Surgical Phase of Care Measure 8 – ACS22
Unplanned Reoperation within the 30 Day Postoperative Period

National Quality Strategy (NQS) Domain: Patient Safety

Measure Type: Outcome

2018 OPP MIPS QUALITY OPTIONS FOR INDIVIDUAL MEASURES:
REGISTRY ONLY

DESCRIPTION:
Percentage of patients aged 18 years and older who had any unplanned reoperation within the 30 day postoperative period.

INSTRUCTIONS:
This measure is to be reported each time a procedure for an unplanned reoperation within the 30 day postoperative period is performed during the performance period ending November 30th. There is no diagnosis associated with this measure. This measure may be reported by eligible clinicians who perform the quality actions described in the measure based on the services provided and the measure-specific denominator coding.

Measure Reporting via Registry:
The listed denominator criteria is used to identify the intended patient population. The numerator options included in this specification are used to submit the quality actions allowed by the measure.

DENOMINATOR:
Patients aged 18 years and older undergoing an operative procedure.

Denominator Criteria (Eligible Cases):
All patients aged 18 years and older
AND
Patients undergoing an operative procedure
AND
One of the following CPT codes for the patient encounter during the reporting period (see appendix 1):

NUMERATOR:
Unplanned return to the operating room for a surgical procedure, for any reason, within 30 days of the principal operative procedure.

Numerator Instructions:
INVERSE MEASURE - A lower calculated performance rate for this measure indicates better clinical care or control. The “Performance Not Met” numerator option for this measure is the representation of the better clinical quality or control. Reporting that numerator option will produce a performance rate that trends closer to 0%, as quality increases. For inverse measures a rate of 100% means all of the denominator eligible patients did not receive the appropriate care or were not in proper control.
NUMERATOR NOTE: This measure intent is to capture patients who go back to the operating room within 30 days for a follow-up procedure based on complications of the principal (denominator eligible) operative procedure. Examples: Breast biopsies (19101) with return for re-excisions or insertion of port-a-cath for chemotherapy would not be considered an unplanned return to the operating room for a surgical procedure. If this patient had an open, incisional biopsy of breast tissue (19101) and subsequently had an appendectomy performed this would not be considered an unplanned return to the operating room for a surgical procedure.

The return to the OR may occur at any hospital or surgical facility.

Numerator Quality-Data Coding Options for Reporting Satisfactorily:

Unplanned return to the operating room for a surgical procedure, for complications of the principal operative procedure, within 30 days of the principal operative procedure.

Performance Met: Unplanned return to the operating room for a surgical procedure, for complications of the principal operative procedure, within 30 days of the principal operative procedure.

OR

No return to the operating room for a surgical procedure for complications of the principal operative procedure within 30 days of the principal operative procedure.

Performance Not Met: No return to the operating room for a surgical procedure for complications of the principal operative procedure within 30 days of the principal operative procedure.

RISK ADJUSTMENT:
Risk adjusted in-hospital unplanned re-operation rates will be calculated by adjusting for the variables listed in the following table. Thus, these patient characteristics must be reported.

<table>
<thead>
<tr>
<th>Age</th>
<th>ASA Class</th>
<th>Emergent/Urgent Operation</th>
<th>Functional Status</th>
<th>Wound Class</th>
<th>Preoperative Sepsis</th>
<th>Dyspnea</th>
<th>Ascites</th>
<th>Surgical Approach</th>
</tr>
</thead>
</table>
RATIONAL
This is an adverse surgical outcome, which is often a preventable cause of harm, thus it is important to measure and report. It is feasible to collect the data and produces reliable and valid results about the quality of care. It is useful and understandable to stakeholders. As highlighted earlier, this measure was developed in a collaborative effort by the American College of Surgeons and the American Board of Surgery. This measure addresses the National Quality Strategy Priorities, and was identified by an expert panel of physician providers to be a critical outcome for this procedure. This measure addresses a high-impact condition as it is one of the most common procedures performed in the U.S. The measure aligns well with the intended use. The care settings include Acute Care Facilities/Hospitals. Data are being collected in a clinical registry that has been in existence for over 5 years, with over 4000 current users. Thus, we are requesting consideration of this measure in the “Registry Reporting” option. The level of analysis is the clinician/individual. All populations are included, except children. The measure allows measurement across the person-centered episode of care out to 30 days after the procedure whether an inpatient, outpatient, or readmitted. The measure addresses disparities in care. The risk adjustment is performed with a parsimonious dataset and aims to allow efficient data collection resources and data reporting. Measures have been harmonized when possible.

SUPPORTING EVIDENCE:
A modified-Delphi methodology using an expert panel of surgeons who are Directors of the American Board of Surgery identified this to be a critical outcome for this surgical procedure (Surgeon Specific Registry Report on Project for ABS MOC Part IV. Unpublished study by the American College of Surgeons in conjunction with the American Board of Surgery, 2011).