Perioperative Care Measures Group

This measures group is to be reported **each time** a surgical procedure with the indications for a prophylactic antibiotic (including first or second generation cephalosporin) and VTE prophylaxis is performed for patients aged 18 years and older.

You will need to report G-code G8492 once to indicate your intent to report on the Perioperative Care Measures Group. Once you have reported the G-code, you should begin reporting using one of the patient sample methods listed below.

The following 2011 Physician Quality Reporting System measures are included in the Perioperative Care Measures Group:

#20. Timing of Antibiotic Prophylaxis — Ordering Physician

Measure Description:

Percentage of surgical patients aged 18 years and older undergoing procedures with the indications for prophylactic parenteral antibiotics, who have an order for prophylactic parenteral antibiotic to be given within one hour (if fluoroquinolone or vancomycin, two hours), prior to the surgical incision (or start of procedure when no incision is required)

#21. Selection of Prophylactic Antibiotic — First OR Second Generation Cephalosporin

Measure Description

Percentage of surgical patients aged 18 years and older undergoing procedures with the indications for a first OR second generation cephalosporin prophylactic antibiotic, who had an order for cefazolin OR cefuroxime for antimicrobial prophylaxis

#22. Discontinuation of Prophylactic Antibiotics (Non-Cardiac Procedures)

Measure Description

Percentage of non-cardiac surgical patients aged 18 years and older undergoing procedures with the indications for prophylactic parenteral antibiotics AND who received a prophylactic parenteral antibiotic, who have an order for discontinuation of prophylactic parenteral antibiotics within 24 hours of surgical end time

#23. Venous Thromboembolism (VTE) Prophylaxis (When Indicated in ALL Patients)

Measure Description

Percentage of patients aged 18 years and older undergoing procedures for which VTE prophylaxis is indicated in all patients, who had an order for 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

This measures group can be reported by one of the following patient sample methods:

- 30 Patient Sample Method 30 unique Medicare Part B FFS (fee for service) patients meeting patient sample criteria for the measures group.
- 50% Patient Sample Method via Claims or 80% Patient
 Sample Method via Registry All patients meeting patient
 sample criteria for the measure group during the entire
 reporting period (January 1 through December 31, 2011
 OR July 1 through December 31, 2011). For the 12-month
 reporting period, a minimum of 15 Medicare Part B FFS
 patients must meet the measures group patient sample
 criteria to report satisfactory. For the 6-month reporting
 period, a minimum of 8 Medicare Part B FFS patients
 must meet the measures group patient sample criteria to
 report satisfactory.

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