Measure Information Form for Risk-standardized complication rate (RSCR) eCQM

Project Title:
Measuring Outcomes in Orthopedics Routinely (MOOR)

Project Overview:
The Centers for Medicare & Medicaid Services (CMS) has provided funding to Brigham and Women’s Hospital’s (BWH) Center for Patient Safety, Research, and Practice to: convert two existing NQF-endorsed measures related to orthopedics to eCQMs, develop three new eCQMs related to orthopedics, and develop a new PRO-PM related to Care Goal Achievement following a total hip arthroplasty (THA) and/or total knee arthroplasty (TKA). The cooperative agreement name is Measuring Outcomes in Orthopedics Routinely (MOOR). The cooperative agreement number is: 1V1CMS331637-01-00.

1. Measure Name (Measure Title De.2.)
Risk-standardized complication rate (RSCR) following elective primary total hip arthroplasty (THA) and/or total knee arthroplasty (TKA) electronic clinical quality measure (eCQM).

2. Descriptive Information
This measure estimates the risk-standardized complication rate (RSCR) following elective primary total hip arthroplasty (THA) and/or total knee arthroplasty (TKA) at the clinician group level for adults 18 years and older. The outcome (complication) is defined as any one of the specified complications (Grosso et al., 2012;) occurring from the date of index admission to 90 days’ post date of the index admission (or procedure encounter if the procedure is done on an outpatient basis). Because this is a MIPS measure, the target population is patients 18 and over across all payers.

2.1 Measure Type (NQF Submission Form De.1.)
Outcome

2.2 Brief Description of Measure (NQF Submission Form De.3.)
This measure estimates the RSCR following elective primary THA and/or TKA at the clinician group level for adults 18 years and older. The outcome (complication) is defined as any one of the specified complications (Grosso et al., 2012) occurring from the date of index admission to 90 days post date of the index admission (or procedure encounter if the procedure is done on an outpatient basis). Because this is a MIPS the target population is patients 18 and over across all payers.

References:

2. (2019). "Hospital-level risk-standardized complication rate (RSCR) following elective primary total hip arthroplasty (THA) and/or total knee arthroplasty (TKA)." National Quality Forum. Retrieved July 9, 2019, from: http://www.qualityforum.org/Qps/MeasureDetails.aspx?standardID=1550&print=0&entityTypeID=1

2.3 If Paired or Grouped (NQF Submission Form De.4.)

This measure is not formally paired or grouped with another measure.

3. Measure Specifications

3.1 Measure-specific Web Page (NQF Submission Form S.1.)

This measure is under development and does not have a measure-specific webpage.

3.2 If this is an eCQM (NQF Submission Form S.2a.)

This measure is an eCQM. We developed the measure in the eMeasure Authoring Tool (MAT). Please see the attached zipped output from the MAT.

3.3 Data Dictionary, Code Table, or Value Sets (NQF Submission Form S.2b.)

To identify patients with a qualifying elective primary total hip and/or total knee arthroplasty, we utilized the following Value Set from the Joint Commission.

- Total Hip, Total Knee Replacement: OID # 2.16.840.1.113762.1.4.1029.96

For the denominator exclusions and complications, we developed and published the following value sets in the VSAC:

<table>
<thead>
<tr>
<th>Value Set Name</th>
<th>OID Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonprimary Total Hip, Total Knee Replacement</td>
<td>2.16.840.1.113762.1.4.1206.5</td>
</tr>
<tr>
<td>Fracture Exclusions for Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.2</td>
</tr>
<tr>
<td>Malignant Neoplasm Complications Related to Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.7</td>
</tr>
<tr>
<td>Sepsis Complications Related to Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.4</td>
</tr>
<tr>
<td>Pneumonia Complications Related to Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.6</td>
</tr>
</tbody>
</table>
Surgical Site Bleeding and Other Surgical Site Complications 2.16.840.1.113762.1.4.1206.10
Pulmonary Embolism Complications Related to Hip and Knee Procedures 2.16.840.1.113762.1.4.1206.3
Mechanical Complications Related to Hip and Knee Procedures 2.16.840.1.113762.1.4.1206.1
Periprosthetic Joint Infection/Wound Infection and Other Wound Complications 2.16.840.1.113762.1.4.1206.8
Procedures Resulted from Periprosthetic Joint Infection/Wound Infections 2.16.840.1.113762.1.4.1206.9
Procedures Resulted from Surgical Site Bleeding and Other Surgical Site Complications 2.16.840.1.113762.1.4.1206.11

3.4 For Instrument-Based Measure (NQF Submission Form S.2c)

Not Applicable

3.5 For Endorsement Maintenance (NQF Submission Form S.3.1. and S.3.2.)

Not applicable (this eCQM is not endorsed)

3.6 Numerator Statement (NQF Submission Form S.4.)

The outcome for this measure is any complication occurring during the index admission to 90 days’ following discharge. If the elective primary THA or TKA are done outpatient, the outcome is any complication that occurred during a period of 90 days following procedure. Complications are counted in the measure if they occur during the index hospital admission or within the 90-day period following discharge or procedure (if outpatient). The complication outcome is a dichotomous (yes/no) outcome. If a patient experiences one or more of these complications in the applicable period, the complication outcome for that patient is counted in the measure as a “yes”.

3.7 Numerator Details (NQF Submission Form S.5.)

The complication is a dichotomous outcome (yes for any complication(s); no for no complications). Therefore, if a patient experiences one or more complications, the outcome variable will be coded as a "yes". Complications are counted in the measure if they occur during the index hospital admission or procedure encounter if done as outpatient (and are not present on admission) or within the 90-day post-date of admission or procedure (outpatient).
The complications captured in the numerator are identified during the index admission OR up to 90 days post-date of index admission (inpatient or outpatient), depending on the complication. The follow-up period for complications from date of index admission is as follows:

The follow-up period for AMI, pneumonia, and sepsis/septicemia/shock is seven days from the date of index admission because these conditions are more likely to be attributable to the procedure if they occur within the first week after the procedure. Additionally, analyses indicated a sharp decrease in the rate of these complications after seven days (Grosso et al., 2012).

Death, surgical site bleeding, and pulmonary embolism are followed for 30 days following admission because clinical experts agree these complications are still likely attributable to the hospital performing the procedure during this period and rates for these complications remained elevated until roughly 30 days’ post admission (Grosso et al., 2012).

The measure follow-up period is 90 days after admission for mechanical complications and periprosthetic joint infection/wound infection. Experts agree that mechanical complications and periprosthetic joint infection/wound infections due to the index THA/TKA occur up to 90 days following THA/TKA (Grosso et al., 2012).

The measure counts all complications occurring during an index admission regardless of when they occur. For example, if a patient experiences an AMI on day 10 of the index admission, the measure will count the AMI as a complication, although the specified follow-up period for AMI is seven days. Clinical experts agree with this approach, as such complications likely represent the quality of care provided during the index admission (Grosso et al., 2012).

<table>
<thead>
<tr>
<th>Complication</th>
<th>Time Frame (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute Myocardial Infarction</td>
<td>7</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>7</td>
</tr>
<tr>
<td>Sepsis</td>
<td>7</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>30</td>
</tr>
<tr>
<td>Surgical site bleeding</td>
<td>30</td>
</tr>
<tr>
<td>Death</td>
<td>30</td>
</tr>
<tr>
<td>Wound infection/Periprosthetic joint infection</td>
<td>90</td>
</tr>
<tr>
<td>Mechanical Complication</td>
<td>90</td>
</tr>
</tbody>
</table>

**Table 1:** TJA Complications and follow-up time frames (inpatient and outpatient) (Grosso et al., 2012).

The measure does not count complications that are coded as present on admission (POA) during the index admission; this prevents identifying a condition as a complication of care if it was present on admission for the THA/TKA procedure.
For full list of SNOMED CT/ICD-10 codes defining complications, see the Data Dictionary attached in the Appendix.

See following value sets in the VSAC:

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<th>Steward</th>
</tr>
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<td>Total Hip, Total Knee Replacement</td>
<td>2.16.840.1.113762.1.4.1029.96</td>
<td>The Joint Commission</td>
</tr>
<tr>
<td>Nonprimary Total Hip, Total Knee Replacement</td>
<td>2.16.840.1.113762.1.4.1206.5</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Fracture Exclusions for Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.2</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Malignant Neoplasm Complications Related to Hip and Knee Procedures</td>
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<td>2.16.840.1.113762.1.4.1206.4</td>
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<td>2.16.840.1.113762.1.4.1206.10</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Pulmonary Embolism Complications Related to Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.3</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Mechanical Complications Related to Hip and Knee Procedures</td>
<td>2.16.840.1.113762.1.4.1206.1</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Periprosthetic Joint Infection/Wound Infection and Other Wound Complications</td>
<td>2.16.840.1.113762.1.4.1206.8</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Procedures Resulted from Periprosthetic Joint Infection/Wound Infections</td>
<td>2.16.840.1.113762.1.4.1206.9</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
<tr>
<td>Procedures Resulted from Surgical Site Bleeding and Other Surgical Site Complications</td>
<td>2.16.840.1.113762.1.4.1206.11</td>
<td>Brigham &amp; Women’s Hospital</td>
</tr>
</tbody>
</table>

References:


3.8 Denominator Statement (NQF Submission Form S.6.)
The target population for this eCQM includes adults 18 years of age or older undergoing elective inpatient or outpatient primary THA and/or TKA procedures (all adult patients, all payers).

3.9 Denominator Details (NQF Submission Form S.7.)

To be included in the measure cohort used in public reporting, patients must meet the following inclusion criteria:

1. Aged 18 or older on the date of procedure
2. Having a qualifying elective primary THA/TKA procedure (inpatient or outpatient); elective primary THA/TKA procedures are defined as those procedures without any of the following:
   - Femur, hip, or pelvic fractures coded in the billing diagnosis field from any hospital encounters within 3 months prior to the date of procedure.
   - Partial hip arthroplasty (PHA) procedures (with a concurrent THA/TKA); partial knee arthroplasty procedures are not distinguished by ICD9 codes and are currently captured by the THA/TKA measure
   - Revision procedures with a concurrent THA/TKA
   - Resurfacing procedures with a concurrent THA/TKA
   - Mechanical complication coded in the billing diagnosis field from any hospital encounters within 3 months prior to the date of procedure.
   - Malignant neoplasm of the pelvis, sacrum, coccyx, lower limbs, or bone/bone marrow or a disseminated malignant neoplasm coded in the billing diagnosis field from any hospital encounters within 3 months prior to the date of procedure.
   - Removal of implanted devises/prostheses
   - Transfer status from another acute care facility for the THA/TKA

Patients are eligible for inclusion in the denominator if they had an elective primary THA and/or a TKA when they were 18 or over. Please refer to the Data Dictionary (attached in Appendix) for the SNOMED CT/ICD 10 codes used to define the cohort for each measure. The ICD 10 codes will allow clinician group practices to run the eCQM using a standard report against EHR data (October 2015 and later) to explore trends over time.

3.10 Denominator Exclusions (NQF Includes “Exception” in the “Exclusion” Field) (NQF Submission Form S.8.)

This measure will exclude patients:

1. Who were discharged against medical advice (AMA); or,
2. Who had more than two THA/TKA procedure codes during the index hospitalization.
If a patient has more than one eligible admission in a calendar year after applying these exclusion criteria (in addition to denominator exclusion conditions and concurrent non-primary THA/TKA procedures), we use the first eligible admission and exclude the other eligible admissions in that year.

3.11 Denominator Exclusion Details  
(NQF Includes “Exception” in the “Exclusion” Field)(NQF Submission Form S.9.)

This measure excludes index admissions/procedures for patients:

1. Who were discharged against medical advice (AMA);
   - Rationale: Providers did not have the opportunity to deliver full care and prepare the patient for discharge.
2. Who had more than two THA/TKA procedure codes during the index hospitalization
   - Rationale: Although clinically possible, it is highly unlikely that patients would receive more than two elective THA/TKA procedures in one hospitalization, which may reflect a coding error.

3.12 Stratification Details/Variables (NQF Submission Form S.10.)

N/A

3.13 Risk Adjustment Type (NQF Submission Form S.11.)

Conceptual Model for Relationship Between Social/Behavioral Determinants of Health and Outcomes

Adapted from Dawson AZ, Walker RJ, Gregor C, Egede LE. Relationship between social determinants of health and systolic blood pressure in United States immigrants. *International Journal of Cardiology Hypertension*, Volume 2, August 2019, 100011

**Figure 1: Conceptual Model for Relationship Between Social/Behavioral Determinants of Health and Outcomes**
We adapted Dawson’s conceptual model for relationships between social determinants of health and systolic blood pressure to describe the relationship between antecedents, predisposing, enabling factors, need for healthcare factors and patient outcomes. For this eCQM the outcome we are measuring is complications after THA/TKA. In terms of antecedents, regional variation will be important when looking at complication rates nationally. Rosenberg and colleagues (2016) found that the amount of variability in health outcomes in the U.S. is large even after accounting for differences in population, co-morbidities, and health system factors. African American/Black race is an important outcome antecedent in the total joint replacement population. Pfefferle et al. (2014) noted poorer outcomes for African American patients (particularly African American women) after TKA. They found that African American women under the age of 60 had the greatest incidence of manipulation after TKA due to stiffness and decreased range of motion. Stone et al. (2019) also found that African American patients had longer lengths of stay, more complications (e.g., sepsis, manipulation under anesthesia) and were less likely to be discharged home than Caucasian patients after total joint surgery.

Patient demographics including age, sex and household income may be important predisposing factors of post-surgical outcomes. Basilico and colleagues (2008) found that older age was an important risk factor for complications following total joint replacement surgery. As noted above, younger African American women have the greatest incidence of manipulation post TKA. Dy and colleagues found that younger age and lower income (e.g., Medicaid) increased the risk of undergoing early revision THA. Kremers et al. (2015) did not find associations between marital status and educational attainment and postoperative complications.

English proficiency may be an important enabling factor for patients undergoing THA/TKA. De Oliveira et al. (2015) found that THA/TKA patients had a high prevalence of inadequate health literacy (60%) that may be associated with poor comprehension of discharge instructions and could potentially impact post-surgical outcomes.

Comorbidity and smoking status are important factors that increase the need for healthcare and may contribute to poorer outcomes in patients undergoing total joint replacement. Kremers et al (2015) explored social and behavioral factors in THA/TKA and found that a positive smoking status was associated with higher rates of post-surgical infections. In addition, literature has shown that obesity is associated with higher rates of peri-operative complications, joint and wound infections, mechanical complications, deep vein thrombosis, blood loss, operative time, and need for revision surgery following primary total joint arthroplasty (Haynes et al 2017). Currently, more than one third of Americans are classified as obese (BMI ≥ 30kg/m$^2$) and morbidly obese patients have significantly higher risk of complications noted above and they undergo total knee arthroplasty at an average age of 13 years younger than non-obese patients due to rapid progression of osteoarthritis (Changulani et al 2008).
3.14 Type of Score (NQF Submission Form S.12.)
Rate/proportion

3.15 Interpretation of Score (NQF Submission Form S.13.)
Better quality=lower score

3.16 Calculation Algorithm/Measure Logic (NQF Submission Form S.14.)

**Step 1: Define the Initial Population**

Identify all patients aged 18 years or older, covered by any healthcare payer, who received an elective primary THA and/or TKA within the measurement period.

**Step 2: Define the Denominator**

Apply the denominator exclusion criteria to all the patients from the initial population and determine the denominator population. For the full list of denominator exclusions, please refer to section 3.9, 3.10, and 3.11.

**Step 3: Define the Numerator**

Identify all patients from the denominator who had a complication during the index admission or within the 90-day post-date of admission period/procedure (inpatient or outpatient). The complication is a dichotomous outcome (yes for any complication(s); no for no complications).

**Step 4: Calculate the Complication Rate**

Divide the number of patients in the numerator (step 3) by the number of patients in the denominator (step 2) and multiply by 100. The measure is reported as a percentage: XX out of 100.

The measure estimates clinician group-level risk standardized complication rates (RSCRs) following elective primary THA/TKA using hierarchical logistic regression models. In brief, the approach simultaneously models data at the patient and clinician group levels to account for variance in patient outcomes within and between clinician groups (Normand et al., 2007; Dimick et al., 2010; Krell et al., 2014; MacKenzie et al., 2015). At the patient level, it models the log-odds of a complication occurring within 90 days of the index admission (or procedure if done as outpatient) using age, sex, selected clinical covariates (see Figure 1 above, “Conceptual model for relationship between Social/behavioral determinants of health” for list of covariates) and a clinician group-specific random intercept. At the clinician group level, it models the clinician group-specific intercepts as arising from a normal distribution. The clinician group intercept represents the underlying risk of a complication at the clinician group level, after accounting for
patient risk. The clinician group-specific intercepts are given a distribution to account for the clustering (non-independence) of patients within the same clinician group. If there were no differences among clinician groups, then after adjusting for patient risk, the clinician group intercepts should be identical across all clinician groups.

The RSCR is calculated as the ratio of the number of “predicted” to the number of “expected” episodes with a complication (inpatient or outpatient), multiplied by the national observed complication rate. For each clinician group, the numerator of the ratio is the number of complications within 90 days predicted based on the clinician group’s performance with its observed case mix, and the denominator is the number of complications expected based on the nation’s performance with that clinician group’s case mix. This approach is analogous to a ratio of “observed” to “expected” used in other types of statistical analyses. It conceptually allows for a comparison of a clinician group’s performance given its case mix to an average clinician group’s performance with the same case mix. Thus, a lower ratio indicates lower-than-expected complication rates or better quality, and a higher ratio indicates higher-than-expected complication rates or worse quality.

In summary, the hierarchical logistic regression model is on the patient level and contains the patient characteristics (see Figure 1) as covariates (with fixed regression coefficients for these covariates that are common over all clinician groups) as well as a random effect for clinician group into which the patient’s clinician belongs. The fixed regression coefficients of the risk factors are estimated using maximum likelihood with numerical quadrature to form the marginal likelihood integrated over the random clinician group-specific intercepts (Lange K, 1999). The random clinician group-specific intercepts are then estimated using an empirical Bayes approach (Schall R, 1991).

The “predicted” number of encounters with a complication (the numerator) is calculated by using the coefficients estimated by regressing the risk factors and the clinician group-specific random intercept on the risk of having an encounter with a complication. The estimated clinician group-specific intercept is added to the sum of the estimated regression coefficients multiplied by the patient characteristics. The results are transformed from the logit scale back to the probability scale (using the anti-logit transformation) and summed over all patients attributed to a clinician group to get a predicted value. The “expected” number of encounters with a complication (the denominator) is obtained in the same manner, but a common intercept using all clinician groups in our sample is added. The results are log transformed and summed over all patients in the clinician group to get an expected value. Thus, the risk factors for the patients and their common fixed regression coefficients are used in both the ‘predicted’ and the ‘expected’, but the expected is based solely on the patient characteristics, whereas the predicted includes the clinician group effect. Note, though, the clinician group random effect can be considered the residual clinician group effect after controlling for the patient risk factors. If there are strong clinician group effects after controlling for patient risk factors, then the RSCRs can be much different across clinician groups; if the clinician group effects are weak after controlling for the patient risk factors, then
the RSCRs will be close to one. To assess clinician group performance for each reporting period, we re-estimate the model coefficients using the years of data in that period. This approach has been used in clinical papers by the statistician on our team (Wakeam et al., 2017; Krimphove et al., 2019)

References:


3.17 Sampling (NQF Submission Form S.15.)
N/A. This measure is not based on a sample.

3.18 Survey/Patient-Reported Data (NQF Submission Form S.16.)
N/A. This measure is not based on a survey or patient-reported data.

3.19 Data Source (NQF Submission Form S.17.)
Electronic Health Records data for calculating this eCQM

3.20 Data Source or Collection Instrument (NQF Submission Form S.18.)
Data sources

1. Electronic health record data

References:

2. Suter LG., et al. (2014). “Procedure Specific Complication Measure Updates and Specifications Report: Elective Primary Total Hip Arthroplasty (THA) and/or Total Knee Arthroplasty (TKA) Risk-Standardized Complication Measure (Version 3.0).”

3.21 Data Source or Collection Instrument (Reference) (NQF Submission Form S.19.)

No data collection instrument provided

3.22 Level of Analysis (NQF Submission Form S.20.)

Clinician group

3.23 Care Setting (NQF Submission Form S.21.)

Inpatient/Hospital/Outpatient.

Our eCQM will consider complications that occur in both inpatient and outpatient settings based on the recommendations of our TEP. Our TEP members warned that clinicians may use “observation status” in hospitals to avoid a “readmission” penalty. TEP members also advised to include both inpatient and outpatient THA/TKA procedures as more of these procedures will be done on an outpatient basis.

3.24 Composite Performance Measure (NQF Submission Form S.22.)

N/A. This measure is not a composite performance measure.