorax Rate (PSI 6)
0120
TUBERCULOUS PLEURISY
TUBERCULOUS PLEURISY, UNSPECIFIED 01201
TP, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE
01202
TP, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN
TP, TUBERCLE BACILI FOUND BY MICROSCOPY 01204
TP, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE
01205
TP, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY
01206
TP, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER METHODS
1972 SECOND MALIG NEO PLEURA
ICD9-CM Diaphragmatic surgery repair codes:
537
ABD REPAIR-DIAPHR HERNIA
LAPAROSCOPIC REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH (OCT08) 5372
OTHER AND OPEN REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH (OCT08)
5375
REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH, NOS (OCT08)
THOR REP-DIAPH HERN NOS 5381
DIAPHRAGMATIC PLICATION
5382
PARASTERN HERNIA REPAIR
5583
LAPAROSCOPIC REPAIR OF DIAPHRAGMATIC HERNIA, WITH THORACIC APPROACH (OCT08) 5584
OTHER AND OPEN REPAIR OF DIAPHRAGMATIC HERNIA, WITH THORACIC APPROACH (OCT08)
ICD-9-CM Thoracic surgery procedure codes:
0522
SYMPATHECTOMY CERVICAL
0523 SYMPATHECTOMY LUMBAR
0529
OTHER SYMPATHECTOMY AND GANGLIONECTOMY
THYMECTOMY, NOT OTHERWISE SPECIFIED
0781 OTHER PARTIAL EXCISION OF THYMUS
0782
OTHER TOTAL EXCISION OF THYMUS
THORACOSCOPIC PARTIAL EXCISION OF THYMUS 0784
0704

0346 latrogenic Pneumothorax Rate (PSI 6)
THORACOSCOPIC TOTAL EXCISION OF THYMUS
3121
MEDIASTINAL TRACHEOSTOMY
3145 OPEN BIOPSY OF LARYNX OR TRACHEA
3173
CLOSURE OF OTHER FISTULA OF TRACHEA
3179
OTHER REPAIR AND PLASTIC OPERATIONS ON TRACHEA 3199
OTHER OPERATIONS ON TRACHEA
3209
OTHER LOCAL EXCISION OR DESTRUCTION OF LESION OR TISSUE OF BRONCHUS
321 OTHER EXCISION OF BRONCHUS
3220
THORAC EXC LUNG LESION
Local excision or destruction of lesion or tissue of lung
3221
PLICATION OF EMPHYSEMATIOUS BLEB
3222
LUNG VOLUME REDUCTION SURGERY 3223
OPEN ABLTN LUNG LES/TISS (OCT06)
3224
PERC ABLTN LUNG LES/TISS (OCT06)
THOR ABLTN LUNG LES/TISS (OCT06) 3226
ABLTN LUNG TISS NEC/NOS (OCT06)
3227
BRNC THRMPLSTY, ABLT MSCL
3228
ENDOSCOPIC EXCISION OR DESTRUCTION OF LESION OR TISSUE OF LUNG
3229 OTHER LOCAL EXCISION OR DESTRUCTION OF LESION OR TISSUE OF LUNC
OTHER LOCAL EXCISION OR DESTRUCTION OF LESION OR TISSUE OF LUNG 323
SEGMENTAL RESECTION OF LUNG
3230
THORAC SEG LUNG RESECT
3239
OTH SEG LUNG RESECT NOS
324
LOBECTOMY OF LUNG
3241 THORAC LOBECTOMY LUNG
3249
OTHER LOBECTOMY OF LUNG
325
COMPLETE PNEUMONECTOMY
3250
THORACOSPC PNEUMONECTOMY
3259

0346 latrogenic Pneumothorax Rate (PSI 6)
OTHER PNEUMONECTOMY NOS
326
RADICAL DISSECTION OF THORACIC STRUCTURES
329
OTHER EXCISION OF LUNG 330
INCISION OF BRONCHUS
331
INCISION OF LUNG
3320
THORACOSCOPC LUNG BIOPSY
3325
OPEN BIOPSY OF BRONCHUS
CLOSED ENDOSCOPIC BIOPSY OF LUNG 3331
DESTRUCTION OF PHRENIC NERVE FOR COLLAPSE OF LUNG (NO LONGER PERFORMED)
3332
ARTIFICIAL PNEUMOTHORAX FOR COLLAPSE OF LUNG
3334
THORACOPLASTY
3339
OTHER SURGICAL COLLAPSE OF LUNG
Repair and plastic operation on lung and bronchus
3341 SUTURE OF LACERATION OF BRONCHUS
3342
CLOSURE OF BRONCHIAL FISTULA
3343
CLOSURE OF LACERATION OF LUNG
3348
OTHER REPAIR AND PLASTIC OPERATIONS ON BRONCHUS
3349
OTHER REPAIR AND PLASTIC OPERATIONS ON LUNG Lung transplant
335
LUNG TRANSPLANTATION
3350
LUNG TRANSPLANTATION, NOS
3351
UNILATERAL LUNG TRANSPLANTATION
3352 BILATERAL LUNG TRANSPLANTATION
336
COMBINED HEART-LUNG TRANSPLANTATION
3392
LIGATION OF BRONCHUS
3393
PUNCTURE OF LUNG
3398
OTHER OPERATIONS ON BRONCHUS 3399
OTHER OPERATIONS ON LUNG

0346 latrogenic Pneumothorax Rate (PSI 6)
3329 OTHER DIAGNOSTIC PROCEDURE ON LUNG AND BRONCHUS
3333
PNEUMOPERITONEUM FOR COLLAPSE OF LUNG
3401
INCISION OF CHEST WALL 3402
EXPLORATORY THORACOTOMY
3403
REOPENING OF RECENT THORACOTOMY SITE
3405 CREATION OF PLEUROPERITONEAL SHUNT
3409
OTHER INCISION OF PLEURA
341
INCISION OF MEDIASTINUM
Diagnostic procedures on chest wall, pleura, mediastinum, and diaphragm 3420
THORACOSCOPIC PLEURAL BX
3421
TRANSPLEURAL THORACOSOCOPY
3422 MEDIASTINOSCOPY
3423
BIOPSY OF CHEST WALL
3425
CLOSED [PERCUTANEOUS][NEEDLE] BIOPSY OF MEDIASTINUM
3426 OPEN BIOPSY OF MEDIASTINUM
3427
BIOPSY OF DIAPHRAGM
3428
OTHER DIAGNOSTIC PROCEDURES ON CHEST WALL, PLEURA, AND DIAPHRAGM 3429
OTHER DIAGNOSTIC PROCEDURES ON MEDIASTINUM
343
EXCISION OR DESTRUCTION OF LESION OR TISSUE OF MEDIASTINUM
344 EXCISION OR DESTRUCTION OF LESION OF CHEST WALL
3451
DECORTICATION OF LUNG
3452
THORACOSCOPC DECORT LUNG
3459 OTHER EXCISION OF PLEURA
Repair of chest wall
3471
SUTURE OF LACERATION OF CHEST WALL
3472 CLOSURE OF THORACOSTOMY
3473
CLOSURE OF OTHER FISTULA OF THORAX
3474

0346 latrogenic Pneumothorax Rate (PSI 6)
REPAIR OF PECTUS DEFORMITY
3479
OTHER REPAIR OF CHEST WALL
Operations on diaphragm 3481
EXCISION OF LESION OR TISSUE OF DIAPHRAGM
3482
SUTURE OF LACERATION OF DIAPHRAGM
3483
CLOSURE OF FISTULA OF DIAPHRAGM
3484
OTHER REPAIR OF DIAPHRAGM
3485 IMPLANTATION OF DIAPHRAGMATIC PACEMAKER
3489
OTHER OPERATIONS ON DIAPHRAGM
3493
REPAIR OF PLEURA
3499
OTHER OPERATIONS ON THORAX, OTHER
Operations on thoracic duct 4061
CANNULATION OF THORACIC DUCT
4062
FISTULIZATION OF THORACIC DUCT
4063
CLOSURE OF FISTULA OF THORACIC DUCT
LIGATION OF THORACIC DUCT 4069
OTHER OPERATIONS ON THORACIC DUCT
Esophagotomy
4201
INCISION OF ESOPHAGEAL WEB
4209
OTHER INCISION OF ESOPHAGUS
4210 ESOPHAGOSTOMY, NOS
4211
CERVICAL ESOPHAGOSTOMY
4212
EXTERIORIZATION OF ESOPHAGEAL POUCH
OTHER EXTERNAL FISTULIZATION OF ESOPHAGUS 4221
OPERATIVE ESOPHAGOSCOPY BY INCISION
4225
OPEN BIOPSY OF ESOPHAGUS
4231
LOCAL EXCISION OF ESOPHAGEAL DIVERTICULUM
LOCAL EXCISION OF OTHER LESION OR TISSUE OF ESOPHAGUS Excision of esophagus

	0346 latrogenic Pneumothorax Rate (PSI 6)
	4239
	OTHER DESTRUCTION OF LESION OR TISSUE OF ESOPHAGUS 4240
	ESOPHAGECTOMY, NOS
	4241
	PARTIAL ESOPHAGECTOMY
	4242
	TOTAL ESOPHAGECTOMY Intrathoracic anastomosis of exophagus
	4251
	INTRATHORACIC ESOPHAGOESOPHAGOSTOMY
	4252
	INTRATHORACIC ESOPHAGOGASTROSTOMY
	4253 Initrationacic esopulaceal anastomosis w/interprosition of small power
	INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF SMALL BOWEL 4254
	OTHER INTRATHORACIC ESOPHAGOENTEROSTOMY
	4255
	INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF COLON
	OTHER INTRATHORACIC ESOPHAGOCOLOSTOMY 4258
	INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ OTHER INTERPOSITION
	4259
	OTHER INTRATHORACIC ANASTOMOSIS OF ESOPHAGUS
	Antesternal anastomosis
	4261 ANTESTERNAL ESOPHAGOESOPHAGOSTOMY
	4262
	ANTESTERNAL ESOPHAGOGASTROSTOMY
	4263
	ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF SMALL BOWEL 4264
	OTHER ANTESTERNAL ESOPHAGOENTEROSTOMY
	4265
	ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF COLON
	4266
	OTHER ANTESTERNAL ESOPHAGOCOLOSTOMY 4268
	0THER ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION
	4269
	OTHER ANTESTERNAL ANASTOMOSIS OF ESOPHAGUS
	Other repair of esophagus
	ESOPHAGOMYOTOMY 4281
	INSERTION OF PERMANENT TUBE INTO ESOPHAGUS
1	4282
	SUTURE OF LACERATION OF ESOPHAGUS
	CLOSURE OF ESOPHAGOSTOMY 4284
	REPAIR OF ESOPHAGEAL FISTULA, NEC
L	

0346 latrogenic Pneumothorax Rate (PSI 6)
4285
REPAIR OF ESOPHAGEAL STRICTURE
PRODUCTION OF SUBCUTANEOUS TUNNEL W/O ESOPHAGEAL ANASTOMOSIS 4287
OTHER GRAFT OF ESOPHAGUS
4289
OTHER REPAIR OF ESOPHAGUS
435
PROXIMAL GASTRECTOMY
4399
TOTAL GASTRECTOMY NEC
ESOPHAGOGASTROPLASTY
4466 OTHER PROCEDURES FOR CREATION OF ESOPHAGOGASTRIC SPHINCTERIC COMPETENCE
4467
LAP CREAT ESOPH SPHINCT
7781
OTH CHEST CAGE OSTECTOMY
7791
TOT CHEST CAGE OSTECTOMY
8104
DORSAL AND DORSO-LUMBAR FUSION, ANTERIOR TECHNIQUE
8134 DEFLISION OF DODSAL AND DODSOLUMBAD SDINE, ANTEDIOD TECHNIQUE
REFUSION OF DORSAL AND DORSOLUMBAR SPINE, ANTERIOR TECHNIQUE ICD-9-CM Lung or pleural biopsy procedure codes:
3326
CLOSED [PERCUTANEOUS] [NEEDLE] BIOPSY OF LUNG
3328
OPEN BIOPSY OF LUNG
3424
PLEURAL BIOPSY
ICD9-CM Cardiac procedure codes:
3510 ODEN LIEADT VALVILLODI ASTV WITHOUT DEDI ACEMENT, UNSDECIFIED VALVIE
OPEN HEART VALVULOPLASTY WITHOUT REPLACEMENT, UNSPECIFIED VALVE 3511
OPEN HEART VALVULOPLASTY OF AORTIC VALVE WITHOUT REPLACEMENT
3512
OPEN HEART VALVULOPLASTY OF MITRAL VALVE WITHOUT REPLACEMENT
3513
OPEN HEART VALVULOPLASTY OF PULMONARY VALVE WITHOUT REPLACEMENT
3514
OPEN HEART VALVULOPLASTY OF TRICUSPID VALVE WITHOUT REPLACEMENT
REPLACEMENT OF UNSPECIFIED HEART VALVE
3521 REPLACEMENT OF AORTIC VALVE WITH TISSUE GRAFT
3522
OTHER REPLACEMENT OF AORTIC VALVE
3523
REPLACEMENT OF MITRAL VALVE WITH TISSUE GRAFT
3524

0346 latrogenic Pneumothorax Rate (PSI 6)
OTHER REPLACEMENT OF MITRAL VALVE
3525 REPLACEMENT OF PULMONARY VALVE WITH TISSUE GRAFT
3526
OTHER REPLACEMENT OF PULMONARY VALVE
3527 DEDIAGEMENT OF TRICUCRID VALVE WITH TICCHE ODAET
REPLACEMENT OF TRICUSPID VALVE WITH TISSUE GRAFT 3528
OTHER REPLACEMENT OF TRICUSPID VALVE 3531
OPERATIONS ON PAPILLARY MUSCLE 3532
OPERATIONS ON CHORDAE TENDINEAE
3533
ANNULOPLASTY 3534
INFUNDIBULECTOMY
3535
OPERATIONS ON TRABECULAE CARNEAE CORDIS 3539
OPERATIONS ON OTHER STRUCTURES ADJACENT TO VALVES OF HEART
3550
REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART WITH PROSTHESIS
3551 REPAIR OF ATRIAL SEPTAL DEFECT WITH PROSTHESIS, OPEN TECHNIQUE
3553
REPAIR OF VENTRICULAR SEPTAL DEFECT WITH PROSTHESIS, OPEN TECHNIQUE
3554 REPAIR OF ENDOCARDIAL CUSHION DEFECT WITH PROSTHESIS
3560
REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART WITH TISSUE GRAFT
3561 REPAIR OF ATRIAL SEPTAL DEFECT WITH TISSUE GRAFT
3562
REPAIR OF VENTRICULAR SEPTAL DEFECT WITH TISSUE GRAFT
3563 REPAIR OF ENDOCARDIAL CUSHION DEFECT WITH TISSUE GRAFT
3570
OTHER AND UNSPECIFIED REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART
3571 OTHER AND UNSPECIFIED REPAIR OF ATRIAL SEPTAL DEFECT
3572
OTHER AND UNSPECIFIED REPAIR OF VENTRICULAR SEPTAL DEFECT
3573
OTHER AND UNSPECIFIED REPAIR OF ENDOCARDIAL CUSHION DEFECT 3581
TOTAL REPAIR OF TETRALOGY OF FALLOT
3582
TOTAL REPAIR OF TOTAL ANOMALOUS PULMONARY VENOUS CONNECTION 3583
TOTAL REPAIR OF TRUNCUS ARTERIOSUS
3584
TOTAL CORRECTION OF TRANSPOSITION OF GREAT VESSELS, NOT ELSEWHERE CLASSIFIED

0346 latrogenic Pneumothorax Rate (PSI 6)
 3591
INTERATRIAL TRANSPOSITION OF VENOUS RETURN 3592
CREATION OF CONDUIT BETWEEN RIGHT VENTRICLE AND PULMONARY ARTERY
3593
CREATION OF CONDUIT BETWEEN LEFT VENTRICLE AND AORTA
3594 CREATION OF CONDUIT BETWEEN ATRIUM AND PULMONARY ARTERY
3595
REVISION OF CORRECTIVE PROCEDURE ON HEART
3597
PERC MTRL VLV REPR W IMP
3598 OTHER OPERATIONS ON SEPTA OF HEART
3599
OTHER OPERATIONS ON VALVES OF HEART
3603
OPEN CHEST CORONARY ARTERY ANGIOPLASTY
3610 AORTOCORONARY BYPASS FOR HEART REVASCULARIZATION, NOT OTHERWISE SPECIFIED
3611
(AORTO)CORONARY BYPASS OF ONE CORONARY ARTERY
(AORTO)CORONARY BYPASS OF TWO CORONARY ARTERIES
3613 (AORTO)CORONARY BYPASS OF THREE CORONARY ARTERIES
3614
(AORTO)CORONARY BYPASS OF FOUR OR MORE CORONARY ARTERIES
3615 CINCLE INTERNAL MANMARY CORONARY ARTERY RYRACC
SINGLE INTERNAL MAMMARY-CORONARY ARTERY BYPASS 3616
DOUBLE INTERNAL MAMMARY-CORONARY ARTERY BYPASS
3617
ABDOMINAL -CORONARY ARTERY BYPASS
3619 OTHER BYPASS ANASTOMOSIS FOR HEART REVASCULARIZATION
362
HEART REVASCULARIZATION BY ARTERIAL IMPLANT
3631
OPEN CHEST TRANSMYOCARDIAL REVASCULARIZATION
3632 OTHER TRANSMYOCARDIAL REVASCULARIZATION
3639
OTHER HEART REVASCULARIZATION
3691
REPAIR OF ANEURYSM OF CORONARY VESSEL
3699 OTHER OPERATIONS ON VESSELS OF HEART
370
PERICARDIOCENTESIS
INCISION OF HEART, NOT OTHERWISE SPECIFIED
3711

0346 latrogenic Pneumothorax Rate (PSI 6)
CARDIOTOMY
3712
PERICARDIOTOMY
3731 PERICARDIECTOMY
3732
EXCISION OF ANEURYSM OF HEART
3733
EXCISION OR DESTRUCTION OF OTHER LESION OR TISSUE OF HEART, OPEN APPROACH
3735
PARTIAL VENTRICULECTOMY
3736 EXCISION OR DESTRUCTION OF LEFT ATRIAL APPENDAGE (LAA) (OCT08)
3737
EXC/DEST HRT LES, THRSPC
3741
IMPLANTATION OF PROSTHETIC CARDIAC SUPPORT DEVICE AROUND THE HEART
3749 OTHER REPAIR OF HEART AND PERICARDIUM
3751
HEART TRANSPLANTATION
3752
IMPLANTATION OF TOTAL REPLACEMENT HEART SYSTEM
3753
REPLACEMENT OF REPAIR OF THORACIC UNIT OF TOTAL REPLACEMENT HEART SYSTEM 3754
REPLACEMENT OR REPAIR OF OTHER IMPLANTABLE COMPONENT OF TOTAL REPLACEMENT HEART SYSTEM
3755
REMOVAL OF INTERNAL BIVENTRICULAR HEART REPLACEMENT SYSTEM (OCT08)
3760
IMPLANTATION OR INSERTION OF BIVENTRICULAR EXTERNAL HEART ASSIST SYSTEM (OCT08) 3761
IMPLANT OF PULSATION BALLOON
3762
INSERTION OF NON-IMPLANTABLE HEART ASSIST SYSTEM
3763
REPAIR OF HEART ASSIST SYSTEM 3764
REMOVAL OF HEART ASSIST SYSTEM
3765
IMPLANT OF EXTERNAL HEART ASSIST SYSTEM
3766
INSERTION OF IMPLANTABLE HEART ASSIST SYSTEM
3767 IMPLANTATION OF CARDIOMYOSTIMULATION SYSTEM
3791
OPEN CHEST CARDIAC MASSAGE
3804
INCISION OF VESSEL, AORTA
RESECTION OF ABDOMINAL AORTA WITH GRAFT REPLACEMENT
INCISION OF VESSEL, OTHER THORACIC 3844

45 SECTION OF THORACIC VESSEL WITH GRAFT REPLACEMENT 64 (CISION OF LESION OF AORTA 65 CISION OF LESION OTHER THORACIC VESSEL 84 35 GATION, DIVISION OF AORTA 85 GATION, DIVISION OF OTHER THORACIC VESSELS 0 15 15 15 15 15 15 15 15 15 15
ICISION OF LESION OF AORTA 65 ICISION OF LESION OTHER THORACIC VESSEL 84 65 GATION , DIVISION OF AORTA 85 GATION, DIVISION OF OTHER THORACIC VESSELS 0 'STEMIC TO PULMONARY ARTERY SHUNT 21 IVAL-PULMONARY ARTERY ANASTOMOSIS 22 10 NRTA-SUBCLAVIAN-CAROTID BYPASS 23 HER INTRATHORACIC VASCULAR SHUNT OR BYPASS 23 HER INTRATHORACIC VASCULAR SHUNT OR BYPASS 23 24 25 26 27 27 27 27 28 29 29 29 20 20 20 20 20 20 20 20 20 20
CISION OF LESION OTHER THORACIC VESSEL 84 GATION , DIVISION OF AORTA 85 GATION, DIVISION OF OTHER THORACIC VESSELS 0 'STEMIC TO PULMONARY ARTERY SHUNT 21 VVAL-PULMONARY ARTERY ANASTOMOSIS 22 DRTA-SUBCLAVIAN-CAROTID BYPASS 23 'HER INTRATHORACIC VASCULAR SHUNT OR BYPASS 23 'HER INTRATHORACIC VASCULAR SHUNT OR BYPASS 23 HIER INTRATHORACIC VASCULAR SHUNT OR BYPASS 24 atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest a, hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
GATION , DIVISION OF AORTA 85 GATION, DIVISION OF OTHER THORACIC VESSELS 0 'STEMIC TO PULMONARY ARTERY SHUNT 21 VAL-PULMONARY ARTERY ANASTOMOSIS 22 0RTA-SUBCLAVIAN-CAROTID BYPASS 23 THER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data D) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. X Female
GATION, DIVISION OF OTHER THORACIC VESSELS 0 (STEMIC TO PULMONARY ARTERY SHUNT 21 VAL-PULMONARY ARTERY ANASTOMOSIS 22 DRTA-SUBCLAVIAN-CAROTID BYPASS 23 THER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest a, hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
STEMIC TO PULMONARY ARTERY SHUNT 21 VAL-PULMONARY ARTERY ANASTOMOSIS 22 DRTA-SUBCLAVIAN-CAROTID BYPASS 23 THER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest a., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
WAL-PULMONARY ARTERY ANASTOMOSIS 22 DRTA-SUBCLAVIAN-CAROTID BYPASS 23 THER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest e, nospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
DRTA-SUBCLAVIAN-CAROTID BYPASS 23 THER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
HER INTRATHORACIC VASCULAR SHUNT OR BYPASS atistical risk model e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (D) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate. x Female
e predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and variates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference pulation used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data ID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest e, nospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected e, multiplied by the reference population rate.
DRG 416 DRG 504 DRG 510 DRG 601 DRG 602 DRG 1103 DRG 1801 DRG 1807 DC 1 DC 6 DC 8 DC 25
DPRDAY Procedure Days Data Not Available DMORB HTN_C DMORB NEURO DMORB CHRNLUNG DMORB DM DMORB DM DMORB DMCX DMORB METS DMORB OBESE DMORB WGHTLOSS DMORB DRUG RL http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/Risk%20Adjustment%20Tables%20PSI%204.3.pdf Not

	0346 latrogenic Pneumothorax Rate (PSI 6)
Type Score	Rate/proportion better quality = lower score
Algorithm	Each indicator is expressed as a rate, is defined as outcome of interest / population at risk or numerator / denominator. The AHRQ Quality Indicators (AHRQ QI) software performs six steps to produce the rates. 1) Discharge-level data is used to mark inpatient records containing the outcome of interest and 2) the population at risk. For provider indicators, the population at risk is also derived from hospital discharge records; for area indicators, the population at risk is derived from U.S. Census data. 3) Calculate observed rates. Using output from steps 1 and 2, rates are calculated for user-specified combinations of stratifiers. 4) Calculate expected rates. Regression coefficients from a reference population database are applied to the discharge records and aggregated to the provider or area level. For indicators that are not risk-adjusted, this is the reference population rate. 5) Calculate risk-adjusted rate. Use the indirect standardization to account for case-mix. For indicators that are not risk-adjusted rates. The shrinkage estimate reflects a reliability adjustment unique to each indicator URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable
	0348 latrogenic Pneumothorax Rate (PDI 5)
Steward	Agency for Healthcare Research and Quality
Description	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code of iatrogenic pneumothorax in any secondary diagnosis field
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code of iatrogenic pneumothorax in any secondary diagnosis field
Numerator Details	Time Window: User may specify the time window; generally one calendar year ICD-9-CM latrogenic pneumothorax diagnosis code:
	5121 IATROGENIC PNEUMOTHORAX
Donominator	
Statement	Discharges, age under 18 years, defined by specific surgical and medical DRGs
Denominator Details	Time Window: All surgical and medical discharges under age 18 defined by specific DRGs or MS-DRGs See Pediatric Quality Indicators Appendices: - Appendix B – Surgical Discharge DRGs - Appendix C – Surgical Discharge MS-DRGs - Appendix D – Medical Discharge DRGs - Appendix E – Medical Discharge MS-DRGs Link to PDI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf
Exclusions	Exclude cases: - neonates with birth weight less than 2500 grams (Birth Weight Category 1-8) - with principal diagnosis of iatrogenic pneumothorax or secondary diagnosis present on admission - with any diagnosis code of chest trauma or pleural effusion - with an ICD-9-CM procedure code of thoracic surgery, lung or pleural biopsy, diaphragmatic surgery repair, OR cardiac surgery

	0348 latrogenic Pneumothorax Rate (PDI 5)
	- normal newborn
	- MDC 14 (pregnancy, childbirth, and puerperium)
	- with missing discharge gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or
	principal diagnosis (DX1=missing)
Exclusion	See Pediatric Quality Indicators Appendices:
Details	- Appendix I – Definitions of Neonate, Newborn, Normal Newborn, and Outborn
	- Appendix L- Low Birth Weight Categories
	Link to PDI appendices:
	http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf
	ICD-9-CM Chest trauma diagnosis codes:
	80700 FRACTURE RIB NOS-CLOSED
	80701
	FRACTURE ONE RIB-CLOSED
	80702
	FRACTURE TWO RIBS-CLOSED
	80703
	FRACTURE THREE RIBS-CLOS
	80704
	FRACTURE FOUR RIBS-CLOSE
	80705
	FRACTURE FIVE RIBS-CLOSE
	80706
	FRACTURE SIX RIBS-CLOSED 80707
	FRACTURE SEVEN RIBS-CLOS
	80708
	FX EIGHT/MORE RIB-CLOSED
	80709
	FX MULT RIBS NOS-CLOSED
	80710
	FRACTURE RIB NOS-OPEN
	80711
	FRACTURE ONE RIB-OPEN 80712
	FRACTURE TWO RIBS-OPEN
	80713
	FRACTURE THREE RIBS-OPEN
	80714
	FRACTURE FOUR RIBS-OPEN
	80715
	FRACTURE FIVE RIBS-OPEN
	FRACTURE SIX RIBS-OPEN
	80717 FRACTURE SEVEN RIBS-OPEN
	80718
	FX EIGHT/MORE RIBS-OPEN
	80719
	FX MULT RIBS NOS-OPEN
	8072
	FRACTURE OF STERNUM-CLOS
	8073

0348 latrogenic Pneumothorax Rate (PDI 5)
FRACTURE OF STERNUM-OPEN
8074
FLAIL CHEST
8075
FX LARYNX/TRACHEA-CLOSED
8076
FX LARYNX/TRACHEA-OPEN
8090
FRACTURE TRUNK BONE-CLOS
8091 FRACTURE TRUNK BONE-OPEN
8600
TRAUM PNEUMOTHORAX-CLOSE
8601
TRAUM PNEUMOTHORAX-OPEN
8602
TRAUM HEMOTHORAX-CLOSED
8603
TRAUM HEMOTHORAX-OPEN
8604
TRAUM PNEUMOHEMOTHOR-CL
8605
TRAUM PNEUMOHEMOTHOR-OPN
86100
HEART INJURY NOS-CLOSED
86101
HEART CONTUSION-CLOSED
86102 HEART LACERATION-CLOSED
86103
HEART CHAMBER LACERAT-CL
86110
HEART INJURY NOS-OPEN
86111
HEART CONTUSION-OPEN
86112
HEART LACERATION-OPEN
86113
HEART CHAMBER LACER-OPN
LUNG INJURY NOS-CLOSED
86121 LUNG CONTUSION-CLOSED
86122
LUNG LACERATION-CLOSED
86130
LUNG INJURY NOS-OPEN
86131
LUNG CONTUSION-OPEN
86132
LUNG LACERATION-OPEN
8620
DIAPHRAGM INJURY-CLOSED

0348 latrogenic Pneumothorax Rate (PDI 5)
8621
DIAPHRAGM INJURY-OPEN
86221 BRONCHUS INJURY-CLOSED
86222
ESOPHAGUS INJURY-CLOSED
86229
INTRATHORACIC INJ NEC-CL
86231
BRONCHUS INJURY-OPEN
ESOPHAGUS INJURY-OPEN 86239
INTRATHORAC INJ NEC-OPEN
8628
INTRATHORACIC INJ NOS-CL
8629
INTRATHORAC INJ NOS-OPEN
8750
OPEN WOUND OF CHEST 8751
OPEN WOUND CHEST-COMPL
8760
OPEN WOUND OF BACK
8761
OPEN WOUND BACK-COMPL
9010
INJURY THORACIC AORTA
9011 INJ INNOMIN/SUBCLAV ART
9012
INJ SUPERIOR VENA CAVA
9013
INJ INNOMIN/SUBCLAV VEIN
90140
INJ PULMONARY VESSEL NOS
90141 INJURY PULMONARY ARTERY
90142
INJURY PULMONARY VEIN
90181
INJ INTERCOSTAL ART/VEIN
90182
INJ INT MAMMARY ART/VEIN
90183 INJ MULT THORACIC VESSEL
90189
INJ THORACIC VESSEL NEC
9019
INJ THORACIC VESSEL NOS
9110
ABRASION TRUNK
9111

0348 latrogenic Pneumothorax Rate (PDI 5)
ABRASION TRUNK-INFECTED
9118
SUPERFIC INJU TRUNK NEC
9119
SUPERFIC INJU TRUNK NEC-INF
9220
CONTUSION OF BREAST
9221
CONTUSION OF CHEST WALL
9223 DACK CONTUSION
BACK CONTUSION 92231
BACK CONTUSION
92233
INTERSCPLR REG CONTUSION
9228
MULIPLE CONTUSION TRUNK
9229
CONTUSION OF TRUNK
92611
CRUSHING INJURY BACK
92619
CRUSHING INJ TRUNK NEC
9268
MULT CRUSHING INJ TRUNK
9269
CRUSHING INJ TRUNK NOS
9290
CRUSH INJ MULT SITE NEC
CRUSHING INJURY NOS 9541
INJ SYMPA NERVE NEC
9548
INJURY TRUNK NERVE NEC
9549
INJURY TRUNK NERVE NOS
95911
INJURY OF CHEST WALL NEC
95919
TRUNK INJURY-SITES NEC
9599
INJURY-SITE NOS
ICD-9-CM Pleural effusion diagnosis codes:
0101
TUBERCULOUS PLEURISY IN PRIMARY PROGRESSIVE TUBERCULOSIS
TUBERCULOUS PLEURISY IN PRIMARY PROGRESSIVE TUBERCULOSIS, UNSPECIFIED
01011 TRIPPT RACTERIAL OR LICTOL OCICAL EVANANOT RONE
TPIPPT, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE
01012 TPIPPT, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN
01013
01013

0348 latrogenic Pneumothorax Rate (PDI 5)
TPIPPT, TUBERCLE BACILI FOUND BY MICROSCOPY
01014 TPIPPT, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE
01015 TPIPPT, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY
01016 TPIPPT, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER METHODS 0117
0117 TUBRCULOUS PNEUMOTHORAX 01170
TUBRCULOUS PNEUMOTHORAX, UNSPECIFIED 01171
TPNEU, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE 01172
TPNEU, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN 01173
TPNEU, TUBERCLE BACILI FOUND BY MICROSCOPY 01174
TPNEU, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE 01175
TPNEU, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY 01176
TPENU, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER METHODS 0120
TUBERCULOUS PLEURISY 01200
TUBERCULOUS PLEURISY, UNSPECIFIED 01201
TP, BACTERIAL OR HISTOLOGICAL EXAM NOT DONE 01202
TP, BACTERIAL OR HISTOLOGICAL EXAM UNKNOWN 01203
TP, TUBERCLE BACILI FOUND BY MICROSCOPY 01204
TP, TUBERCLE BACILI NOT FOUND BY MICROSCOPY BUT BY BACTERIAL CULTURE 01205
TP, TUBERCLE BACILI NOT FOUND BY BACTERIOLOGICAL BUT CONFIRMED HISTOLOGICALLY 01206
TP, TUBERCLE BACILI NOT FOUND BY BACTERILOGICAL OR HISTOLOGICAL BUT CONFIRMED OTHER METHODS 1972
SECOND MALIG NEO PLEURA 5111
WITH EFUSION, WITH MENTION OF A BACTERIAL CAUSE OTHER THAN TUBERCULOSIS 5118
OTHER SPECIFIED FORM OF EFFUSION, EXCEPT TUBERCULOUS 51181
MALIGNANT PLEURAL EFFUSION (OCT08) 51189
OTHER SPECIFIED FORMS OF EFFUSION, EXCEPT TUBERCULOSIS (OCT08) 5119
UNSPECIFIED PLEURAL EFFUSION

0348 latrogenic Pneumothorax Rate (PDI 5)
ICD-9-CM Thoracic surgery procedure codes:
0522
SYMPATHECTOMY CERVICAL
0523 SYMPATHECTOMY LUMBAR
0529
OTHER SYMPATHECTOMY AND GANGLIONECTOMY
0780
THYMECTOMY, NOT OTHERWISE SPECIFIED
0781 OTHER PARTIAL EXCISION OF THYMUS
0782
OTHER TOTAL EXCISION OF THYMUS
0783
THORACOSCOPIC PARTIAL EXCISION OF THYMUS
THORACOSCOPIC TOTAL EXCISION OF THYMUS 3121
MEDIASTINAL TRACHEOSTOMY
3145
OPEN BIOPSY OF LARYNX OR TRACHEA
3173
CLOSURE OF OTHER FISTULA OF TRACHEA 3179
OTHER REPAIR AND PLASTIC OPERATIONS ON TRACHEA
3199
OTHER OPERATIONS ON TRACHEA
3209
OTHER LOCAL EXCISION OR DESTRUCTION OF LESION OR TISSUE OF BRONCHUS 321
OTHER EXCISION OF BRONCHUS
3220
THORAC EXC LUNG LESION
Local excision or destruction of lesion or tissue of lung
3221 PLICATION OF EMPHYSEMATIOUS BLEB
3222
LUNG VOLUME REDUCTION SURGERY
3223
OPEN ABLTN LUNG LES/TISS (OCT06)
PERC ABLTN LUNG LES/TISS (OCT06) 3225
THOR ABLTN LUNG LES/TISS (OCT06)
3226
ABLTN LUNG TISS NEC/NOS (OCT06)
BRNC THRMPLSTY, ABLT MSCL 3228
ENDOSCOPIC EXCISION OR DESTRUCTION OF LESION OR TISSUE OF LUNG
3229
OTHER LOCAL EXCISION OR DESTRUCTION OF LESION OR TISSUE OF LUNG
323

0348 latrogenic Pneumothorax Rate (PDI 5)
SEGMENTAL RESECTION OF LUNG
3230
THORAC SEG LUNG RESECT
3239 OTH SEG LUNG RESECT NOS
324
LOBECTOMY OF LUNG
3241
THORAC LOBECTOMY LUNG
OTHER LOBECTOMY OF LUNG 325
COMPLETE PNEUMONECTOMY
3250
THORACOSPC PNEUMONECTOMY
3259
OTHER PNEUMONECTOMY NOS 326
RADICAL DISSECTION OF THORACIC STRUCTURES
329
OTHER EXCISION OF LUNG
330
INCISION OF BRONCHUS
331 INCISION OF LUNG
3320
THORACOSCOPC LUNG BIOPSY
3325
OPEN BIOPSY OF BRONCHUS
3327 CLOSED ENDOSCOPIC BIOPSY OF LUNG
3328
OPEN BIOPSY OF LUNG
3331
DESTRUCTION OF PHRENIC NERVE FOR COLLAPSE OF LUNG (NO LONGER PERFORMED)
ARTIFICIAL PNEUMOTHORAX FOR COLLAPSE OF LUNG 3334
THORACOPLASTY
3339
OTHER SURGICAL COLLAPSE OF LUNG
Repair and plastic operation on lung and bronchus
3341 SUTURE OF LACERATION OF BRONCHUS
3342
CLOSURE OF BRONCHIAL FISTULA
3343
CLOSURE OF LACERATION OF LUNG
3348 OTHER REPAIR AND PLASTIC OPERATIONS ON BRONCHUS
3349
OTHER REPAIR AND PLASTIC OPERATIONS ON LUNG
Lung transplant

	0348 latrogenic Pneumothorax Rate (PDI 5)
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0348 latrogenic Pneumothorax Rate (PDI 5)
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3481 EXCISION OF LESION OR TISSUE OF DIAPHRAGM
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SUTURE OF LACERATION OF DIAPHRAGM
3483
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3484
OTHER REPAIR OF DIAPHRAGM 3485
3489
OTHER OPERATIONS ON DIAPHRAGM
3493
REPAIR OF PLEURA
3499 OTHER OPERATIONS ON THORAX, OTHER
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CANNULATION OF THORACIC DUCT
4062
FISTULIZATION OF THORACIC DUCT
4063
CLOSURE OF FISTULA OF THORACIC DUCT 4064
LIGATION OF THORACIC DUCT
4069
OTHER OPERATIONS ON THORACIC DUCT
Esophagotomy
INCISION OF ESOPHAGEAL WEB 4209
4207

0348 latrogenic Pneumothorax Rate (PDI 5)
OTHER INCISION OF ESOPHAGUS
4210
ESOPHAGOSTOMY, NOS
4211 CERVICAL ESOPHAGOSTOMY
4212
EXTERIORIZATION OF ESOPHAGEAL POUCH
4219
OTHER EXTERNAL FISTULIZATION OF ESOPHAGUS 4221
OPERATIVE ESOPHAGOSCOPY BY INCISION
4225
OPEN BIOPSY OF ESOPHAGUS 4231
LOCAL EXCISION OF ESOPHAGEAL DIVERTICULUM 4232
LOCAL EXCISION OF OTHER LESION OR TISSUE OF ESOPHAGUS
Excision of esophagus
4239
OTHER DESTRUCTION OF LESION OR TISSUE OF ESOPHAGUS
4240
ESOPHAGECTOMY, NOS
4241 PARTIAL ESOPHAGECTOMY
4242
TOTAL ESOPHAGECTOMY
Intrathoracic anastomosis of exophagus
4251
INTRATHORACIC ESOPHAGOESOPHAGOSTOMY
INTRATHORACIC ESOPHAGOGASTROSTOMY 4253
INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF SMALL BOWEL
4254
OTHER INTRATHORACIC ESOPHAGOENTEROSTOMY
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INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF COLON
OTHER INTRATHORACIC ESOPHAGOCOLOSTOMY 4258
14258 INTRATHORACIC ESOPHAGEAL ANASTOMOSIS W/ OTHER INTERPOSITION
4259
OTHER INTRATHORACIC ANASTOMOSIS OF ESOPHAGUS
Antesternal anastomosis
4261
ANTESTERNAL ESOPHAGOESOPHAGOSTOMY
4262 ANTESTEDNAL ESODUACOCASTDOSTOMY
ANTESTERNAL ESOPHAGOGASTROSTOMY 4263
ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF SMALL BOWEL
4264
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4265

	0348 latrogenic Pneumothorax Rate (PDI 5)
	ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION OF COLON
	4266
	OTHER ANTESTERNAL ESOPHAGOCOLOSTOMY
	4268
	OTHER ANTESTERNAL ESOPHAGEAL ANASTOMOSIS W/ INTERPOSITION
	4269
	OTHER ANTESTERNAL ANASTOMOSIS OF ESOPHAGUS Other repair of esophagus
	427
	ESOPHAGOMYOTOMY
	4281
	INSERTION OF PERMANENT TUBE INTO ESOPHAGUS
	4282
	SUTURE OF LACERATION OF ESOPHAGUS
	CLOSURE OF ESOPHAGOSTOMY 4284
	REPAIR OF ESOPHAGEAL FISTULA, NEC
	4285
	REPAIR OF ESOPHAGEAL STRICTURE
	4286
	PRODUCTION OF SUBCUTANEOUS TUNNEL W/O ESOPHAGEAL ANASTOMOSIS
	4287
	OTHER GRAFT OF ESOPHAGUS
	4289 OTHER REPAIR OF ESOPHAGUS
	435
	PROXIMAL GASTRECTOMY
	4399
	TOTAL GASTRECTOMY NEC
	4465
	ESOPHAGOGASTROPLASTY
	4466 OTHER PROCEDURES FOR CREATION OF ESOPHAGOGASTRIC SPHINCTERIC COMPETENCE
	4467
	LAP CREAT ESOPH SPHINCT
	7781
	OTH CHEST CAGE OSTECTOMY
	7791
	TOT CHEST CAGE OSTECTOMY
	8104 DORSAL AND DORSO-LUMBAR FUSION, ANTERIOR TECHNIQUE
	8134
	REFUSION OF DORSAL AND DORSOLUMBAR SPINE, ANTERIOR TECHNIQUE
	ICD-9-CM Lung or pleural biopsy procedure codes:
	3326
	CLOSED [PERCUTANEOUS] [NEEDLE] BIOPSY OF LUNG
	OPEN BIOPSY OF LUNG
	3424 PLEURAL BIOPSY
	ICD9-CM Diaphragmatic surgery repair codes:
	537
L	

0348 latrogenic Pneumothorax Rate (PDI 5)
ABD REPAIR-DIAPHR HERNIA
5371
LAPAROSCOPIC REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH (OCT08) 5372
OTHER AND OPEN REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH (OCT08)
5375
REPAIR OF DIAPHRAGMATIC HERNIA, ABDOMINAL APPROACH, NOS (OCT08)
THOR REP-DIAPH HERN NOS 5381
DIAPHRAGMATIC PLICATION
5382
PARASTERN HERNIA REPAIR
5583 LAPAROSCOPIC REPAIR OF DIAPHRAGMATIC HERNIA, WITH THORACIC APPROACH (OCT08)
5584
OTHER AND OPEN REPAIR OF DIAPHRAGMATIC HERNIA, WITH THORACIC APPROACH (OCT08)
ICD9-CM Cardiac procedure codes:
3510 ODEN LIEADT VALVUU ODLASTV MUTUOLIT DEDLAGEMENT, UNSDEGLEIED VALVE
OPEN HEART VALVULOPLASTY WITHOUT REPLACEMENT, UNSPECIFIED VALVE 3511
OPEN HEART VALVULOPLASTY OF AORTIC VALVE WITHOUT REPLACEMENT
3512
OPEN HEART VALVULOPLASTY OF MITRAL VALVE WITHOUT REPLACEMENT
3513 OPEN HEART VALVULOPLASTY OF PULMONARY VALVE WITHOUT REPLACEMENT
3514
OPEN HEART VALVULOPLASTY OF TRICUSPID VALVE WITHOUT REPLACEMENT
REPLACEMENT OF UNSPECIFIED HEART VALVE 3521
REPLACEMENT OF AORTIC VALVE WITH TISSUE GRAFT
3522
OTHER REPLACEMENT OF AORTIC VALVE
3523 REPLACEMENT OF MITRAL VALVE WITH TISSUE GRAFT
3524
OTHER REPLACEMENT OF MITRAL VALVE
3525
REPLACEMENT OF PULMONARY VALVE WITH TISSUE GRAFT
3526 OTHER REPLACEMENT OF PULMONARY VALVE
3527
REPLACEMENT OF TRICUSPID VALVE WITH TISSUE GRAFT
3528 OTHER REPLACEMENT OF TRICUS RID VALVE
OTHER REPLACEMENT OF TRICUSPID VALVE 3531
OPERATIONS ON PAPILLARY MUSCLE
3532
OPERATIONS ON CHORDAE TENDINEAE
3533 ANNULOPLASTY
3534

0348 latrogenic Pneumothorax Rate (PDI 5)
INFUNDIBULECTOMY
3535 OPERATIONS ON TRABECULAE CARNEAE CORDIS
3539
OPERATIONS ON OTHER STRUCTURES ADJACENT TO VALVES OF HEART
3550 Dedaid of linedecified sedtal defect of lifeadt with deoctlifese
REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART WITH PROSTHESIS 3551
REPAIR OF ATRIAL SEPTAL DEFECT WITH PROSTHESIS, OPEN TECHNIQUE
REPAIR OF VENTRICULAR SEPTAL DEFECT WITH PROSTHESIS, OPEN TECHNIQUE 3554
REPAIR OF ENDOCARDIAL CUSHION DEFECT WITH PROSTHESIS
3560
REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART WITH TISSUE GRAFT 3561
REPAIR OF ATRIAL SEPTAL DEFECT WITH TISSUE GRAFT
3562
REPAIR OF VENTRICULAR SEPTAL DEFECT WITH TISSUE GRAFT 3563
REPAIR OF ENDOCARDIAL CUSHION DEFECT WITH TISSUE GRAFT
3570
OTHER AND UNSPECIFIED REPAIR OF UNSPECIFIED SEPTAL DEFECT OF HEART
3571 OTHER AND UNSPECIFIED REPAIR OF ATRIAL SEPTAL DEFECT
3572
OTHER AND UNSPECIFIED REPAIR OF VENTRICULAR SEPTAL DEFECT
3573 OTHER AND UNSPECIFIED REPAIR OF ENDOCARDIAL CUSHION DEFECT
3581
TOTAL REPAIR OF TETRALOGY OF FALLOT
3582 TOTAL REPAIR OF TOTAL ANOMALOUS PULMONARY VENOUS CONNECTION
3583
TOTAL REPAIR OF TRUNCUS ARTERIOSUS
3584 TOTAL CORRECTION OF TRANSPOSITION OF GREAT VESSELS, NOT ELSEWHERE CLASSIFIED
3591
INTERATRIAL TRANSPOSITION OF VENOUS RETURN
3592 CREATION OF CONDUIT BETWEEN RIGHT VENTRICLE AND PULMONARY ARTERY
3593
CREATION OF CONDUIT BETWEEN LEFT VENTRICLE AND AORTA
3594
CREATION OF CONDUIT BETWEEN ATRIUM AND PULMONARY ARTERY 3595
REVISION OF CORRECTIVE PROCEDURE ON HEART
PERC MTRL VLV REPR W IMP 3598
OTHER OPERATIONS ON SEPTA OF HEART
OTHER OPERATIONS ON VALVES OF HEART

0348 latrogenic Pneumothorax Rate (PDI 5)
OPEN CHEST CORONARY ARTERY ANGIOPLASTY 3610
AORTOCORONARY BYPASS FOR HEART REVASCULARIZATION, NOT OTHERWISE SPECIFIED
3611 (AORTO)CORONARY BYPASS OF ONE CORONARY ARTERY
3612 (AORTO)CORONARY BYPASS OF TWO CORONARY ARTERIES
3613 (AORTO)CORONARY BYPASS OF THREE CORONARY ARTERIES
3614 (AORTO)CORONARY BYPASS OF FOUR OR MORE CORONARY ARTERIES
3615 SINGLE INTERNAL MAMMARY-CORONARY ARTERY BYPASS
3616 DOUBLE INTERNAL MAMMARY-CORONARY ARTERY BYPASS
3617 ABDOMINAL -CORONARY ARTERY BYPASS
3619 OTHER BYPASS ANASTOMOSIS FOR HEART REVASCULARIZATION
362 HEART REVASCULARIZATION BY ARTERIAL IMPLANT
3631 OPEN CHEST TRANSMYOCARDIAL REVASCULARIZATION
3632 OTHER TRANSMYOCARDIAL REVASCULARIZATION
3639 OTHER HEART REVASCULARIZATION
3691 REPAIR OF ANEURYSM OF CORONARY VESSEL
3699 OTHER OPERATIONS ON VESSELS OF HEART
370 PERICARDIOCENTESIS
3710 INCISION OF HEART, NOT OTHERWISE SPECIFIED
3711 CARDIOTOMY
3712 PERICARDIOTOMY
3731 PERICARDIECTOMY
3732 EXCISION OF ANEURYSM OF HEART
3733 EXCISION OR DESTRUCTION OF OTHER LESION OR TISSUE OF HEART, OPEN APPROACH
3735 PARTIAL VENTRICULECTOMY
3736 EXCISION OR DESTRUCTION OF LEFT ATRIAL APPENDAGE (LAA) (OCT08)
3737 EXC/DEST HRT LES, THRSPC
3741

	0348 latrogenic Pneumothorax Rate (PDI 5)
	IMPLANTATION OF PROSTHETIC CARDIAC SUPPORT DEVICE AROUND THE HEART
	3749
	OTHER REPAIR OF HEART AND PERICARDIUM
	3751
	HEART TRANSPLANTATION
	3752
	IMPLANTATION OF TOTAL REPLACEMENT HEART SYSTEM 3753
	REPLACEMENT OF REPAIR OF THORACIC UNIT OF TOTAL REPLACEMENT HEART SYSTEM 3754
	REPLACEMENT OR REPAIR OF OTHER IMPLANTABLE COMPONENT OF TOTAL REPLACEMENT HEART SYSTEM 3755
	REMOVAL OF INTERNAL BIVENTRICULAR HEART REPLACEMENT SYSTEM (OCT08)
	3760
	IMPLANTATION OR INSERTION OF BIVENTRICULAR EXTERNAL HEART ASSIST SYSTEM (OCT08)
	3761
	IMPLANT OF PULSATION BALLOON
	3762
	INSERTION OF NON-IMPLANTABLE HEART ASSIST SYSTEM
	3763
	REPAIR OF HEART ASSIST SYSTEM 3764
	REMOVAL OF HEART ASSIST SYSTEM
	3765
	IMPLANT OF EXTERNAL HEART ASSIST SYSTEM
	3766
	INSERTION OF IMPLANTABLE HEART ASSIST SYSTEM
	3767
	IMPLANTATION OF CARDIOMYOSTIMULATION SYSTEM
	3791
	OPEN CHEST CARDIAC MASSAGE
	3804 INCISION OF VESSEL, AORTA
	3805
	INCISION OF VESSEL, OTHER THORACIC
1	3844
	RESECTION OF ABDOMINAL AORTA WITH GRAFT REPLACEMENT
	3845
1	RESECTION OF THORACIC VESSEL WITH GRAFT REPLACEMENT
1	
	EXCISION OF LESION OF AORTA
	3865 EXCISION OF LESION OTHER THORACIC VESSEL
	3884
1	LIGATION, DIVISION OF AORTA
1	3885
	LIGATION, DIVISION OF OTHER THORACIC VESSELS
	390
	SYSTEMIC TO PULMONARY ARTERY SHUNT
1	3921
	CAVAL-PULMONARY ARTERY ANASTOMOSIS
	3922
1	AORTA-SUBCLAVIAN-CAROTID BYPASS

	0348 latrogenic Pneumothorax Rate (PDI 5)
	3923 OTHER INTRATHORACIC VASCULAR SHUNT OR BYPASS
Risk Adjustment	Statistical risk model The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, birthweight (500g groups), age in days (29-60, 61-90, 91+), age in years (in 5-year age groups), modified CMS DRG and AHRQ CCS comorbities. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 43 states and approximately 6 million pediatric discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Age in Years 13 to 18 Age in Years 1 to 13 URL http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/Risk%20Adjustment%20Tables%20PDI%204.3.pdf Not applicable
Stratification	Not applicable
Type Score	Rate/proportion better quality = lower score
Algorithm	Each indicator is expressed as a rate, is defined as outcome of interest / population at risk or numerator / denominator. The AHRQ Quality Indicators (AHRQ QI) software performs six steps to produce the rates. 1) Discharge-level data is used to mark inpatient records containing the outcome of interest and 2) the population at risk. For provider indicators, the population at risk is also derived from hospital discharge records; for area indicators, the population at risk is derived from U.S. Census data. 3) Calculate observed rates. Using output from steps 1 and 2, rates are calculated for user-specified combinations of stratifiers. 4) Calculate expected rates. Regression coefficients from a reference population database are applied to the discharge records and aggregated to the provider or area level. For indicators that are not risk-adjusted, this is the reference population rate. 5) Calculate risk-adjusted rate. Use the indirect standardization to account for case-mix. For indicators that are not risk-adjusted rates. The shrinkage estimate reflects a reliability adjustment unique to each indicator URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable

	0349 Transfusion Reaction (PSI 16)
Steward	Agency for Healthcare Research and Quality
Description	The count of medical and surgical discharges for patients age greater than or equal to 18 or in MDC 14 with ICD-9-CM code for transfusion reaction in any secondary diagnosis field.
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges 18 years and older or in MDC 14 with ICD-9-CM codes for transfusion reaction in any secondary diagnosis field of all medical and surgical discharges defined by specific DRGs or MS-DRGs See Patient Safety Indicators Appendices: - Appendix B – Medical Discharge DRGs - Appendix C – Medical Discharge MS-DRGs - Appendix D – Surgical Discharge DRGs - Appendix E – Surgical Discharge MS-DRGs Link to PSI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf

	0349 Transfusion Reaction (PSI 16)
	Exclude cases: -with principal diagnosis of transfusion reaction or secondary diagnosis present on admission -with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)
Numerator Details	Time Window: User may specify the time window; generally one calendar year
	ICD-9-CM Transfusion reaction diagnosis codes: 9996
	ABO INCOMPATIBILITY REACTION 99960
	ABO INCOMPATIBILITY REACTION NOS 99961
	ABO INCOMP/HTR NEC 99962
	ABO INCOMPAT/ACUTE HTR 99963
	ABO INCOMPAT/DELAY HTR 99969
	ABO INCOMPAT REACTN NEC 9997
	RH INCOMPATIBILITY REACTION 99970
	RH INCOMPAT REACTION NOS 99971
	RH INCOMP/HTR NEC 99972
	RH INCOMPAT/ACUTE HTR 99973
	RH INCOMPAT/DELAY HTR 99974
	RH INCOMPAT REACTION NEC
	E8760 MISMATCHED BLOOD IN TRANSFUSION
Denominator Statement	Not applicable
Denominator Details	Time Window: Not applicable
	Not applicable
Exclusions	Not applicable
Exclusion Details	Not applicable
Risk Adjustment	No risk adjustment or risk stratification Not applicable
Stratification	Not applicable
Type Score	Count better quality = lower score
Algorithm	Identify cases meeting the target outcome. Exclude cases meeting the exclusion criteria. Count the number of case at the hospital level. URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable
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	0350 Transfusion Reaction (PDI 13)
Steward	Agency for Healthcare Research and Quality
Description	The count of medical and surgical discharges for patients age less than 18 and not in MDC 14 with ICD-9-CM code for transfusion reaction in any secondary diagnosis field.
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges under age 18 with ICD-9-CM codes for transfusion reaction in any secondary diagnosis field of all medical and surgical discharges defined by specific DRGs or MS-DRGs with the exclusion of neonates, cases in MDC 14 and instances with the outcome of interest was present on admission. See Pediatric Quality Indicators Appendices: - Appendix B – Surgical DRGs - Appendix C – Surgical MS-DRGs - Appendix D – Medical DRGs - Appendix E – Medical MS-DRGs - Appendix I – Definitions of, Neonate, Newborn, Normal Newborn, and Outborn Link to PDI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf Cases excluded with missing gender (SEX=missig, age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)
Numerator	Time Window: User may specify the time window; generally one calendar year
Details	ICD-9-CM Transfusion reaction diagnosis codes: 9996 ABO INCOMPATIBILITY REACTION 99960 ABO INCOMPATIBILITY REACTION 99961 ABO INCOMPATIBILITY REACTION 99963 ABO INCOMPAT/ACUTE HTR 99969 ABO INCOMPAT REACTN NEC 99970 RH INCOMPAT REACTION 99970 RH INCOMPATIBILITY REACTION 99971 RH INCOMPATIBILITY REACTION 99972 RH INCOMPATIBILITY REACTION 99973 RH INCOMPAT/ACUTE HTR 99973 RH INCOMPAT/DELAY HTR 99974 RH INCOMPAT REACTION NEC E8760

	0350 Transfusion Reaction (PDI 13)
	MISMATCHED BLOOD IN TRANSFUSION
Denominator Statement	Not applicable
Denominator Details	Time Window: Not applicable
	Not applicable
Exclusions	Not applicable
Exclusion Details	Not applicable
	No risk adjustment or risk stratification Not applicable
Stratification	Not applicable
Type Score	Count better quality = lower score
	Identify cases meeting the target outcome. Exclude cases meeting the exclusion criteria. Count the number of case at the hospital level. URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable

	0362 Foreign Body left after procedure (PDI 3)
Steward	Agency for Healthcare Research and Quality
Description	Count of discharges with foreign body left in during procedure in medical and surgical discharges among patients less than 18 years and not MDC 14 (pregnancy, childbirth, and puerperium)
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges under age 18 with ICD-9-CM codes for foreign body left in during procedure in any secondary diagnosis field of medical and surgical discharges defined by specific DRGs or MS-DRGs where several exclusions are applied to the numerator. (Details of the numerator, medical and surgical discharges DRGs and MS-DRGs, and exclusions appear in 2a1.3).
Numerator Details	Time Window: User may specify the time window; generally one calendar year
	ICD-9-CM Foreign body left in during procedure diagnosis codes: 9984
	FOREIGN BODY ACCIDENTALLY LEFT DURING A PROCEDURE 9987
	ACUTE REACTIONS TO FOREIGN SUBSTANCE ACCIDENTALLY LEFT DURING A PROCEDURE Foreign body left in during: E8710
	SURGICAL OPERATION E8711 INFUSION OR TRANSFUSION
	E8712 KIDNEY DIALYSIS OR OTHER PERFUSION
	E8713 INJECTION OR VACCINATION

0362 Foreign Body left after procedure (PDI 3)	
E8714	
ENDOSCOPIC EXAMINATION	
E8715	
ASPIRATION OF FLUID OR TISSUE, PUNCTURE, AND CATHETERIZATION E8716	
HEART CATHETERIZATION	
E8717	
REMOVAL OF CATHETER OR PACKING	
E8718	
OTHER SPECIFIED PROCEDURES	
E8719	
UNSPECIFIED PROCEDURE	
See Pediatric Quality Indicators Appendices: - Appendix B – Surgical DRGs	
- Appendix B – Surgical DRGs	
- Appendix D – Medical DRGs	
- Appendix E – Medical MS-DRGs	
Numerator exclusions:	
- with ICD-9-CM codes for foreign body left in during procedure in the principal diagnosis field or secondary diagnosis field	eld if
present on admission	
 normal newborn newborns weighing less than 500 grams (Birth Weight Category 1) 	
- MDC 14 (pregnancy, childbirth, and puerperium)	
- with missing discharge gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) o	r
principal diagnosis (DX1=missing)	
See Pediatric Quality Indicators Appendices:	
- Appendix I – Definitions of Neonate, Newborn, Normal Newborn, and Outborn	
- Appendix L – Low Birth Weight Categories	
Link to PDI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PDI%20Appendices.pdf	
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nominator Time Window: Not applicable	
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Not applicable	
clusions Not applicable	
clusion Not applicable	
tails	
sk No risk adjustment or risk stratification	
justment Not applicable	
ratification Not applicable	
pe Score Count better quality = lower score	
gorithm Identify cases meeting the target outcome. Exclude cases meeting the exclusion criteria. Count the number of case at	the
hospital level. URL Not applicable	
	lf

	0363 Foreign Body Left During Procedure (PSI 5)
Steward	Agency for Healthcare Research and Quality
Description	Count of discharges with foreign body left in during procedure in medical and surgical discharges among patients 18 years

	0363 Foreign Body Left During Procedure (PSI 5)
	and older or MDC 14 (pregnancy, childbirth, and puerperium)
Туре	Outcome
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable
Level	Facility
Setting	Hospital/Acute Care Facility
Numerator Statement	Discharges, 18 years and older or MDC 14 (pregnancy, childbirth, and puerperium), with ICD-9-CM codes for foreign body left in during procedure in any secondary diagnosis field of medical and surgical discharges defined by specific DRGs or MS- DRGs. (Details of medical and surgical discharges defined by specific DRGs or MS-DRGs and exclusions noted in 2a1.3).
Numerator Details	Time Window: User may specify the time window; generally one calendar year
	ICD-9-CM Foreign body left in during procedure diagnosis codes: 9984
	FOREIGN BODY ACCIDENTALLY LEFT DURING A PROCEDURE 9987
	ACUTE REACTIONS TO FOREIGN SUBSTANCE ACCIDENTALLY LEFT DURING A PROCEDURE Foreign body left in during: E8710
	SURGICAL OPERATION E8711
	INFUSION OR TRANSFUSION E8712
	KIDNEY DIALYSIS OR OTHER PERFUSION E8713
	INJECTION OR VACCINATION E8714
	ENDOSCOPIC EXAMINATION E8715
	ASPIRATION OF FLUID OR TISSUE, PUNCTURE, AND CATHETERIZATION E8716 HEART CATHETERIZATION
	E8717 REMOVAL OF CATHETER OR PACKING
	E8718 OTHER SPECIFIED PROCEDURES
	E8719 UNSPECIFIED PROCEDURE
	See Patient Safety Indicators Appendices: - Appendix B – Medical Discharge DRGs
	- Appendix C – Medical Discharge MS-DRGs - Appendix D – Surgical Discharge DRGs - Appendix E – Surgical Discharge MS-DRGs
	Link to PSI appendices: http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf Numerator exclusions include:
	 - cases with the outcome of interest noted as present on admission - cases with the following missing variables: gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal diagnosis (DX1=missing)
Denominator	Not applicable

	0363 Foreign Body Left During Procedure (PSI 5)
Statement	
Denominator Details	Time Window: Not applicable
	Not applicable
Exclusions	Not applicable
Exclusion Details	Not applicable
	No risk adjustment or risk stratification Not applicable
Stratification	Not applicable
Type Score	Count better quality = lower score
Algorithm	Identify cases meeting the target outcome. Exclude cases meeting the exclusion criteria. Count the number of case at the hospital level. URL Not applicable http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf
Copyright	Not applicable

	0371 Venous Thromboembolism Prophylaxis
Steward	The Joint Commission
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. This measure is part of a set of six nationally implemented prevention and treatment measures that address VTE (VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: Incidence of Potentially-Preventable VTE) that are used in The Joint Commission's accreditation process.
Туре	Process
Data Source	Administrative claims, Electronic Clinical Data : Electronic Health Record, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. The vendor may not offer the measure set to hospitals until verification has been passed. Attachment VTE 4.0 ManuaLF-634469565251741848.pdf
Level	Facility, Population : National
Setting	Hospital/Acute Care Facility
Numerator Statement	 Patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given: the day of or the day after hospital admission the day of or the day after surgery end date for surgeries that start the day of or the day after hospital admission
Numerator Details	 Time Window: Episode of Care Five data elements are used to calculate the numerator: Reason for No VTE Prophylaxis – Hospital Admission - Documentation why mechanical or pharmacologic VTE prophylaxis was not administered at hospital admission. Allowable values: Yes or No/UTD. Surgery End Date - The date the surgical procedure ended after hospital admission. Surgical Procedure - A surgical procedure was performed using general or neuraxial anesthesia the day of or the day after hospital admission. Allowable values: Yes or No/UTD VTE Prophylaxis - The type of venous thromboembolism (VTE) prophylaxis documented in the medical record. Allowable values: 1 - 7 or A - None of the above, not documented or UTD. VTE Prophylaxis Date - The month, day, and year that the initial VTE prophylaxis (mechanical and/or

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	pharmacologic) was administered after hospital admission.
Denominator Statement	All discharged hospital inpatients
Denominator Details	Time Window: Episode of care
	Eleven data elements are used to calculate the denominator:
	1. Admission Date – The month, day and year of admission to acute inpatient care.
	 Birthdate - The month, day and year the patient was born. Clinical Trial - Documentation that during this hospital stay the patient was enrolled in a clinical trial in which patients
	with VTE were being studied. Allowable values: Yes or No/UTD
	4. Comfort Measures Only - Physician/advanced practice nurse/physician assistant (physician/APN/PA) documentation
	of comfort measures only. Commonly referred to as "palliative care" in the medical community and "comfort care" by the
	general public. Palliative care includes attention to the psychological and spiritual needs of the patient and support for the
	dying patient and the patient's family. Comfort Measures Only are not equivalent to the following: Do Not Resuscitate (DNR),
	living will, no code, no heroic measure. Allowable values represent the earliest physician/APN/PA documentation: Day 0 or 1, Day 2 or after Timing unclear or Nat Desumented//JTD
	Day 2 or after, Timing unclear or Not Documented/UTD. 5. Discharge Date – The month day and year the patient was discharged from acute care, left against medical advice
	or expired during the stay.
	6. ICD-9-CM Other Diagnosis Codes - The International Classification of Diseases, Ninth Revision, Clinical Modification
	(ICD-9-CM) codes associated with the diagnosis for this hospitalization.
	7. ICD-9-CM Principal Diagnosis Code - The International Classification of Diseases, Ninth Revision, Clinical Modification
	(ICD-9-CM) code associated with the diagnosis established after study to be chiefly responsible for occasioning the admission
	of the patient for this hospitalization. 8. ICD-9-CM Principal Procedure Code - The International Classification of Diseases, Ninth Revision, Clinical
	Modification (ICD-9-CM) code that identifies the principal procedure performed during this hospitalization. The principal
	procedure is the procedure performed for definitive treatment rather than diagnostic or exploratory purposes, or which is
	necessary to take care of a complication.
	9. ICU Admission Date - The day, month and year that the order was written for the patient to be directly admitted or
	transferred (from a lower level of care) to the intensive care unit (ICU).
	10. ICU Admission or Transfer - Documentation that the patient was admitted or transferred to the intensive care unit (ICU) at this hospital. 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.
	11. ICU Discharge Date - The day, month and year that the order was written to discharge the patient from the intensive
	care unit (ICU), left against medical advice (AMA) or expired.
Exclusions	• Patients less than 18 years of age
	 Patients who have a length of stay (LOS) less than two days and greater than 120 days Patients with Comfort Measures Only documented on day of or day after hospital arrival
	 Patients with comon measures only documented on day of or day after hospital anival Patients enrolled in clinical trials
	• 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 LOS greater than or equal to one day
	Patients with ICD-9-CM Principal Diagnosis Code of Mental Disorders or Stroke as defined in Appendix A, Table
	7.01, 8.1 or 8.2
	• Patients with ICD-9-CM Principal or Other Diagnosis Codes of Obstetrics or VTE as defined in Appendix A, Table
	 7.02, 7.03 or 7.04 Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected
	surgeries as defined in Appendix A, Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, 5.24
Exclusion	The patient age in years is equal to the Admission Date minus the Birthdate. Patients less than 18 years are
Details	excluded.
	 Length of stay (LOS) in days is equal to the Discharge Date minus the Admission Date. If the LOS is greater than
	120 days or equal to or less than 2 days, the patient is excluded.

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	Patients with Comfort Measures Only allowable value of 1 (Day 0 or 1) are excluded.
	Patients are excluded if "Yes" is selected for Clinical Trial.
	The data element ICU Admission or Transfer is used to determine if the patient was admitted to the ICU. If "Yes" is
	selected, the case flows to the ICU Admission Date. If the ICU Admission Date is equal to the hospital admission or the ICU
	Admission Date is the day after the hospital admission date, the ICU Admission and ICU Discharge Date are used to
	determine if the patient was in the ICU for one or more days. If the LOS is less than one day, the patient is excluded from
	VTE-1. In addition, if the patient's ICU Admission Date is prior to the hospital admission day, the patient is excluded (direct
i	admit to ICU).
	Patients with ICD-9-CM Principal Diagnosis Code of Mental Disorders or Stroke are excluded. Deticate with ICD-9-CM Principal or Other Diagnosis Codes of Obstatrice or VTE are evoluded
	 Patients with ICD-9-CM Principal or Other Diagnosis Codes of Obstetrics or VTE are excluded. Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected
	surgeries are excluded.
	5
	No risk adjustment or risk stratification
÷	Not applicable
	Not Applicable, the measure is not stratified.
51	Rate/proportion better quality = higher score
3	1. Start processing. Run cases that are included in the VTE Initial Patient Population and pass the edits defined in the
	Transmission Data Processing Flow: Clinical through this measure.
	2. Calculate Length of Stay. Length of Stay, in days, is equal to the Discharge Date minus the Admission Date.
	3. Check Length of Stay
	a. If Length of Stay is less than 2 days, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing.
	b. If Length of Stay is greater than or equal to 2 days, continue processing and proceed to ICD-9-CM Principal
	Diagnosis Code.
	4. Check ICD-9-CM Principal Diagnosis Code
	a. If the ICD-9-CM Principal Diagnosis Code is on Table 7.01, 8.1, or 8.2, the case will proceed to a Measure Category
	Assignment of B and will not be in the Measure Population. Stop processing.
	b. If the ICD-9-CM Principal Diagnosis Code is not on Table 7.01, 8.1, or 8.2, continue processing and proceed to ICD-
	9-CM Principal or Other Diagnosis Code.
	5. Check ICD-9-CM Principal or Other Diagnosis Code
	a. If at least one of the ICD-9-CM Principal or Other Diagnosis Code is on Table 7.02, 7.03, or 7.04, the case will
	proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing.
	b. If none of the ICD-9-CM Principal or Other Diagnosis Code is on Table 7.02, 7.03, or 7.04, continue processing and
	proceed to ICD-9-CM Principal Procedure Code.
	 Check ICD-9-CM Principal Procedure Code If the ICD-9-CM Principal Procedure Code is on Table 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24, the case will
	a. If the ICD-9-CM Principal Procedure Code is on Table 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24, the case will proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing.
	b. If the ICD-9-CM Principal Procedure Code is missing or not on Table 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24,
	continue processing and proceed to Comfort Measures Only.
	7. Check Comfort Measures Only
	a. If Comfort Measures Only is missing, the case will proceed to a Measure Category Assignment of X and will be
	rejected. Stop processing.
1	b. If Comfort Measures Only equals 1, the case will proceed to a Measure Category Assignment of B and will not be in
1	the measure population. Stop processing.
	c. If Comfort Measures Only equals 2, 3, or 4, continue processing and proceed to Clinical Trial.
	8. Check Clinical Trial
	a. If Clinical Trial is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
	processing.
	b. If Clinical Trial equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing
	measure population. Stop processing. c. If Clinical Trial equals No, continue processing and proceed to VTE Prophylaxis.
	 If Clinical Trial equals No, continue processing and proceed to VTE Prophylaxis. Check ICU Admission or Transfer

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a. If ICU Admission or Transfer is missing, the case will proceed to a Measure Category Assignment of X and will be
rejected. Stop processing.
b. If ICU Admission or Transfer is equal to 2 or 3, the case will proceed to VTE Prophylaxis.
c. If ICU Admission or Transfer is equal to 1, continue processing and proceed to ICU Admission Date.
10. Check ICU Admission Date
a. If ICU Admission Date is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
Stop processing.
b. If ICU Admission Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of D
and will be in the Measure Population. Stop processing.
c. If ICU Admission Date equals a Non Unable to Determine Value, continue processing and proceed to the Initial ICU
Day calculation.
11. Calculate Initial ICU Day. Initial ICU Day, in days, is equal to ICU Admission Date minus Admission Date.
12. Check Initial ICU Day
a. If the Initial Day is less than 0 days, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing.
b. If the Initial Day is equal to 0 days or 1 day, the case will proceed to ICU Discharge Date.
c. If the Initial Day is greater than or equal to 2 days, continue processing and proceed to VTE Prophylaxis.
 Check ICU Discharge Date only if Initial ICU Day is less than or equal to 1 day
a. If the ICU Discharge Date is missing, the case will proceed to a Measure Category Assignment of X and will be
rejected. Stop processing.
b. If the ICU Discharge Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of
D and will be in the Measure Population. Stop processing.
c. If the ICU Discharge Date equals a Non Unable to Determine Value, continue processing and proceed to the ICU
LOS calculation.
14. Calculate ICU LOS. ICU LOS is equal to ICU Discharge Date minus ICU Admission Date.
15. Check ICU LOS
a. If ICU LOS is less than zero days, the case will proceed to a Measure Category Assignment of X and will be
rejected. Stop processing.
b. If ICU LOS is greater than or equal to 1 day, the case will proceed to a Measure Category Assignment of B and will
not be in the Measure Population. Stop processing.
c. If ICU LOS is equal to zero days, the case will proceed to VTE Prophylaxis.
16. Check VTE Prophylaxis
a. If VTE Prophylaxis is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
Stop processing.
 If VTE Prophylaxis is equal to A, continue processing and proceed to check Reason for No VTE Prophylaxis – Hospital Admission.
1. If Reason for No VTE Prophylaxis - Hospital Admission is missing, the case will proceed to a Measure Category
Assignment of X and will be rejected. Stop processing.
2. If Reason for No VTE Prophylaxis – Hospital Admission equals No, the case will proceed to a Measure Category
Assignment of D and will be in the Measure Population. Stop processing.
3. If Reason for No VTE Prophylaxis - Hospital Admission equals Yes, the case will proceed to a Measure Category
Assignment of E and will be in the Numerator Population. Stop processing.
c. If VTE Prophylaxis is equal to 1,2,3,4,5,6,7 and not equal to A, continue processing and proceed to VTE Prophylaxis
Date.
17. Check VTE Prophylaxis Date
a. If the VTE Prophylaxis Date is missing, the case will proceed to a Measure Category Assignment of X and will be
rejected. Stop processing.
b. If the VTE Prophylaxis Date equals Unable to Determine, the case will proceed to a Measure Category Assignment
of D and will be in the Measure Population. Stop processing.
c. If the VTE Prophylaxis Date equals a Non Unable to Determine Value, continue processing and proceed to the Initial
Prophylaxis Day calculation.
18. Calculate Initial Prophylaxis Day. Initial Prophylaxis Day, in days, is equal to the VTE Prophylaxis Date minus the
Admission Date.

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	19. Check Initial Prophylaxis Day
	a. If Initial Prophylaxis Day is less than zero days, the case will proceed to a Measure category Assignment of X and
	will be rejected. Stop processing.
	b. If Initial Prophylaxis Day is equal to zero days or 1 day, the case will proceed to a Measure Category Assignment of E and will be in the Numerator Population. Stop processing.
	c. If Initial Prophylaxis Day is greater than or equal to 2 days, continue processing and proceed to Surgical Procedure.
	20. Check Surgical Procedure
	a. If Surgical Procedure is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
	Stop processing.
	b. If Surgical Procedure equals No, the case will proceed to a Measure Category Assignment of D and will be in the
	Measure Population. Stop processing.
	c. If Surgical Procedure equals Yes, continue processing and proceed to Surgery End Date.
	21. Check Surgery End Date
	a. If the Surgery End Date is missing, the case will proceed to a Measure Category Assignment of X and will be
	rejected. Stop processing.
	b. If the Surgery End Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing.
	c. If the Surgery End Date equals a Non Unable to Determine Value, continue processing and proceed to the Initial
	Surgical Prophylaxis Day calculation.
	22. Calculate Initial Surgical Prophylaxis Day. Initial Surgical Prophylaxis Day, in days, is equal to the VTE Prophylaxis
	Date minus Surgery End Date.
	23. Check Initial Surgical Prophylaxis Day
	a. If the Initial Surgical Prophylaxis Day is greater than or equal to 2 days, the case will proceed to a Measure Category
	Assignment of D and will be in the Measure Population. Stop processing. If the Initial Surgical Prophylaxis Day is equal to zero days or 1 day, the case will proceed to a Measure Category Assignment
	of E and will be in the Numerator Population. Stop processing. Attachment 2zq_VTE1.pdf
Copyright	The Specifications Manual for National Hospital Inpatient Quality Measures (Specifications Manual) is the result of the
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	set of national hospital quality measures. A primary objective of this collaborative effort is to promote and enhance the utility of
	these measures for all hospitals.
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	accreditation; including performance measures systems; are required to update their software and associated documentation based on the published manual production timelines.
	Example Acknowledgement: The Specifications Manual for National Hospital Inpatient Quality Measures [Version xx, Month,
	Year] is the collaborative work of the Centers for Medicare & Medicaid Services and The Joint Commission. The Specifications
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	Specifications Manual for National Hospital Inpatient Quality Measures must update their software and associated
	documentation based on the published manual production timelines.

	0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
Steward	The Joint Commission
	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 the initial admission (or transfer) to the Intensive Care Unit (ICU) or surgery end date for surgeries that start the day of or the day after ICU admission (or transfer). This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring by Protocol, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: VTE Incidence of Potentially-Preventable VTE).
Туре	Process

	0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
Data Source	Administrative claims, Electronic Clinical Data, Paper Records Each data element in the data dictionary includes suggested
	data sources.
	The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools
	based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and
	conformance of the data collection tool with the measure specifications. The vendor may not offer the measure set to hospitals
	until verification has been passed. Attachment VTE 4.0 ManuaLF-634469622988616848.pdf
Level	Facility, Population : National
Setting	Hospital/Acute Care Facility
Numerator Statement	Patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given:
Statement	the day of or the day after ICU admission (or transfer)
	 the day of or the day after surgery end date for surgeries that start the day of or the day after ICU admission (or
	transfer)
Numerator	Time Window: Episode of Care
Details	
	Six data elements are used to calculate the numerator:
	1. Anesthesia Start Date The date the anesthesia for the procedure started.
	2. ICU VTE Prophylaxis The type of venous thromboembolism (VTE) prophylaxis that was initially administered in the
	ICU. Allowable values: 1 - 7 or A – None of the above, not documented or UTD.
	 ICU VTE Prophylaxis Date The day, month and year that the initial VTE prophylaxis (mechanical and/or pharmacologic) option was administered after admission/transfer to the intensive care unit (ICU).
	4. Reason for No VTE Prophylaxis – ICU Admission Documentation why mechanical or pharmacologic VTE
	prophylaxis was not administered at ICU admission/transfer. Allowable values: Yes or No/UTD.
	5. Surgery End Date – ICU Admission The date the surgical procedure ended after ICU admission or transfer.
	6. Surgical Procedure – ICU Admission A surgical procedure was performed using general or neuraxial anesthesia the
	day of or the day after ICU admission or transfer. Allowable values: Yes or No/UTD
Denominator	Patients directly admitted or transferred to ICU
Statement	
Denominator Details	Time Window: Episode of care
Dottano	Eleven data elements are used to calculate the denominator:
	1. Admission Date – The month, day and year of admission to acute inpatient care.
	2. Birthdate - The month, day and year the patient was born.
	3. Clinical Trial - Documentation that during this hospital stay the patient was enrolled in a clinical trial in which patients
	with the same condition as the measure set were being studied. Allowable values: Yes or No/UTD
	4. Comfort Measures Only - Physician/advanced practice nurse/physician assistant (physician/APN/PA) documentation
	of comfort measures only. Commonly referred to as "palliative care" in the medical community and "comfort care" by the
	general public. Palliative care includes attention to the psychological and spiritual needs of the patient and support for the dying patient and the patient's family. Comfort Measures Only are not equivalent to the following: Do Not Resuscitate (DNR),
	living will, no code, no heroic measure. Allowable values represent the earliest physician/APN/PA documentation: Day 0 or 1,
	Day 2 or after, Timing unclear or Not Documented/UTD.
	5. Discharge Date – The month day and year the patient was discharged from acute care, left against medical advice
	or expired during the stay.
	6. ICD-9-CM Other Diagnosis Codes - The International Classification of Diseases, Ninth Revision, Clinical
	Modification (ICD-9-CM) codes associated with the diagnosis for this hospitalization.
	7. ICD-9-CM Principal Diagnosis Code - The International Classification of Diseases, Ninth Revision, Clinical
	Modification (ICD-9-CM) code associated with the diagnosis established after study to be chiefly responsible for occasioning the admission of the patient for this hospitalization.
	8. ICD-9-CM Principal Procedure Code - The International Classification of Diseases, Ninth Revision, Clinical
	Modification (ICD-9-CM) code that identifies the principal procedure performed during this hospitalization. The principal
	procedure is the procedure performed for definitive treatment rather than diagnostic or exploratory purposes, or which is

	0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
	 necessary to take care of a complication. 9. ICU Admission Date - The day, month and year that the order was written for the patient to be directly admitted or transferred (from a lower level of care) to the intensive care unit (ICU). 10. ICU Admission or Transfer - Documentation that the patient was admitted or transferred to the intensive care unit (ICU) at this hospital. 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. 11. ICU Discharge Date - The day, month and year that the order was written to discharge the patient from the intensive care unit (ICU), left against medical advice (AMA) or expired. Please note: The majority of general data elements that are missing data cause the EOC record to be rejected.
	 Patients less than 18 years of age Patients who have a hospital length of stay (LOS) less than two days and greater than 120 days Patients with Comfort Measures Only documented on day of or day after hospital arrival Patients enrolled in clinical trials Patients with ICU LOS less than one day without VTE prophylaxis administered and documentation for no VTE prophylaxis Patients with ICD-9-CM Principal or Other Diagnosis Code of Obstetrics or VTE as defined in Appendix A, Table 7.02, 7.03, or 7.04 Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected surgeries as defined in Appendix A, Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, 5.24 that start the day of or the day after ICU admission or transfer
	 The patient age in years is equal to the Admission Date minus the Birthdate. The month and day portion of the admission date and birthdate are used to yield the most accurate age. Patients less than 18 years are excluded. Length of stay (LOS) in days is equal to the Discharge Date minus the Admission Date. If the LOS is greater than 120 days or equal to or less than 2 days, the patient is excluded. Patients with Comfort Measures Only allowable value of 1 (Day 0 or 1) are excluded. Patients are excluded if "Yes" is selected for Clinical Trial. The data element ICU Admission or Transfer is used to determine if the patient was admitted to the ICU. If "Yes" is selected, the case flows to the ICU Admission Date and ICU Discharge Date. The ICU Admission and ICU Discharge Date are used to determine if the patient was in the ICU for one or more days. If the ICU LOS is less than one day, the patient is excluded from VTE-2. Patients with ICD-9-CM Principal or Other Diagnosis Codes of Obstetrics or VTE are excluded. Patients with ICD-9-CM Principal Procedure Code of Surgical Care Improvement Project (SCIP) VTE selected surgeries are excluded if the surgery started the day of or the day after ICU admission or transfer.
	No risk adjustment or risk stratification
	Not applicable Not Applicable, the measure is not stratified.
	Rate/proportion better quality = higher score
Algorithm	 Start processing. Run cases that are included in the VTE Initial Patient Population and pass the edits defined in the Transmission Data Processing Flow: Clinical through this measure. Calculate Length of Stay. Length of Stay, in days, is equal to the Discharge Date minus the Admission Date. Check Length of Stay If Length of Stay is less than 2 days, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing. If Length of Stay is greater than or equal to 2 days, continue processing and proceed to ICD-9-CM Principal or Other Diagnosis Code Check ICD-9-CM Principal or Other Diagnosis Code is on Table 7.02, 7.03, or 7.04, the case will proceed to a Measure Category Assignment of B and will not be in the Measure Category Assignment of B and will not be in the Diagnosis Code is on Table 7.02, 7.03, or 7.04, the case will proceed to a Measure Category Assignment of B and will not be in the Diagnosis Code is on Table 7.02, 7.03, or 7.04, the case will proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing.

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5. Check Comfort Measures Only
a. If Comfort Measures Only is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
 b. If Comfort Measures Only equals 1, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing.
c. If Comfort Measures Only equals 2, 3, or 4, continue processing and proceed to Clinical Trial.
 Check Clinical Trial If Clinical Trial is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
processing. b. If Clinical Trial equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the measure
population. Stop processing. c. If Clinical Trial equals No, continue processing and proceed to ICU Admission or Transfer.
7. Check ICU Admission or Transfer
a. If ICU Admission or Transfer is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
b. If ICU Admission or Transfer is equal to 2 or 3, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing.
c. If ICU Admission or Transfer is equal to 1, continue processing and proceed to ICU Admission Date.
 Check ICU Admission Date If ICU Admission Date is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
processing.
b. If ICU Admission Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing.
c. If ICU Admission Date equals a Non Unable to Determine Value, continue processing and proceed to the ICD-9-CM Principal Procedure Code.
9. Check ICD-9-CM Principal Procedure Code
a. If ICD-9-CM Principal Procedure Code is on Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24, the case will proceed to Anesthesia Start Date.
b. If ICD-9-CM Principal Procedure Code is missing or not on Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24, the case will proceed to ICU Discharge Date.
10. Check Anesthesia Start Date only if ICD-9-CM Principal Procedure Code is on Tables 5.17, 5.19, 5.20, 5.21, 5.22, 5.23, or 5.24.
a. If Anesthesia Start Date is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
 b. If Anesthesia Start Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing.
c. If Anesthesia Start Date equals a Non Unable to Determine Value, continue processing and proceed to the ICU Initial
Surgery Day calculation. 11. Calculate ICU Initial Surgery Day. ICU Initial Surgery Day, in days, is equal to the Anesthesia Start Date minus the ICU
Admission Date. 12. Check ICU Initial Surgery Day
a. If ICU Initial Surgery Day is less than or equal to 1 day, the case will proceed to a Measure Category Assignment of B and
will not be in the Measure Population. Stop processing. b. If Initial Surgery Day is greater than or equal to 2 days, continue processing and proceed to ICU Discharge Date.
13. Check ICU Discharge Date a. If ICU Discharge Date is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
processing.
b. If ICU Discharge Date equals Unable to Determine, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing.
c. If ICU Discharge Date equals a Non Unable to Determine Value, continue processing and proceed to the ICU VTE Prophylaxis.
14. Check ICU VTE Prophylaxis
a. If ICU VTE Prophylaxis is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop

	0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
	processing. b. If ICU VTE Prophylaxis is equal to A, continue processing and proceed to Reason for No VTE Prophylaxis – ICU Admissi
	c. If ICU VTE Prophylaxis is equal to 1, 2, 3, 4, 5, 6, 7 and not equal to A, continue processing and proceed to ICU VTE Prophylaxis Date.
	15. Check Reason for No VTE Prophylaxis – ICU Admission only if ICU VTE Prophylaxis is equal to A.
	a. If Reason for No VTE Prophylaxis – ICU Admission is missing, the case will proceed to a Measure Category Assignment
	X and will be rejected. Stop processing.
	b. If Reason for No VTE Prophylaxis - ICU Admission equals Yes, the case will proceed to a Measure Category Assignmen
	of E and will be in the Numerator Population. Stop processing.
	c. If Reason for No VTE Prophylaxis – ICU Admission equals No, continue processing and proceed to the ICU LOS
	calculation.
	16. Calculate ICU LOS. ICU LOS is equal to ICU Discharge Date minus ICU Admission Date.
	 Check ICU LOS If ICU LOS is less than zero days, the case will proceed to a Measure Category Assignment of X and will be rejected. Sto
	processing.
	b. If ICU LOS is equal to zero days, the case will proceed to a Measure Category Assignment of B and will not be in the
	Measure Population. Stop processing.
	c. If ICU LOS is greater than or equal to 1 day, the case will proceed to a Measure Category Assignment of D and w
	be in the Measure Population. Stop processing.
	18. Check ICU VTE Prophylaxis Date
	a. If ICU VTE Prophylaxis Date is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	 b. If ICU VTE Prophylaxis Date equals Unable to Determine, the case will proceed to a Measure Category Assignm
	of D and will be in the Measure Population. Stop processing.
	c. If ICU VTE Prophylaxis Date equals a Non Unable to Determine Value, continue processing and proceed to the I
	Initial Prophylaxis Day calculation.
	19. Calculate ICU Initial Prophylaxis Day. ICU Initial Prophylaxis Day, in days, is equal to ICU VTE Prophylaxis Date minus
	ICU Admission Date.
	20. Check ICU Initial Prophylaxis Day
	a. If ICU Initial Prophylaxis Day is less than zero days, the case will proceed to a Measure Category Assignment of X and v be rejected. Stop processing.
	b. If ICU Initial Prophylaxis Day is equal to zero days or 1 day, the case will proceed to a Measure Category Assignment of
	and will be in the Numerator Population. Stop processing.
	c. If ICU Initial Prophylaxis Day is greater than or equal to 2 days, continue processing and proceed to Surgical Procedure
	ICU Admission.
	21. Check Surgical Procedure – ICU Admission
	a. If Surgical Procedure ICU Admission is missing, the case will proceed to a Measure Category Assignment of X and will b
	rejected. Stop processing.
	b. If Surgical Procedure ICU Admission equals No, the case will proceed to a Measure Category Assignment of D and will be the Measure Deputation.
	in the Measure Population. Stop processing. c. If Surgical Procedure ICU Admission equals Yes, continue processing and proceed to Surgery End Date - ICU Admissio
	22. Check Surgery End Date - ICU Admission
	a. If Surgery End Date - ICU Admission is missing, the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the case will proceed to a Measure Category Assignment of X and will the category Assignment of X an
	rejected. Stop processing.
	b. If Surgery End Date - ICU Admission equals Unable to Determine, the case will proceed to a Measure Category Assignment
	of D and will be in the Measure Population. Stop processing.
	c. If Surgery End Date - ICU Admission equals a Non Unable to Determine Value, continue processing and proceed to the
	Initial Surgical Prophylaxis Day calculation.
	23. Calculate ICU Initial Surgical Prophylaxis Day. ICU Initial Surgical Prophylaxis Day, in days, is equal to the ICU VTE
	Prophylaxis Date minus Surgery End Date - ICU Admission.
-	24. Check ICU Initial Surgical Prophylaxis Day
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	Comments due by March 19, 2012 by 6:00pm ET
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	0372 Intensive Care Unit Venous Thromboembolism Prophylaxis
	 a. If ICU Initial Surgical Prophylaxis Day is greater than or equal to 2 days, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing. b. If ICU Initial Surgical Prophylaxis Day is equal to zero days or 1 day, the case will proceed to a Measure Category Assignment of E and will be in the Numerator Population. Stop processing. Attachment 2zr_VTE2.pdf
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	0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
Steward	The Joint Commission
Description	This measure assesses the number of patients diagnosed with confirmed VTE who received an overlap of Parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulation and warfarin therapy. For patients who received less than five days of overlap therapy, they should be discharged on both medications and have a Reason for Discontinuation of Overlap Therapy. Overlap therapy should be administered for at least five days with an international normalized ratio (INR) greater than or equal to 2 prior to discontinuation of the parenteral anticoagulation therapy, or INR less than 2 but discharged on both medications or have a Reason for Discontinuation of Overlap Therapy. This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: Incidence of Potentially-Preventable VTE).
Туре	Process
Data Source	Administrative claims, Electronic Clinical Data, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. Verification must be completed and passed before the vendor can offer the data collection tool to hospitals. Attachment VTE 4.0 ManuaLF-634469519104709898.pdf
Level	Facility, Population : National
Setting	Hospital/Acute Care Facility
Numerator Statement	 Patients who received overlap therapy: Included Populations: Patients who received warfarin and parenteral anticoagulation: Five or more days, with an INR greater than or equal to 2 prior to discontinuation of parenteral therapy OR Five or more days, with an INR less than 2 and discharged on overlap therapy OR Less than five days and discharged on overlap therapy OR With documentation of reason for discontinuation of overlap therapy OR With documentation of a reason for no overlap therapy
Numerator	Time Window: Episode of care
Details	Six data elements are used to calculate the numerator: 1. INR Value - Documentation of an international normalized ratio (INR) value greater than or equal to 2 prior to

0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
 discontinuation of the parenteral anticoagulation therapy. Allowable Value (AV): Yes or No/UTD Overlap Therapy - Documentation that parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulation therapy and warfarin were both administered on the same day or a reason is documented why overlap therapy was not initiated. Allowable Value: 1- There was documentation of overlap therapy; 2 -There is a reason for no overlap therapy; or 3- There was no overlap therapy and no reason/UTD. Overlap Therapy Start Date - The first date that the parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulation therapy and warfarin were administered. Parenteral Anticoagulant End Date - The last date that a parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulant medication was administered. Parenteral Anticoagulant Prescribed at Discharge - Documentation that a parenteral (intravenous [IV] or subcutaneous [subcu]) anticoagulant medication was prescribed at discharge. Allowable Value: Yes or No/UTD Reason for Discontinuation of Overlap Therapy - Documentation of a reason for discontinuation of the overlap therapy by a physician/advanced practice nurse/physician assistant or pharmacist (physician/APN/PA or pharmacist). Allowable Value: Yes or No/UTD
Patients with confirmed VTE who received warfarin. The target population includes patients discharged with an ICD-9-CM Principal or Other Diagnosis Codes for VTE as defined in Table 7.03 or Table 7.04.
Time Window: Episode of Care
 Eleven data elements are used to calculate the denominator: Admission Date – The month, day and year of admission to acute inpatient care. Birthdate - The month, day and year the patient was born. Clinical Trial - Documentation that during this hospital stay the patient was enrolled in a clinical trial in which patients with the same condition as the measure set were being studied. Allowable values: Yes or No/UTD Comfort Measures Only - Physician/advanced practice nurse/physician assistant (physician/APN/PA) documentation of confort measures only. Commonly referred to as 'palliative care' in the medical community and 'comfort care' by the general public. Palliative care includes attention to the psychological and spiritual needs of the patient and support for the dying patient and the patient's family. Comfort Measures Only are not equivalent to the following: Do Not Resuscitate (DNR), living will, no code, and no heroic measure. Allowable values represent the earliest physician/APN/PA documentation: (AV 1) Day 0 or 1, (AV 2) Day 2 or after, (AV 3) Timing unclear or (AV 4) Not Documented/UTD. Discharge Date – The month day and year the patient was discharged on the day of discharge. Allowable values: 1-8. ICD-9-CM Other Diagnosis Codes - The International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM) code associated with the secondary diagnoses for this hospitalization. ICD-9-CM Principal Diagnosis Code - The International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM) code associated with the diagnosis established after study to be chiefly responsible for occasioning the adiagnosis of VTE [deep vein thrombosis (DVT) and/or pulmonary embolism (PE)] was confirmed in a defined location. Allowable values: Yes or No/UTD VTE Diagnostic Test – Documentation that a diagnostic test for VTE was performed. Allowable values: Yes or No/UTD Warfarin Administration - Documenta
 Patients less than 18 years of age Patients who have a length of stay greater than 120 days Patients with Comfort Measures Only documented Patients enrolled in clinical trials Patients discharged to a health care facility for hospice care Patients discharged to home for hospice care Patients who expired Patients who left against medical advice

	0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
	Patients discharged to another hospital
	Patients without warfarin therapy during hospitalization
	 Patients without VTE confirmed by diagnostic testing
Exclusion Details	 Patient without a Principal or Other ICD-9-CM Diagnosis Code on Table 7.03 or 7.04 are excluded. The patient age in years is equal to the Admission Date minus the Birthdate. Patients less than 18 years are
	excluded.
	Length of stay (LOS) in days is equal to the Discharge Date minus the Admission Date. If the LOS is greater than
	120 days, the patient is excluded.
	 Patients with AV 1,2 or 3 for Comfort Measures Only are excluded.
	Patients are excluded if "Yes" is selected for Clinical Trial.
	The allowable values (AV) 2, 3, 4, 6 and 7 for Discharge Disposition exclude patients who are discharged to a heal facility for boarding care boarded and a facility for boarded and a fa
	 care facility for hospice care, home to hospice care, expired, left against medical advice, or to another hospital. Patients are excluded if "No" is selected for Warfarin Administration.
	 Patients are excluded if "No" is selected for VTE Diagnostic Test.
	 Patients are excluded if "No" is selected for VTE Confirmed.
Risk	No risk adjustment or risk stratification
	Not Applicable
-	Not Applicable, the measure is not stratified.
Type Score	Rate/proportion better quality = higher score
Algorithm	1. Start processing. Run cases that are included in the VTE Initial Patient Population and pass the edits defined in
	Transmission Data Processing Flow: Clinical through this measure.
	2. Check ICD-9-CM Principal or Other Diagnosis Code
	a. If none of the ICD-9-CM Principal or Other Diagnosis Code is on Table 7.03 or 7.04 (VTE, Obstetrics-VTE), the cas
	will proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing.
	b. If at least one of the ICD-9-CM Principal or Other Diagnosis Code is on Table 7.03 or 7.04, continue processing an proceed to Comfort Measures Only.
	3. Check Comfort Measures Only
	a. If Comfort Measures Only is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	b. If Comfort Measures Only equals 1, 2 or 3, the case will proceed to a Measure Category Assignment of B and will
	not be in the measure population. Stop processing.
	c. If Comfort Measures Only equals 4, continue processing and proceed to Clinical Trial.
	4. Check Clinical Trial
	a. If Clinical Trial is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
	processing.
	b. If Clinical Trial equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the
	measure population. Stop processing.
	 c. If Clinical Trial equals No, continue processing and proceed to Discharge Disposition. 5. Check Discharge Disposition
	a. If Discharge Disposition equals 2, 3, 4, 6, 7, the case will proceed to a Measure Category Assignment of B and will
	not be in the measure population. Stop processing.
	b. If Discharge Disposition equals 1, 5, 8, continue processing and proceed to VTE Diagnostic Test.
	6. Check VTE Diagnostic Test
	a. If VTE Diagnostic Test is missing, the case will proceed to a Measure Category Assignment of X and will be
	rejected. Stop processing.
	b. If VTE Diagnostic Test equals No, the case will proceed to a Measure Category Assignment of B and will not be in
	the measure population. Stop processing.
	c. If VTE Diagnostic Test equals Yes, continue processing and proceed to VTE Confirmed.
	7. Check VTE Confirmed
	a. If VTE Confirmed is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
	Stop processing.
	b. If VTE Confirmed equals No, the case will proceed to a Measure Category Assignment of B and will not be in the

0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
measure population. Stop processing.
c. If VTE Confirmed equals Yes, continue processing and proceed to Warfarin Administration.
8. Check Warfarin Administration
a. If Warfarin Administration is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
Stop processing.
If Warfarin Administration equals No, the case will proceed to a Measure Category Assignment of B and will not be in the
measure population. Stop processing.
c. If Warfarin Administration equals Yes, continue processing and proceed to Overlap Therapy.
9. Check Overlap Therapy
a. If Overlap Therapy is missing, the case will proceed to a Measure Category Assignment of X and will be rejected.
Stop processing.
b. If Overlap Therapy equals 2, the case will proceed to a Measure Category Assignment of B and will not be in the
Measure Population. Stop processing.
c. If Overlap Therapy equals 3, the case will proceed to a Measure Category Assignment of D and will be in the
Measure Population. Stop processing.
d. If Overlap Therapy equals 1, continue processing and proceed to the Overlap Therapy Start Date.
10. Check Overlap Therapy Start Date
a. If Overlap Therapy Start Date is missing, the case will proceed to a Measure Category Assignment of X and will be
rejected. Stop processing.
b. If the Overlap Therapy Start Date equals Unable to Determine, the case will proceed to a Measure Category
Assignment of D and will be in the Measure Population. Stop processing.
c. If Overlap Therapy Start Date equals a Non Unable to Determine Value, continue processing and proceed to the
Parenteral Anticoagulant End Date.
11. Check Parenteral Anticoagulant End Date
a. If Parenteral Anticoagulant End Date is missing, the case will proceed to a Measure Category Assignment of X and
will be rejected. Stop processing.
b. If the Parenteral Anticoagulant End Date equals Unable to Determine, the case will proceed to a Measure Category
Assignment of D and will be in the Measure Population. Stop processing.
c. If Parenteral Anticoagulant End Date equals a Non Unable to Determine Value, continue processing and proceed to
the Overlap Therapy Days calculation.
12. Calculate Overlap Therapy Days. Overlap Therapy Days, in days, is equal to Parenteral Anticoagulant End Date
minus Overlap Therapy Start Date.
13. Check Overlap Therapy Days
a. If Overlap Therapy Days is less than 0 days, the case will proceed to a Measure Category Assignment of D and will
be in the Measure Population. Stop processing.
b. If Overlap Therapy Days is greater than or equal to 4 days, continue processing and proceed to INR Value.
a. If INR Value is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop
processing.
b. If INR Value equals Yes, the case will proceed to a Measure Category Assignment of E and will be in the Numerator
Population. Stop processing.
c. If INR Value equals No, continue processing and proceed to Parenteral Anticoagulant Prescribed at Discharge.
c. If Overlap Therapy Days is greater than or equal to zero days and less than 4 days, continue processing and
proceed to Parenteral Anticoagulant Prescribed at Discharge.
14. Check Parenteral Anticoagulant Prescribed at Discharge
a. If Parenteral Anticoagulant Prescribed at Discharge is missing, the case will proceed to a Measure Category
Assignment of X and will be rejected. Stop processing.
b. If Parenteral Anticoagulant Prescribed at Discharge equals Yes, the case will proceed to a Measure Category
Assignment of E and will be in the Numerator Population. Stop processing.
c. If Parenteral Anticoagulant Prescribed at Discharge equals No, continue processing and proceed to Reason for
Discontinuation of Overlap Therapy.
15. Check Reason for Discontinuation of Overlap Therapy
a. If Reason for Discontinuation of Overlap Therapy is missing, the case will proceed to a Measure Category
Assignment of X and will be rejected. Stop processing.
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	0373 Venous Thromboembolism Patients with Anticoagulant Overlap Therapy
	 b. If Reason for Discontinuation of Overlap Therapy equals Yes, the case will proceed to a Measure Category Assignment of E and will be in the Numerator Population. Stop processing. c. If Reason for Discontinuation of Overlap Therapy equals No, the case will proceed to a Measure Category Assignment of D and will be in the Measure Population. Stop processing. Attachment 2zs_VTE3.pdf
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	0376 Incidence of Potentially Preventable Venous Thromboembolism
Steward	The Joint Commission
Description	This measure assesses the number of patients with confirmed venous thromboembolism (VTE) during hospitalization (not present at admission) who did not receive VTE prophylaxis between hospital admission and the day before the VTE diagnostic testing order date. This measure is part of a set of six prevention and treatment measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count Monitoring by Protocol, and VTE-5: VTE Warfarin Therapy Discharge Instructions).
Туре	Outcome
Data Source	Administrative claims, Paper Records Each data element in the data dictionary includes suggested data sources. The data are collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy and conformance of the data collection tool with the measure specifications. Verification must be completed and passed before the vendor can offer the data collection tool to hospitals. Attachment VTE 4.0 ManuaLF-634469532965647398.pdf
Level	Facility, Population : National
Setting	Hospital/Acute Care Facility
Numerator Statement	Patients who received no VTE prophylaxis prior to the VTE diagnostic test order date
Numerator Details	Time Window: Episode of Care One data element is used to calculate the numerator: VTE Prophylaxis Status - Documentation of VTE prophylaxis (mechanical and/or pharmacologic) administration between the hospital admission date and the day before the VTE diagnostic test order date. Allowable Value (AV): 1 There is documentation that VTE prophylaxis was administered between the day of admission and the day before the VTE diagnostic test order date, 2 There is no documentation that VTE prophylaxis was administered between the day of admission and the day before the VTE diagnostic test order date or unable to determine from medical record documentation, or 3 There is physician/advanced practice nurse/physician assistant (physician/APN/PA) or pharmacist documentation of a reason for not administering mechanical and pharmacological VTE prophylaxis during hospitalization.
Denominator	Patients who developed confirmed VTE during hospitalization. The target population includes patients discharged with an ICD-

	0376 Incidence of Potentially Preventable Venous Thromboembolism
Statement	9-CM Secondary Diagnosis Codes for VTE as defined in Table 7.03 or Table 7.04.
Denominator Details	Time Window: Episode of Care
	Ten data elements are used to calculate the denominator:
	1. Admission Date – The month, day and year of admission to acute inpatient care.
	2. Birthdate - The month, day and year the patient was born.
	Clinical Trial - Documentation that during this hospital stay the patient was enrolled in a clinical trial in which patients with the same condition as the measure set were being studied. Allowable values: Yes or No/UTD
	4. Comfort Measures Only - Physician/advanced practice nurse/physician assistant (physician/APN/PA) documentation of
	comfort measures only. Commonly referred to as "palliative care" in the medical community and "comfort care" by the general
	public. Palliative care includes attention to the psychological and spiritual needs of the patient and support for the dying patient
	and the patient's family. Comfort Measures Only are not equivalent to the following: Do Not Resuscitate (DNR), living will, no code, no heroic measure. Allowable values represent the earliest physician/APN/PA documentation: (AV 1) Day 0 or 1, (AV 2) Day 2 or after, (AV 3) Timing unclear or (AV 4) Not Documented/UTD.
	5. Discharge Date – The month day and year the patient was discharged from acute care, left against medical advice or expired during the stay.
	6. ICD-9-CM Other Diagnosis Codes - The International Classification of Diseases, Ninth Revision, Clinical Modification (ICD- 9-CM) codes associated with the Secondary diagnoses for this hospitalization.
	7. ICD-9-CM Principal Diagnosis Code - The International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM) code associated with the diagnosis established after study to be chiefly responsible for occasioning the admission of the patient for this hospitalization.
	8. VTE Confirmed – Documentation by a physician/advanced practice nurse/physician assistant (physician/APN/PA) that a diagnosis of VTE [deep vein thrombosis (DVT) and/or pulmonary embolism (PE)] was confirmed in a defined location.
	Allowable values: Yes or No/UTD
	 VTE Diagnostic Test – Documentation that a diagnostic test for VTE was performed. Allowable values: Yes or No/UTD VTE Present at Admission - Documentation by a physician/advanced practice nurse/physician assistant (physician/APN/PA) that VTE was diagnosed or suspected on admission. Allowable values: Yes or No/UTD.
Exclusions	. Patients less than 18 years of age
Exclusions	Patients who have a length of stay greater than 120 days
	Patients with Comfort Measures Only documented
	Patients enrolled in clinical trials
	Patients with ICD-9-CM Principal Diagnosis Code of VTE as defined in Appendix A, Table 7.03 or 7.04 Detients with VTE Present at Administra
	 Patients with VTE Present at Admission Patients with reasons for not administering mechanical and pharmacologic prophylaxis
	Patients with reasons for not administering mechanical and pharmacologic prophylaxis Patients without VTE confirmed by diagnostic testing
Exclusion	The patient age in years is equal to the Admission Date minus the Birthdate. The month and day portion of the admission date
Details	and birthdate are used to yield the most accurate age. Patients less than 18 years are excluded. • Length of stay (LOS) in days is equal to the Discharge Date minus the Admission Date. If the LOS is greater than 120 days,
	the patient is excluded.
	Patients are excluded if allowable value 1, 2 or 3 is selected for Comfort Measures Only. Detions are excluded if "Very is calculated for Clinical Trial.
	 Patients are excluded if "Yes" is selected for Clinical Trial. Patients with a Principal ICD-9-CM Diagnosis Code on Table 7.03 or 7.04. are excluded.
	• Patients are excluded if "Yes" is selected for VTE Present at Admission.
	• Patients are excluded if allowable value "3" is selected for VTE Prophylaxis Status.
	Patients are excluded if "No" is selected for VTE Diagnostic Test.
	Patients are excluded if "No" is selected for VTE Confirmed.
Risk Adjustment	No risk adjustment or risk stratification No risk adjustment or risk stratification as intermediate outcome
Stratification	Not Applicable
Type Score	Rate/proportion better quality = lower score
Algorithm	1. Start processing. Run cases that are included in the VTE Initial Patient Population and pass the edits defined in the Transmission Data Processing Flow: Clinical through this measure.
	NOF REVIEW DRAFT—DO NOT CITE OR QUOTE

	0376 Incidence of Potentially Preventable Venous Thromboembolism
	 Check ICD-9-CM Principal Diagnosis Code If the ICD-9-CM Principal Diagnosis Code is on Table 7.03 or 7.04 (VTE, Obstetrics-VTE), the case will proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing. If the ICD-9-CM Principal Diagnosis Code is not on Table 7.03 or 7.04, continue processing and proceed to ICD-9-CM Other Diagnosis Codes Check ICD-9-CM Other Diagnosis Codes If all ICD-9-CM Other Diagnosis Codes are missing or none of them on Table 7.03 or 7.04, the case will proceed to a Measure Category Assignment of B and will not be in the Measure Population. Stop processing. If at least one of the ICD-9-CM Other Diagnosis Codes is on Table 7.03 or 7.04, continue processing and proceed to VTE Present at Admission. Check VTE Present at Admission a. If VTE Present at Admission is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing. If VTE Present at Admission equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the measure Category Assignment of B and will not be in the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing. If VTE Present at Admission equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing. If VTE Present at Admission equals No, continue processing and proceed to Comfort Measures Only.
	 Check Comfort Measures Only a. If Comfort Measures Only is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	 b. If Comfort Measures Only equals 1, 2, or 3, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing. c. If Comfort Measures Only equals 4, continue processing and proceed to Clinical Trial.
	 6. Check Clinical Trial a. If Clinical Trial is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	 b. If Clinical Trial equals Yes, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing. c. If Clinical Trial equals No, continue processing and proceed to VTE Diagnostic Test.
	7. Check VTE Diagnostic Test a. If VTE Diagnostic Test is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	b. If VTE Diagnostic Test equals No, the case will proceed to a Measure Category Assignment of B and will not be in the measure population. Stop processing. c. If VTE Diagnostic Test equals Yes, continue processing and proceed to VTE Confirmed.
	 8. Check VTE Confirmed a. If VTE Confirmed is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop processing.
	b. If VTE Confirmed equals No, the case will proceed to a Measure Category Assignment of B and will not be in the measure population.
	 9. Check VTE Prophylaxis Status a. A if VTE Prophylaxis status is missing, the case will proceed to a Measure Category Assignment of X and will be rejected. Stop Processing.
	 b. If VTE confirmed equals 3, the care will proceed to a Measure Category Assignment of B and will not be in the measure Population. Stop Processing. c. If VTE Prophylaxis Status equals 1, the case will proceed to a Measure Category Assignment of D and will be in the
	Measure Population. d. If VTE Confirmed equals 2, the case will proceed to a Measure Category Assignment of E and will be in the Numerator Population. Stop Processing. Attachment 2zv_VTE6.pdf
Copyright	The Specifications Manual for National Hospital Inpatient Quality Measures (Specifications Manual) is the result of the collaborative efforts of the Centers for Medicare & Medicaid Services (CMS) and The Joint Commission to publish a uniform set of national hospital quality measures. A primary objective of this collaborative effort is to promote and enhance the utility of these measures for all hospitals. No royalty or use fee is required for copying or reprinting this manual, but the following are required as a condition of usage: 1)

0376 Incidence of Potentially Preventable Venous Thromboembolism
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	0419 Documentation of Current Medications in the Medical Record
Steward	Centers for Medicare & Medicaid Services
Description	Percentage of specified visits for patients aged 18 years and older for which the eligible professional attests to documenting a list of current medications to the best of his/her knowledge and ability. This list must include ALL prescriptions, over-the-counters, herbals, vitamin/mineral/dietary (nutritional) supplements AND must contain the medications' name, dosage, frequency and route
Туре	Process
Data Source	Administrative claims, Electronic Clinical Data : Registry Medicare Part B claims data URL NQF 0419 Endorsement Summary 012312 zip file of supporting docuementation sent to H. Bossley & A. Lyzenga via email on 01/23/12 due to path submission error Attachment m130_attachment_partb_detail_line_item_format.pdf
Level	Clinician : Individual, Population : National
Setting	Ambulatory Care : Clinician Office, Behavioral Health/Psychiatric : Outpatient, Dialysis Facility, Home Health, Other, Post Acute/Long Term Care Facility : Nursing Home/Skilled Nursing Facility, Post Acute/Long Term Care Facility : Rehabilitation Clinic, Hospital outpatient
Numerator Statement	ALL MEASURE SPECIFICATION DETAILS REFERENCE THE 2012 PHYSICIAN QUALITY REPORTING SYSTEM MEASURE SPECIFICATION. Eligible professional attests to documenting a list of current medications to the best of his/her knowledge and ability. This list must include ALL prescriptions, over-the counters, herbals, vitamin/mineral/dietary (nutritional) supplements AND must contain the medications' name, dosages, frequency and route NUMERATOR NOTE: By reporting G8427, the eligible professional is attesting the documented current medication information is accurate and complete to the best of his/her knowledge and ability at the time of the patient encounter. This code may also be reported if there is documentation that no medications are currently being taken.
Numerator Details	Time Window: This measure is to be reported at each visit during the 12 month reporting period. Eligible professionals meet the intent of this measure by making a best effort to document a current, complete and accurate medication list during each encounter. There is For the purposes of calculating performance, the Numerator(A) is defined by providers reporting the clinical quality action was performed. For this measure, performing the clinical quality action is numerator HCPCS G8427. Current Medications with Name, Dosage, Frequency and Route Documented G8427: List of current medications (includes prescription, over-the-counter, herbals, vitamin/mineral/dietary [nutritional] supplements) documented by the provider, including drug name, dosage, frequency and route
Statement	ALL MEASURE SPECIFICATION DETAILS REFERENCE THE 2012 PHYSICIAN QUALITY REPORTING SYSTEM MEASURE SPECIFICATION. All visits occurring during the 12 month reporting period for patients aged 18 years and older at the time of the encounter where one or more denominator CPT or HCPCS codes AND any of the 3 numerator HCPCS codes are reported on the claims submission for the encounter. All discussed coding is listed in "2a1.7. Denominator Details" section below.
Denominator Details	Time Window: All visits occurring during the 12 month reporting period for patients aged 18 years and older at the time of the encounter.

	0419 Documentation of Current Medications in the Medical Record
	For the purposes of defining the denominator, the Performance Denominator(PD) is defined by the patient's age, encounter date, denominator CPT or HCPCS codes and the provider reported numerator HCPCS codes described below (G8427, G8430 & G8428).
	Patients aged greater than or equal to 18 years on date of encounter AND
	Patient encounter during the reporting period (CPT or HCPCS): 90801, 90802, 90804, 90805, 90806, 90807, 90808, 90809, 90816, 90817, 90818, 90819, 90821, 90822, 90957, 90958, 90959, 90960, 90962, 90965, 90966, 92002, 92004, 92012, 92014, 92541, 92542, 92543, 92544, 92545, 92547, 92548, 92557, 92567, 92568, 92570, 92585, 92588, 92626, 96116, 96150, 96152, 97001, 97002, 97003, 97004, 97802, 97803, 97804, 98960, 98961, 98962, 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99324, 99325, 99326, 99327, 99328, 99334, 99335, 99336, 99337, 99341, 99342, 99343, 99344, 99345, 99347, 99348, 99349, 99350, G0101, G0108, G0270, G0402, G0438, G0439 AND
	Patient encounters with the following numerator HCPCS Code G8427, G8430, G8428.
	Current Medications with Name, Dosage, Frequency and Route Documented G8427: List of current medications (includes prescription, over-the-counter, herbals, vitamin/mineral/dietary [nutritional] supplements) documented by the provider, including drug name, dosage, frequency and route Current Medications with Dosage not Documented, Patient not Eligible G8430: Provider documentation that patient is not eligible for medication assessment Current Medications with Name, Dosage, Frequency, Route not Documented, Reason not Specified G8428: Current medications (includes prescription, over-the-counter, herbals, vitamin/mineral/dietary [nutritional]
	supplements) with drug name, dosage, frequency and route not documented by the provider, reason not specified
	ALL MEASURE SPECIFICATION DETAILS REFERENCE THE 2012 PHYSICIAN QUALITY REPORTING SYSTEM MEASURE SPECIFICATION. A patient is not eligible or excluded (B) from the performance denominator (PD) if one or more of the following reason(s) exist 1. Patient refuses to participate 2. Patient is in an urgent or emergent medical situation where time is of the essence and to delay treatment would jeopardize the patient's health status 3. Patient cognitively impaired and no authorized representative(s), caregiver(s), and or other healthcare resources are available
	For the purposes of identifying performance exclusions, Denominator Exclusions (B) are defined by providers reporting the
Details	exclusion clinical quality action. For this measure, the clinical exclusion code is numerator HCPCS G8430. Current Medications with Dosages not Documented, Patient not Eligible G8430: Provider documentation that patient is not eligible for medication assessment
Risk	No risk adjustment or risk stratification
	N/A
Stratification	This measure is not stratified. All eligible patients are subject to the same numerator criteria.
51	Rate/proportion better quality = higher score
Algorithm	 This section provides details and formulas to calculate Performance and Denominator Exclusions. PERFORMANCE CALCULATION To calculate provider performance, complete a fraction with the following measure components: Numerator (A), Performance Denominator (PD) and Denominator Exclusions (B). Numerator (A): Number of patients meeting numerator criteria Performance Denominator (PD): Number of patients meeting criteria for denominator inclusion Denominator Exclusions (B): Number of patients with valid exclusions The method of performance calculation is determined by the following: 1) identify the patients who meet the eligibility criteria for the denominator CPT or HCPCS codes and numerator HCPCS codes as listed in "2a1.7. Denominator Details". 2) identify which of those patients meet the numerator criteria (G8427) (A) 3) for those patients who do not meet the numerator criteria, determine whether an appropriate exclusion applies (G8430) (B)
	and subtract those patients from the denominator with the following calculation: Numerator (A)/[Performance Denominator

	0419 Documentation of Current Medications in the Medical Record	
	(PD) - Denominator Exclusions (B)] DENOMINATOR EXCLUSIONS The Exclusion Calculation is: Denominator Exclusions (B)/Performance Denominator (PD) Attachment Calculation for Performance.docx	
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	0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)		
Steward	Agency for Healthcare Research and Quality		
Description	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM codes for deep vein thrombosis or pulmonary embolism in any secondary diagnosis field.		
Туре	Outcome		
Data Source	Administrative claims HCUP State Inpatient Databases (SID). Healthcare Cost and Utilization Project (HCUP). Agency for Healthcare Research and Quality, Rockville, MD. URL http://www.hcup-us.ahrq.gov/sidoverview.jsp Not applicable URL http://qualityindicators.ahrq.gov/Downloads/Software/WinQI/V43/AHRQ%20QI%20Software%20Instructions,%20WinQI.pdf Not applicable		
Level	Facility		
Setting	Hospital/Acute Care Facility		
Numerator Statement	Discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM codes for deep vein thrombosis or pulmonary embolism in any secondary diagnosis field.		
Numerator Details	Time Window: User may specify the time window; generally one calendar year ICD-9-CM Deep vein thrombosis diagnosis codes: 45111 PHLEBITIS AND THROMBOSIS OF FEMORAL VEIN (DEEP) (SUPERFICIAL) 45119 PHLEBITIS AND THROMBOPHLEBITIS OF DEEP VESSEL OF LOWER EXTREMITIES – OTHER 4512 PHLEBITIS AND THROMBOPHLEBITIS OF LOWER EXTREMITIES UNSPECIFIED* 45181 PHLEBITIS AND THROMBOPHLEBITIS OF ILIAC VEIN 4519 PHLEBITIS AND THROMBOPHLEBITIS OF OTHER SITES - OF UNSPECIFIED SITE* 45340 DVT-EMBLSM LOWER EXT NOS (OCT04) 45341 DVT-EMBLSM LOWER EXT NOS (OCT04) 45342 DVT-EMB PROX LOWER EXT (OCT04) 45343 DVT-EMB DISTAL LOWER EXT (OCT04) 4538 OTHER VENOUS EMBOLISM AND THROMBOSIS OF OTHER SPECIFIED VEINS* 4539 OTHER VENOUS EMBOLISM AND THROMBOSIS OF UNSPECIFIED SITE* * Does not apply on or after October 1, 2009. ICD-9-CM Pulmonary embolism diagnosis codes: 4151		

NQF REVIEW DRAFT—DO NOT CITE OR QUOTE Comments due by March 19, 2012 by 6:00pm ET

	0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)		
	PULMONARY EMBOLISM AND INFARCTION		
	41511		
	IATROGENIC PULMONARY EMBOLISM AND INFARCTION		
	41519		
	PULMONARY EMBOLISM AND INFARCTION, OTHER		
	All surgical discharges age 18 and older defined by specific DRGs or MS-DRGs and an ICD-9-CM code for an operating room		
Statement	procedure		
Denominator	Time Window: User may specify the time window; generally one calendar year		
Details	See Patient Safety Indicators Appendices:		
	- Appendix A – Operating Room Procedure Codes		
	- Appendix D – Surgical Discharge DRGs		
	- Appendix E – Surgical Discharge MS-DRGs		
	Link to PSI appendices:		
	http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/TechnicalSpecifications/PSI%20Appendices.pdf		
Exclusions	Exclude cases:		
	- with principal diagnosis of deep vein thrombosis or pulmonary embolism or secondary diagnosis present on admission		
	- where a procedure for interruption of vena cava is the only operating room procedure		
	- where a procedure for interruption of vena cava occurs before or on the same day as the first operating room procedure		
	- MDC 14 (pregnancy, childbirth, and puerperium) - with missing gender (SEX=missing), age (AGE=missing), quarter (DQTR=missing), year (YEAR=missing) or principal		
	diagnosis (DX1=missing), age (AGL=missing), quarter (DQTR=missing), year (TEAR=missing) or principal diagnosis (DX1=missing)		
Exclusion	ICD-9-CM Interruption of vena cava procedure code:		
Details	387		
Details	INTERRUPTION OF VENA CAVA		
Risk	Statistical risk model		
Adjustment	Statistical risk model The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, age (in 5-year age groups), modified CMS DRG, and the AHRQ Comorbidity category. The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Data (SID) for the years 2008, a database consisting of 42 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Age 18 to 24 Age 25 to 29 Age 30 to 34 Age 35 to 39 Age 40 to 44 Age 51 to 49 Age 65 to 74 Age 75 to 79 Age 80 to 84 Age 85+ MDRG 101 MDRG 102 MDRG 103 MDRG 104 MDRG 105 MDRG 107 MDRG 108		

0450 Pc	ostoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)
MDRG	402
MDRG	502
MDRG	503
MDRG	505
MDRG	507
MDRG	508
MDRG	509
MDRG	511
MDRG	514
MDRG	519
MDRG	601
MDRG	602
MDRG	603
MDRG	
	604
MDRG	611
MDRG	701
MDRG	705
MDRG	801
MDRG	802
MDRG	804
MDRG	805
MDRG	806
MDRG	807
MDRG	808
MDRG	811
MDRG	815
MDRG	1001
MDRG	1003
MDRG	1006
MDRG	1101
MDRG	1102
MDRG	1103
MDRG	1104
MDRG	1107
MDRG	1109
MDRG	1201
MDRG	1301
MDRG	1302
MDRG	1303
MDRG	1304
MDRG	1707
MDRG	1708
MDRG	1709
MDRG	1801
MDRG	1802
MDRG	2104
MDRG	2406
MDRG	2407
MDRG	2408
MDRG	2501
MDRG	7701
MDRG	7702
MDC	1

0450 Postoperative Pulmonary Embolism or Deep Vein Thrombosis Rate (PSI 12)	
 MDC 4	
MDC 5	
MDC 7	
MDC 11	
MDC 12	
MDC 16	
MDC 17	
MDC 18	
MDC 21	
MDC 22	
MDC 24	
MDC 25	
TRNSFER Transfer-in	
COMORB CHF	
COMORB VALVE	
COMORB PULMCIRC	
COMORB PERIVASC	
COMORB HTN_C	
COMORB PARA	
COMORB NEURO	
COMORB CHRNLUNG	
COMORB DM	
COMORB HYPOTHY	
COMORB RENLFAIL	
COMORB AIDS	
COMORB LYMPH	
COMORB METS	
COMORB TUMOR	
COMORB OBESE	
COMORB WGHTLOSS	
COMORB BLDLOSS	
COMORB ANEMDEF	
COMORB ALCOHOL	
COMORB DRUG	
COMORB PSYCH	
COMORB DEPRESS	
URL http://qualityindicators.ahrq.gov/Downloads/Software/SAS/V43/Risk%20Adjustment%20Tables%20PSI%204.3.pdf Not	
applicable	
Not applicable	
Rate/proportion better quality = lower score	
 Each indicator is expressed as a rate, is defined as outcome of interest / population at risk or numerator / denominator. The	
AHRQ Quality Indicators (AHRQ QI) software performs six steps to produce the rates. 1) Discharge-level data is used to mark	
inpatient records containing the outcome of interest and 2) the population at risk. For provider indicators, the population at risk	
is also derived from hospital discharge records; for area indicators, the population at risk is derived from U.S. Census data. 3)	
Calculate observed rates. Using output from steps 1 and 2, rates are calculated for user-specified combinations of stratifiers.	
4) Calculate expected rates. Regression coefficients from a reference population database are applied to the discharge	
records and aggregated to the provider or area level. For indicators that are not risk-adjusted, this is the reference population	
rate. 5) Calculate risk-adjusted rate. Use the indirect standardization to account for case-mix. For indicators that are not risk-	
adjusted, this is the same as the observed rate. 6) Calculate smoothed rate. A Univariate shrinkage factor is applied to the	
risk-adjusted rates. The shrinkage estimate reflects a reliability adjustment unique to each indicator URL Not applicable	
http://qualityindicators.ahrq.gov/Downloads/Resources/Publications/2011/QI%20Empirical%20Methods%2005-03-11.pdf	
Not applicable	



Patient Safety Measures: Complications Endorsement Maintenance Project

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Measure Number and Title	Description
#0019: Documentation of medication list in the outpatient record	Percentage of patients having a medication list in the medical record.
#0020: Documentation of allergies and adverse reactions in the outpatient record	Percentage of patients having documentation of allergies and adverse reactions in the medical record.
#0021: Annual monitoring for patients on persistent medications	 The percentage of members 18 years of age and older who received at least 180 treatment days of ambulatory medication therapy for a select therapeutic agent during the measurement year and at least one therapeutic monitoring event for the therapeutic agent in the measurement year. For each product line, report each of the four rates separately and as a total rate. Annual monitoring for members on angiotensin converting enzyme (ACE) inhibitors or angiotensin receptor blockers (ARB) Annual monitoring for members on digoxin Annual monitoring for members on anticonvulsants Total rate (the sum of the four numerators divided by the sum of the four denominators)
#0022: Use of High Risk Medications in the Elderly	 a: Percentage of Medicare members 65 years of age and older who received at least one high-risk medication. b: Percentage of Medicare members 65 years of age and older who received at least two different high-risk medications. For both rates, a lower rate represents better performance.
#0035: Fall Risk Management	 a) Discussing Fall Risk. The percentage of Medicare members 75 years of age and older, or 65–74 years of age with balance or walking problems or a fall in the past 12 months, who were seen by a practitioner in the past 12 months and who discussed falls or problems with balance or walking with their current practitioner. b) Managing Fall Risk. The percentage of Medicare members 65 years of age and older who had a fall or had problems with balance or walking in the past 12 months, who were seen by a practitioner in the past 12 months.
#0097: Medication Reconciliation	Percentage of patients aged 65 years and older discharged from any inpatient facility (e.g. hospital, skilled nursing facility, or rehabilitation facility) and seen within 60 days following discharge in the office by the physician providing on-going care who had a reconciliation of the discharge medications with the current medication list in the medical record documented.
#0101: Falls: Screening for Future Fall Risk	Percentage of patients aged 65 years and older who were screened for fall risk (2 or more falls in the past year or any fall with injury in the past year) at least once within 12 months

Measure Number and Title	Description
#0130: Risk-Adjusted Deep Sternal Wound Infection Rate	Percent of patients aged 18 years and older undergoing isolated CABG who, within 30 days postoperatively, develop deep sternal wound infection involving muscle, bone, and/or mediastinum requiring operative intervention
#0136: Heart Failure (HF): Detailed discharge instructions	Percentage of heart failure patients discharged home with written instructions or educational material given to patient or caregiver at discharge or during the hospital stay addressing all of the following: activity level, diet, discharge medications, follow-up appointment, weight monitoring, and what to do if symptoms worsen.
#0138: Urinary catheter- associated urinary tract infection for intensive care unit (ICU) patients	Percentage of intensive care unit patients with urinary catheter- associated urinary tract infections
#0139: Central line catheter-associated blood stream infection rate for ICU and high-risk nursery (HRN) patients	Percentage of ICU and high-risk nursery patients, who over a certain amount of days acquired a central line catheter-associated blood stream infections over a specified amount of line-days
#0140: Ventilator- associated pneumonia for ICU and high-risk nursery (HRN) patients	Percentage of ICU and HRN patients who over a certain amoint of days have ventilator-associated pneumonia
#0141: Patient Fall Rate	All documented falls, with or without injury, experienced by patients on eligible unit types in a calendar quarter. Reported as Total Falls per 1,000 Patient Days and Unassisted Falls per 1000 Patient Days. (Total number of falls / Patient days) X 1000 Measure focus is safety. Target population is adult acute care inpatient and adult rehabilitation patients.
#0181: Increase in number of pressure ulcers	Percentage of patients who had an increase in the number of pressure ulcers
#0200: Death among surgical inpatients with treatable serious complications (failure to rescue)	Percentage of surgical inpatients with complications of care whose status is death
#0201: Pressure ulcer prevalence	The total number of patients that have hospital-acquired (nosocomial) stage II or greater pressure ulcers on the day of the prevalence study.
#0202: Falls with injury	All documented patient falls with an injury level of minor or greater on eligible unit types in a calendar quarter. Reported as Injury falls per 1000 Patient Days. (Total number of injury falls / Patient days) X 1000 Measure focus is safety. Target population is adult acute care inpatient and adult rehabilitation patients.

Measure Number and Title	Description
#0203: Restraint	Total number of patients that have vest and/or limb restraint (upper or
prevalence (vest and limb	lower body or both) on the day of the prevalence study.
only)	
#0239: Venous	Percentage of patients aged 18 years and older undergoing procedures
Thromboembolism (VTE)	for which VTE prophylaxis is indicated in all patients, who had an order for
Prophylaxis	Low Molecular Weight Heparin (LMWH), Low-Dose Unfractionated
	Heparin (LDUH), adjusted-dose warfarin, fondaparinux or mechanical
	prophylaxis to be given within 24 hours prior to incision time or within 24
	hours after surgery end time.
#0263: Patient Burn	Percentage of ASC admissions experiencing a burn prior to discharge
#0266: Patient Fall	Percentage of ASC admissions experiencing a fall in the ASC.
#0267: Wrong Site, Wrong	Percentage of ASC admissions experiencing a wrong site, wrong side,
Side, Wrong Patient, Wrong	wrong patient, wrong procedure, or wrong implant event.
Procedure, Wrong Implant	
#0281: Urinary infections	This measure is used to assess the number of admissions for urinary tract
(PQI 12)	infection per 100,000 population. See Notes.
#0293: Medication	Percentage of patients transferred to another acute hospitals whose
Information	medical record documentation indicated that medication information was
	communicated to the receiving hospital within 60 minutes of departure
#0298: Central Line Bundle	Percentage of intensive care patients with central lines for whom all
Compliance	elements of the central line bundle are documented and in place.
	The central line bundle elements include:
	•Hand hygiene ,
	Maximal barrier precautions upon insertion
	•Chlorhexidine skin antisepsis
	•Optimal catheter site selection, with subclavian vein as the preferred site
	for non-tunneled catheters in patients 18 years and older
#0200. Curreiter Cite	• Daily review of line necessity with prompt removal of unnecessary lines
#0299: Surgical Site	Percentage of surgical site infections occurring within thirty days after
Infection Rate	the operative procedure if no implant is left in place or with one year if an
	implant is in place in patients who had an NHSN operative procedure
	performed during a specified time period and the infection appears to be related to the operative procedure.
#0301: Surgery patients	Percentage of surgery patients with surgical hair site removal with
with appropriate hair	clippers or depilatory or no surgical site hair removal.
removal	cuppers of depilatory of no surgical site fidit removal.
TETHUVAI	

Measure Number and Title	Description
#0302: Ventilator Bundle	Percentage of intensive care unit patients on mechanical ventilation at time of survey for whom all four elements of the ventilator bundle are documented and in place. The ventilator bundle elements are: •Head of bed (HOB) elevation 30 degrees or greater (unless medically contraindicated); noted on 2 different shifts within a 24 hour period •Daily ""sedation interruption" and daily assessment of readiness to extubate; process includes interrupting sedation until patient follow commands and patient is assessed for discontinuation of mechanical ventilation; Parameters of discontinuation include: resolution of reason for intubation; inspired oxygen content roughly 40%; assessment of patients ability to defend airway after extubation due to heavy sedation; minute ventilation less than equal to 15 liters/minute; and respiratory rate/tidal volume less than or equal to 105/min/L(RR/TV< 105) •SUD (peptic ulcer disease) prophylaxis •DVT (deep venous thrombosis) prophylaxis
#0303: Late sepsis or meningitis in neonates (risk-adjusted)	Percentage of infants born at the hospital, whose birth weight is between 401 and 1500 grams OR whose gestational age is between 22 weeks 0 days and 29 weeks 6 days with late sepsis or meningitis with one or more of the following criteria: Bacterial Pathogen, Coagulase Negative Staphylococcus, Fungal Infection
#0304: Late sepsis or meningitis in Very Low Birth Weight (VLBW) neonates (risk-adjusted)	Percentage of infants born at the hospital, whose birth weight is between 401 and 1500 grams OR whose gestational age is between 22 weeks 0 days and 29 weeks 6 days, who have late sepsis or meningitis, with one or more of the following criteria: Bacterial Pathogen, Coagulase Negative Staphylococcus, Fungal Infection
#0329: All-Cause Readmission Index (risk adjusted)	Overall inpatient 30-day hospital readmission rate.
#0337: Pressure Ulcer Rate (PDI 2)	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code of pressure ulcer in any secondary diagnosis field and ICD-9-CM code of pressure ulcer stage III or IV (or unstagable) in any secondary diagnosis field
#0330: Hospital 30-day, all- cause, risk-standardized readmission rate following heart failure hospitalization for patients 18 and older	The measure estimates a hospital 30-day risk-standardized readmission rate (RSRR), defined as readmission for any cause within 30 days after the date of discharge of the index admission for patients 18 and older discharged from the hospital with a principal diagnosis of heart failure (HF).
#0335: PICU Unplanned Readmission Rate	The total number of patients requiring unscheduled readmission to the ICU within 24 hours of discharge or transfer.
#0336: Review of Unplanned PICU Readmissions	Periodic clinical review of unplanned readmissions to the PICU that occurred within 24 hours of discharge or transfer from the PICU.
#0344: Accidental Puncture or Laceration Rate (PDI 1)	Percent of discharges among cases meeting the inclusion and exclusion rules for the denominator with ICD-9-CM code denoting accidental cut, puncture, perforation, or laceration during a procedure in any secondary

	diagnosis field.
Measure Number and Title	Description
#0345: Accidental Puncture	Percent of discharges among cases meeting the inclusion and exclusion
or Laceration Rate (PSI 15)	rules for the denominator with ICD-9-CM code denoting accidental cut,
	puncture, perforation, or laceration during a procedure in any secondary
	diagnosis field.
#0346: latrogenic	Percent of discharges with ICD-9-CM code for iatrogenic pneumothorax in
Pneumothorax Rate (PSI 6)	any secondary diagnosis field among cases meeting the inclusion and
	exclusion rules for the denominator
#0347: Death Rate in Low-	Percent of discharges with disposition of "deceased" (DISP=20) among
Mortality Diagnosis Related	cases meeting the inclusion and exclusion rules for the denominator
Groups (PSI 2)	
#0348: latrogenic	Percent of discharges among cases meeting the inclusion and exclusion
Pneumothorax Rate (PDI 5)	rules for the denominator with ICD-9-CM code of iatrogenic
	pneumothorax in any secondary diagnosis field
#0349: Transfusion	The count of medical and surgical discharges for patients age greater than
Reaction (PSI 16)	or equal to 18 or in MDC 14 with ICD-9-CM code for transfusion reaction
	in any secondary diagnosis field.
#0350: Transfusion	The count of medical and surgical discharges for patients age less than 18
Reaction (PDI 13)	and not in MDC 14 with ICD-9-CM code for transfusion reaction in any
	secondary diagnosis field.
#351: Death among	Percentage of cases having developed specified complications of care
surgical inpatients with	with an in-hospital death.
serious, treatable	
complications (PSI 4)	
#0352: Failure to Rescue In-	Percentage of patients who died with a complications in the hospital.
Hospital Mortality (risk	
adjusted)	
#0353: Failure to Rescue	Percentage of patients who died with a complication within 30 days from
30-Day Mortality (risk	admission.
adjusted)	
#0362: Foreign Body left	Count of discharges with foreign body left in during procedure in medical
after procedure (PDI 3)	and surgical discharges among patients less than 18 years and not MDC
	14 (pregnancy, childbirth, and puerperium)
#0363: Foreign Body Left	Count of discharges with foreign body left in during procedure in medical
During Procedure (PSI 5)	and surgical discharges among patients 18 years and older or MDC 14
1000 CT D	(pregnancy, childbirth, and puerperium)
#0367: Post operative	Percentage of abdominopelvic surgery cases with reclosure of
Wound Dehiscence (PDI 11)	postoperative disruption of abdominal wall.
#0368: Post operative	Percentage of abdominopelvic surgery cases with reclosure of
Wound Dehiscence (PSI 14)	postoperative disruption of abdominal wall.

Measure Number and Title	Description
#0371: Venous	This measure assesses the number of patients who received venous
Thromboembolism	thromboembolism (VTE) prophylaxis or have documentation why no VTE
Prophylaxis	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. This measure is part of a set of six nationally
	implemented prevention and treatment measures that address VTE (VTE-
	2: ICU VTE Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap
	Therapy, VTE-4: VTE Patients Receiving UFH with Dosages/Platelet Count
	Monitoring, VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-
	6: Incidence of Potentially-Preventable VTE) that are used in The Joint
	Commission's accreditation process.
#0372: Intensive Care Unit	This measure assesses the number of patients who received venous
Venous Thromboembolism	thromboembolism (VTE) prophylaxis or have documentation why no VTE
Prophylaxis	prophylaxis was given the day of or the day after the initial admission (or
	transfer) to the Intensive Care Unit (ICU) or surgery end date for surgeries
	that start the day of or the day after ICU admission (or transfer). This
	measure is part of a set of six prevention and treatment measures that
	address VTE (VTE-1: VTE Prophylaxis, VTE-3: VTE Patients with
	Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with
	Dosages/Platelet Count Monitoring by Protocol, VTE-5: VTE Warfarin
	Therapy Discharge Instructions and VTE-6: VTE Incidence of Potentially-
	Preventable VTE).
#0373: Venous	This measure assesses the number of patients diagnosed with confirmed
Thromboembolism Patients	VTE who received an overlap of Parenteral (intravenous [IV] or
with Anticoagulant Overlap	subcutaneous [subcu]) anticoagulation and warfarin therapy. For patients
Therapy	who received less than five days of overlap therapy, they should be
merupy	discharged on both medications and have a Reason for Discontinuation of
	Overlap Therapy. Overlap therapy should be administered for at least five
	days with an international normalized ratio (INR) greater than or equal to
	2 prior to discontinuation of the parenteral anticoagulation therapy, or
	INR less than 2 but discharged on both medications or have a Reason for
	Discontinuation of Overlap Therapy. This measure is part of a set of six
	prevention and treatment measures that address VTE (VTE-1: VTE
	Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-4: VTE Patients Receiving
	UFH with Dosages/Platelet Count Monitoring, VTE-5: VTE Warfarin
	Therapy Discharge Instructions and VTE-6: Incidence of Potentially-
	Preventable VTE).
#0374: Venous	This measure assesses the number of patients diagnosed with confirmed
Thromboembolism Patients	venous thromboembolism (VTE) who received intravenous (IV)
Recieving Unfractionated	unfractionated heparin (UFH) therapy dosages AND had their platelet
Heparin with	counts monitored using defined parameters such as a nomogram or
Dosages/Platelet Count	protocol. This measure is part of a set of six prevention and treatment
Monitoring by Protocol or	measures that address VTE (VTE-1: VTE Prophylaxis, VTE-2: ICU VTE
Nomogram	Prophylaxis, VTE-3: VTE Patients with Anticoagulation Overlap Therapy,
	VTE-5: VTE Warfarin Therapy Discharge Instructions and VTE-6: Incidence
	of Potentially-Preventable VTE).
	of Potentially-Preventable VTE).

NQF-endorsed® Safety Measures

Measure Number and Title	Description
#0375: Venous	This measure assesses the number of patients diagnosed with confirmed
Thrmoboembolism	VTE that
Warfarin Therapy Discharge	are discharged on warfarin to home, home with home health or home
Instructions	hospice with written discharge instructions that address all four criteria:
	compliance issues, dietary advice, follow-up monitoring, and information
	about the potential for adverse drug reactions/interactions. This measure
	is part of a set of six prevention and treatment measures that address VTE
	(VTE-1: VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients
	with Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH
	with Dosages/Platelet Count Monitoring by Protocol and VTE-6: Incidence
	of Potentially-Preventable VTE).
#0376: Incidence of	This measure assesses the number of patients with confirmed venous
Potentially Preventable	thromboembolism (VTE) during hospitalization (not present at admission)
Venous Thromboembolism	who did not receive VTE prophylaxis between hospital admission and the
	day before the VTE diagnostic testing order date. This measure is part of a
	set of six prevention and treatment measures that address VTE (VTE-1:
	VTE Prophylaxis, VTE-2: ICU VTE Prophylaxis, VTE-3: VTE Patients with
	Anticoagulation Overlap Therapy, VTE-4: VTE Patients Receiving UFH with
	Dosages/Platelet Count Monitoring by Protocol, and VTE-5: VTE Warfarin
	Therapy Discharge Instructions).
#0382: Oncology:	Percentage of patients with a diagnosis of cancer receiving 3D conformal
Radiation Dose Limits to	radiation therapy with documentation in medical record that normal
Normal Tissues	tissue dose constraints were established within five treatment days for a
	minimum of one tissue
#0389: Prostate Cancer:	Percentage of patients with a diagnosis of prostate cancer, at low risk of
Avoidance of Overuse	recurrence, receiving interstitial prostate brachytherapy, OR external
Measure – Isotope Bone	beam radiotherapy to the prostate, OR radical prostatectomy, OR
Scan for Staging Low-Risk Patients	cryotherapy who did not have a bone scan performed at any time since
#0419: Documentation of	diagnosis of prostate cancer Percentage of patients aged 18 years and older with a list of current
Current Medications in the	medications (includes prescription, over-the-counter, herbals,
Medical Record	vitamin/mineral/dietary [nutritional] supplements) documented by the
	provider, including drug name, dosage, frequency and route
#0450: Postoperative	Percent of discharges among cases meeting the inclusion and exclusion
Pulmonary Embolism or	rules for the denominator with ICD-9-CM codes for deep vein thrombosis
Deep Vein Thrombosis Rate	or pulmonary embolism in any secondary diagnosis field.
(PSI 12)	, , , , , , , , , , , , , , , , , , , ,
#0464: Anesthesiology and	Percentage of patients who undergo CVC insertion for whom CVC was
Critical Care: Prevention of	inserted with all elements of maximal sterile barrier technique (cap AND
Catheter-Related	mask AND sterile gown AND sterile gloves AND a large sterile sheet AND
Bloodstream Infections	hand hygiene AND 2% chlorhexidine for cutaneous antisepsis) followed
(CRBSI) – Central Venous	
Catheter (CVC) Insertion	
Protocol	

Measure Number and Title	Description
#0478: Nosocomial Blood Stream Infections in Neonates (NQI #3)	Percentage of qualifying neonates with selected bacterial blood stream infections
#0500: Severe Sepsis and Septic Shock: Management Bundle	Initial steps in the management of the patient presenting with infection (severe sepsis or septic shock)
#0501: Confirmation of Endotracheal Tube Placement	Any time an endotracheal tube is placed into an airway in the Emergency Department or an endotraceal tube is placed by an outside provider and that patient arrives already intubated (EMS or hospital transfer) or when an airway is placed after patients arrives to the ED there should be some method attempted to confirm ETT placement
#0503: Anticoagulation for acute pulmonary embolus patients	Number of acute embolus patients who have orders for anticoagulation (heparin or low-molecular weight heparin) for pulmonary embolus while in the ED.
#0504: Pediatric Weight Documented in Kilograms	Percent of emergency department patients < 18 years of age with a current weight in kilograms documented in the ED record
#505: Thirty-day all-cause risk standardized readmission rate following acute myocardial infarction (AMI) hospitalization.	Hospital-specific 30-day all-cause risk standardized readmission rate following hospitalization for AMI among Medicare beneficiaries aged 65 years or older at the time of index hospitalization.
#506: Thirty-day all-cause risk standardized readmission rate following pneumonia hospitalization.	Hospital-specific 30-day all-cause risk standardized readmission rate following hospitalization for pneumonia among Medicare beneficiaries aged 65 years or older at the time of index hospitalization
#0510: Exposure time reported for procedures using fluoroscopy	Percentage of final reports for procedures using fluoroscopy that include documentation of radiation exposure or exposure time
#0515: Ambulatory surgery patients with appropriate method of hair removal	Percentage of ASC admissions with appropriate surgical site hair removal.
#0520: Drug Education on All Medications Provided to Patient/Caregiver During Episode	Percent of patients or caregivers who were instructed during their episode of home health care on how to monitor the effectiveness of drug therapy, how to recognize potential adverse effects, and how and when to report problems
#0523: Pain Assessment Conducted	Percentage of home health episodes of care in which the patient was assessed for pain, using a standardized pain assessment tool, at start/resumption of care.
#0524: Pain Interventions Implemented During Short Term Episodes Of Care	Percentage of short term home health episodes of care during which pain interventions were included in the physician-ordered plan of care and implemented.
#0526: Timely Initiation of Care	Percent of patients with timely start or resumption of home health care
#0530: Mortality for Selected Conditions	A composite measure of in-hospital mortality indicators for selected conditions.

Measure Number and Title	Description
#0531: Patient Safety for	A composite measure of potentially preventable adverse events for
Selected Indicators	selected indicators
#0532: Pediatric Patient	A composite measure of potentially preventable adverse events for
Safety for Selected	selected pediatric indicators
Indicators	
#0537: Multifactor Fall Risk	Percentage of home health episodes of care in which patients 65 and
Assessment Conducted in	older had a multi-factor fall risk assessment at start/resumption of care.
Patients 65 and Older	
#0538: Pressure Ulcer	Percentage of home health episodes of care in which the physician-
Prevention Included in Plan	ordered plan of care includes interventions to prevent pressure ulcers.
of Care	
#0539: Pressure Ulcer	Percentage of short term home health episodes of care during which
Prevention Implemented	interventions to prevent pressure ulcers were included in the physician-
during Short Term Episodes	ordered plan of care and implemented.
of Care	
#0540: Pressure Ulcer Risk	Percentage of home health episodes of care in which the patient was
Assessment Conducted	assessed for risk of developing pressure ulcers at start/resumption of
	care.
#0541: Proportion of Days	The percentage of patients 18 years and older who met the proportion of
Covered (PDC): 5 Rates by	days covered (PDC) threshold of 80% during the measurement year. A
Therapeutic Category	performance rate is calculated seperately for the following medication
	categories: Beta-Blockers (BB), Angiotensin-Converting Enzyme
	Inhibitor/Angiotensin-Receptor Blocker (ACEI/ARB), Calcium-Channel
	Blockers (CCB), Diabetes Medication, Statins.
	The full detailed measure specifications have also been submitted as a
	separate attachment.
#0542: Adherence to	Medication Possession Ratio (MPR) for chronic medications for individuals
Chronic Medications	over 18 years of age
#0553: Care for Older	Percentage of adults 65 years and older who had a medication review
Adults – Medication Review	
(COA)	
#0554: Medication	Percentage of discharges from January 1 to December 1 of the
Reconciliation Post-	measurement year for patients 65 years of age and older for whom
Discharge (MRP)	medications were reconciled on or within 30 days of discharge.
#0555: Monthly INR	Average percentage of monthly intervals in which Part D beneficiaries
Monitoring for	with claims for warfarin do not receive an INR test during the
Beneficiaries on Warfarin	measurement period
#0556: INR for Beneficiaries	Percentage of episodes with an INR test performed 3 to 7 days after a
Taking Warfarin and	newly-started interacting anti-infective medication for Part D
Interacting Anti-Infective	beneficiaries receiving warfarin
Medications	

Measure Number and Title	Description
#0564: Complications within 30 Days Following Cataract Surgery Requiring Additional Surgical Procedures	Percentage of patients aged 18 years and older with a diagnosis of uncomplicated cataract who had cataract surgery and had any of a specified list of surgical procedures in the 30 days following cataract surgery which would indicate the occurrence of any of the following major complications: retained nuclear fragments, endophthalmitis, dislocated or wrong power IOL, retinal detachment, or wound dehiscence.
#0581: Deep Vein Thrombosis Anticoagulation >= 3 Months	This measure identifies patients with deep vein thrombosis (DVT) on anticoagulation for at least 3 months after the diagnosis
#0582: Diabetes and Pregnancy: Avoidance of Oral Hypoglycemic Agents	This measure identifies pregnant women with diabetes who are not taking an oral hypoglycemic agent.
#0586: Warfarin_PT/ INR Test	This measure identifies the percentage of patients taking warfarin during the measurement year who had at least one PT/INR test within 30 days after the first warfarin prescription in the measurement year
#0593: Pulmonary Embolism Anticoagulation >= 3 Months	This measure identifies patients with pulmonary embolism (PE) on anticoagulation for at least 3 months after the diagnosis.
#0612: Warfarin - INR Monitoring	Percentage of patients taking warfarin with PT/INR monitoring
#0646: Reconciled Medication List Received by Discharged Patients (Inpatient Discharges to Home/Self Care or Any Other Site of Care)	Percentage of patients, regardless of age, discharged from an inpatient facility to home or any other site of care, or their caregiver(s), who received a reconciled medication list at the time of discharge including, at a minimum, medications in the specified categories
#0655: Otitis Media with Effusion: Antihistamines or decongestants – Avoidance of inappropriate use	Percentage of patients aged 2 months through 12 years with a diagnosis of OME were not prescribed or recommended to receive either antihistamines or decongestants
#0656: Otitis Media with Effusion: Systemic corticosteroids – Avoidance of inappropriate use	Percentage of patients aged 2 months through 12 years with a diagnosis of OME who were not prescribed systemic corticosteroids
#0657: Otitis Media with Effusion: Systemic antimicrobials – Avoidance of inappropriate use	Percentage of patients aged 2 months through 12 years with a diagnosis of OME who were not prescribed systemic antimicrobials

Measure Number and Title	Description
#0667: Pulmonary CT	Percent of patients undergoing CT pulmonary angiogram for the
Imaging for Patients at Low	evaluation of possible PE who are at low-risk for PE consistent with
Risk for Pulmonary	guidelines(1) prior to CT imaging. (1) Torbicki A, Perrier A, Konstantinides
Embolism	S, et al. Guidelines on the diagnosis and management of acute pulmonary
	embolism: the Task Force for the Diagnosis and Management of Acute
	Pulmonary Embolism of the European Society of Cardiology (ESC). Eur
	Heart J. 2008 Sep;29(18):2276-315
#0674: Percent of	This measure is based on data from all non-admission MDS 3.0
Residents Experiencing One	assessments of long-stay nursing facility residents which may be annual,
or More Falls with Major	quarterly, significant change, significant correction, or discharge
Injury (Long Stay)	assessment. It reports the percent of residents who experienced one or
	more falls with major injury (e.g., bone fractures, joint dislocations, closed
	head injuries with altered consciousness, and subdural hematoma) in the
	last year (12-month period). The measure is based on MDS 3.0 item
	J1900C, which indicates whether any falls that occurred were associated
	with major injury.
#0675: The Percentage of	This measure is based on data from the MDS 3.0 assessment of short-stay
Residents on a Scheduled	nursing facility residents and reports the percentage of those short-stay
Pain Medication Regimen	residents who can self-report and who are on a scheduled pain
on Admission Who Self-	medication regimen at admission (5-day PPS MDS assessment) and who
Report a Decrease in Pain	report lower levels of pain on their discharge MDS 3.0 assessment or their
Intensity or Frequency	14-day PPS MDS assessment (whichever comes first) when compared
(Short-stay)	with the 5-day PPS MDS assessment.
#0676: Percent of	This measure updates CMS' current QM on pain severity for short-stay
Residents Who Self-Report	residents (people who are discharged within 100 days of admission). This
Moderate to Severe Pain	updated measure is based on data from the Minimum Data Set (MDS 3.0)
(Short-Stay)	14-day PPS assessments. This measure reports the percentage of short-
(0	stay residents with a 14-day PPS assessment during a selected quarter (3
	months) who have reported almost constant or frequent pain and at least
	one episode of moderate to severe pain, or any severe or horrible pain, in
	the 5 days prior to the 14-day PPS assessment.
#0677: Percent of	The proposed long-stay pain measure reports the percent of long-stay
Residents Who Self-Report	residents of all ages in a nursing facility who reported almost constant or
Moderate to Severe Pain	frequent pain and at least one episode of moderate to severe pain or any
(Long-Stay)	severe or horrible pain in the 5 days prior to the MDS assessment (which
	may be an annual, quarterly, significant change or significant correction
	MDS) during the selected quarter.
	Long-stay residents are those who have had at least 100 days of nursing
	facility care. This measure is restricted to the long stay population
	because a separate measure has been submitted for the short-stay
	residents (those who are discharged within 100 days of admission).
	residents (those who are discharged within 100 days of admission).

Measure Number and Title	Description
#0678: Percent of Residents with Pressure Ulcers That Are New or Worsened (Short-Stay)	This measure updates CMS' current QM pressure ulcer measure which currently includes Stage 1 ulcers. The measure is based on data from the MDS 3.0 assessment of short-stay nursing facility residents and reports the percentage of residents who have Stage 2-4 pressure ulcers that are new or have worsened. The measure is calculated by comparing the Stage 2-4 pressure ulcer items on the discharge assessment and the previous MDS assessment (which may be an OBRA admission or 5-day PPS assessment). The quality measure is restricted to the short-stay population defined as those who are discharged within 100 days of admission. The quality measure does not include the long-stay residents who have been in the nursing facility for longer than 100 days. A separate measure has been submitted for them.
#0679: Percent of High Risk Residents with Pressure Ulcers (Long Stay)	CMS currently has this measure in their QMs but it is based on data from MDS 2.0 assessments and it includes Stage 1 ulcers. This proposed measure will be based on data from MDS 3.0 assessments of long-stay nursing facility residents and will exclude Stage 1 ulcers from the definition. The measure reports the percentage of all long-stay residents in a nursing facility with an annual, quarterly, significant change or significant correction MDS assessment during the selected quarter (3-month period) who were identified as high risk and who have one or more Stage 2-4 pressure ulcer(s). High risk populations are those who are comatose, or impaired in bed mobility or transfer, or suffering from malnutrition. Long-stay residents are those who have been in nursing facility care for more than 100 days. This measure is restricted to the population that has long-term needs; a separate pressure ulcer measure is being submitted for short-stay populations. These are defined as having a stay that ends with a discharge within the first 100 days.
#0687: Percent of Residents Who Were Physically Restrained (Long Stay)	The measure is based on data from the MDS 3.0 assessment of long-stay nursing facility residents and reports the percentage of all long-stay residents who were physically restrained. The measure reports the percentage of all long-stay residents in nursing facilities with an annual, quarterly, significant change, or significant correction MDS 3.0 assessment during the selected quarter (3-month period) who were physically restrained daily during the 7 days prior to the MDS assessment (which may be annual, quarterly, significant change, or significant correction MDS 3.0 assessment).

Measure Number and Title	Description
#0689: Percent of	This measure updates CMS' current QM on patients who lose too much
Residents Who Lose Too	weight. This measure captures the percentage of long-stay residents who
Much Weight (Long-Stay)	had a weight loss of 5% or more in the last month or 10% or more in the
	last 6 months who were not on a physician-prescribed weight-loss
	regimen noted on an MDS assessment (which may be an annual,
	quarterly, significant change or significant correction MDS assessment)
	during the selected quarter (3-month period).
	In order to address seasonal variation, the proposed measure uses a two-
	quarter average for the facility. Long-stay residents are those who have
	been in nursing care at least 100 days. The measure is restricted to this
	population, which has long-term care needs, rather than the short-stay
	population who are discharged within 100 days of admission.
#0695: Hospital 30-Day	This measure estimates hospital risk-standardized 30-day readmission
Risk-Standardized	rates following PCI in patients at least 65 years of age. As PCI patients may
Readmission Rates	be readmitted electively for staged revascularization procedures, we will
following Percutaneous	exclude such elective readmissions from the measure. The measure uses
Coronary Intervention (PCI)	clinical data available in the National Cardiovascular Disease Registry
	(NCDR) CathPCI Registry for risk adjustment that has been linked with the
	administrative claims data used to identify readmissions.

Measure Number and Title	Description
Measure Number and Title #0704: Proportion of Patients Hospitalized with AMI that have a Potentially Avoidable Complication (during the Index Stay or in the 30-day Post-Discharge Period)	 Percent of adult population aged 18 – 65 years who were admitted to a hospital with acute myocardial infarction (AMI), were followed for one-month after discharge, and had one or more potentially avoidable complications (PACs). PACs may occur during the index stay or during the 30-day post discharge period (Please reference attached document labeled NQF_AMI_PACS_Risk_Adjustment_2.16.10.xls, tabs labeled CIP_Index PAC_Stays and CIP_PAC_Readmission). We define PACs during each time period as one of three types: (A) PACs during the Index Stay (Hospitalization): (I) PACs related to the anchor condition: The index stay is regarded as having a PAC if during the index hospitalization the patient develops one or more complications such as cardiac arrest, ventricular fibrillation, cardiogenic shock, stroke, coma, acute post-hemorrhagic anemia etc. that may result directly due to AMI or its management. (2) PACs due to Comorbidities: The index stay is also regarded as having a PAC if one or more of the patient's controlled comorbid conditions is exacerbated during the hospitalization (i.e. it was not present on admission). Examples of these PACs are diabetic emergency with hypo- or hyperglycemia, tracheostomy, mechanical ventilation, pneumonia, lung complications gastritis, ulcer, GI hemorrhage etc. (3) PACs suggesting Patient Safety Failures: The index stay is regarded as having a PAC if there are one or more complications related to patient safety issues. Examples of these PACs are septicemia, meningitis, other infections, phlebitis, deep vein thrombosis, pulmonary embolism or any of the CMS-defined hospital acquired conditions (HACs). (B) PACs due to Comorbidities: Readmissions and emergency room visits during the 30-day post discharge period: (1) PACs suggesting Patient Safety Failures: Readmissions or emergency room visits during the 30-day post discharge period: (3) PACs suggesting Patient Safety Failures: Readmissions and emergency room visits
	costs associated with each of these types of PACs during the index hospitalization (tab labeled CIP_Index PAC_Stays) and for readmissions

Measure Number and Title	Description
	and emergency room visits during the 30-day post-discharge period (tab labeled CIP_PAC_Readmission). The information is based on a two-year national commercially insured population (CIP) claims database. The database had 4.7 million covered lives and \$95 billion in "allowed amounts" for claims costs. The database was an administrative claims database with medical as well as pharmacy claims. The two tabs demonstrate the most common PACs that occurred in patients hospitalized with AMI.

Measure Number and Title	Description
Measure Number and Title #0705: Proportion of Patients Hospitalized with Pneumonia that have a Potentially Avoidable Complication (during the Index Stay or in the 30-day Post-Discharge Period)	 Percent of adult population aged 18 – 65 years who were admitted to a hospital with Pneumonia, were followed for one-month after discharge, and had one or more potentially avoidable complications (PACs). PACs may occur during the index stay or during the 30-day post discharge period (Please reference attached document labeled NQF Pneumonia PACs Risk Adjustment 2.16.10.xls, tabs labeled CIP_Index PAC_Stays and CIP_PAC_Readmission). We define PACs during each time period as one of three types: (A) PACs related to the anchor condition: The index stay is regarded as having a PAC if during the index hospitalization the patient develops one or more of the avoidable complications that can result from pneumonia, such as respiratory failure, respiratory insufficiency, pneumothorax, pulmonary collapse, or requires respiratory intubation and mechanical ventilation, incision of pleura, thoracocentesis, chest drainage, tracheostomy etc. (2) PACs due to Comorbidities: The index stay is also regarded as having a PAC if one or more of the patient's controlled comorbid conditions is exacerbated during the hospitalization (i.e. it was not present on admission). Examples of these PACs are diabetic emergency with hypo- or hyperglycemia, stroke, coma, gastritis, ulcer, GI hemorrhage, acute renal failure etc. (3) PACs during the 30-day post discharge period: (1) PACs related to the anchor condition: Readmissions and emergency room visits during the 30-day post discharge period are considered PACs if they are for potentially avoidable complications of pneumonia, such as respiratory failure, respiratory insufficiency, pneumorina, such as respiratory failure, respiratory insufficiency, present on admission). Examples of these PACs are infections, sepsis, phlebitis, deep vein thrombosis, pulmonary embolism or any of the CMS-defined hospital acquired conditions (HACs). (B) PACs during the 30-day post discharge period: (1) PACs related to the anchor condition: Readmissions and
	 pneumonia such as respiratory failure, respiratory insufficiency, pneumonia, respiratory intubation, mechanical ventilation, etc. (2) PACs due to Comorbidities: Readmissions and emergency room visits during the 30-day post discharge period are also considered PACs if they are due to an exacerbation of one or more of the patient's comorbid conditions, such as a diabetic emergency with hypo- or hyperglycemia,
	 stroke, coma, gastritis, ulcer, GI hemorrhage, acute renal failure etc. (3) PACs suggesting Patient Safety Failures: Readmissions or emergency room visits during the 30-day post discharge period are considered PACs if they are due to sepsis, infections, phlebitis, deep vein thrombosis, or for any of the CMS-defined hospital acquired conditions (HACs). The enclosed workbook labeled NQF Pneumonia PACs Risk Adjustment 2.16.10.xls, gives the frequency and costs associated with each of these

Measure Number and Title	Description
	types of PACs during the index hospitalization (tab labeled CIP_Index PAC_Stays) and for readmissions and emergency room visits during the 30-day post-discharge period (tab labeled CIP_PAC_Readmission). The information is based on a two-year national commercially insured population (CIP) claims database. The database had 4.7 million covered lives and \$95 billion in "allowed amounts" for claims costs. The database was an administrative claims database with medical as well as pharmacy claims. The two tabs demonstrate the most common PACs that occurred in patients hospitalized with pneumonia.

Measure Number and Title	Description
Measure Number and Title	 (3) PACs suggesting Patient Safety Failures: Emergency room visits, professional and ancillary services for major infections, deep vein thrombosis, adverse drug events, and other patient safety-related events are considered PACs. The summary tab in the enclosed workbook labeled NQF_Chronic_Care_PACs_Risk_Adjustment_2.9.10.xls gives the overview of the frequency and costs associated with each of these types of PACs for each of the six chronic conditions. Detailed drill-down tabs (e.g. DM IP Stay and DM Prof + OP fac) are also provided in the same workbook for each of the six chronic conditions to highlight high-frequency PACs. The information is based on a two-year, national, commercially insured population (CIP), claims database. The database had 4.7 million covered lives and \$95 billion in "allowed amounts" for claims costs. The database was an administrative claims database with medical as well as pharmacy claims. It is important to note that while the overall frequency of PAC
	hospitalizations are low (for all chronic care conditions summed together, PAC frequency was 6.32% of all PAC occurrences), they amount to over 58% of the PAC medical costs.

Measure Number and Title	Description
#0715: Standardized adverse event ratio for children and adults undergoing cardiac catheterization for congenital heart disease	Ratio of observed to expected clinically important preventable and possibly preventable adverse events, risk-adjusted
#0739: Radiation Dose of Computed Tomography (CT)	The measure has two components. Part A is an outcome measure; Part B is a process measure. Both would work together towards improving quality and allowing hospitals and imaging facilities to conduct ongoing quality improvement. Part A: radiation dose associated with computed tomography (CT) examinations of the head, neck, chest, abdomen/pelvis and lumbar spine, obtained in children and adults. Part B: The proportion of CT examinations where a measure of dose is included in the final medical report
#0740: Participation in a Systematic National Dose Index Registry	Participation in a multi-center, standardized data collection and feedback program that will establish national dose index benchmarks for designated examinations. The registry will eventually provide a comparison of practice or facility dose indices such as CTDIvol and DLP for specified examinations relative to national and regional benchmarks. Data is captured electronically from the images of CT examinations using Digital Imaging and Communications in Medicine (DICOM) standards and the Integrating the Healthcare Enterprise (IHE) Radiation Exposure Monitoring (REM) profile.
#0751: Risk Adjusted Urinary Tract Infection Outcome Measure After Surgery	Risk adjusted, case mix adjusted urinary tract infection outcome measure of adults 18+ years after surgical procedure.

Measure Number and Title	Description
#0752: National Healthcare Safety Network (NHSN) Catheter-associated Urinary Tract Infection (CAUTI) Outcome Measure	 Standardized Infection Ratio (SIR) of healthcare-associated, catheter-associated urinary tract infections (CAUTI) will be calculated among patients in the following patient care locations: Intensive Care Units (ICUs) (excluding patients in neonatal ICUs [NICUs: Level II/III and Level III nurseries]) Specialty Care Areas (SCAs) - adult and pediatric: long term acute care, bone marrow transplant, acute dialysis, hematology/oncology, and solid organ transplant locations other inpatient locations (excluding Level I and Level II nurseries). Data from these locations are reported from acute care general hospitals (including specialty hospitals), freestanding long term acute care hospitals, rehabilitation hospitals, and behavioral health hospitals. Only locations where patients reside overnight are included, i.e., inpatient locations. De.3. If included in a composite, please identify the composite measure
	(title and NQF number if endorsed).
#0753: American College of Surgeons – Centers for Disease Control and Prevention (ACS-CDC) Harmonized Procedure Specific Surgical Site Infection (SSI) Outcome Measure	Prototype measure for the facility adjusted Standardized Infection Ratio (SIR) of deep incisional and organ/space Surgical Site Infections (SSI) at the primary incision site among adult patients aged >= 18 years as reported through the ACS National Surgical Quality Improvement Program (ACS-NSQIP) or CDC National Health and Safety Network (NHSN). Prototype also includes a systematic, retrospective sampling of operative procedures in healthcare facilities. This prototype measure is intended for time-limited use and is proposed as a first step toward a more comprehensive SSI measure or set of SSI measures that include additional surgical procedure categories and expanded SSI risk-adjustment by procedure type. This single prototype measure is applied to two operative procedures, colon surgeries and abdominal hysterectomies, and the measure yields separate SIRs for each procedure.
#0754: National Healthcare Safety Network (NHSN) Central line-associated Bloodstream Infection (CLABSI) Outcome Measure	 Standardized Infection Ratio (SIR) of healthcare-associated, central line-associated bloodstream infections (CLABSI) will be calculated among patients in the following patient care locations: Intensive Care Units (ICUs) Specialty Care Areas (SCAs) - adult and pediatric: long term acute care, bone marrow transplant, acute dialysis, hematology/oncology, and solid organ transplant locations other inpatient locations. (Data from these locations are reported from acute care general hospitals (including specialty hospitals), freestanding long term acute care hospitals, rehabilitation hospitals, and behavioral health hospitals. Only locations where patients reside overnight are included, i.e., inpatient locations.