

NQF Cancer Standing Committee Orientation Web Meeting

Melissa Mariñelarena, RN, MPA, CPHQ, Senior Director Tara Murphy, MPAP, Project Manger Mauricio Menendez, MS, Project Analyst

February 6, 2018

Welcome

Project Team



Melissa Mariñelarena, RN, MPA, CPHQ, Senior Director



Tara Murphy, MPAPProject Manager



Mauricio Menendez, MS
Project Analyst

Agenda for the Call

- Standing Committee Introductions
- Discuss Fall 2017 Strategic Work
- Overview of NQF, the Consensus Development Process, and Roles of the Standing Committee, co-chairs, NQF staff
- Overview of NQF's portfolio of Cancer measures
- Review of project activities and timelines
- Overview of NQF's measure evaluation criteria
- SharePoint Tutorial
- Measure Worksheet example
- Next steps

Cancer Standing Committee

Karen Fields, MD, Co-Chair
Shelley Fuld Nasso, MPP, Co-Chair
Gregary Bocsi, DO, FCAP
Brent Braveman, PhD, OTR/L, FAOTA
Jennifer Carney, MD
Steven Chen, MD, MBA, FACS
Crawford Clay
Matthew Facktor, MD, FACS
Heidi Floyd
Jennifer Harvey, MD, FACR
Bradford Hirsch, MD

Jette Hogenmiller, PhD, MN, APRN/ARNP, CDE, NTP, TNCC, CEE Joseph Laver, MD, MHA J. Leonard Lichtenfeld, MD, MACP **Stephen Lovell** Jennifer Malin, MD, MACP Jodi Maranchie, MD, FACS Benjamin Movsas, MD Diane Otte, RN, MS, OCN **Beverly Reigle, PhD, RN** David J. Sher, MD, MPH Danielle Ziernicki, PharmD

Strategic Discussion Topics

Strategic Topics Fall 2017

- No measures were ready for Committee review in the Fall 2017 cycle.
- In lieu of measure evaluation, the Committee will convene to discuss broader portfolio topics.
- The Committee will build on the Summer 2017 measure prioritization criteria and measurement gaps work.

Questions?

Overview of NQF, the CDP, and Roles

The National Quality Forum: A Unique Role

Established in 1999, NQF is a nonprofit, nonpartisan, membership-based organization that brings together public- and private-sector stakeholders to reach consensus on healthcare performance measurement. The goal is to make healthcare in the U.S. better, safer, and more affordable.

Mission: To lead national collaboration to improve health and healthcare quality through measurement

- An Essential Forum
- Gold Standard for Quality Measurement
- Leadership in Quality

NQF Activities in Multiple Measurement Areas

Performance Measure Endorsement

- 600+ NQF-endorsed measures across multiple clinical areas
- 15 empaneled expert standing committees

Measure Applications Partnership (MAP)

Advises HHS on selecting measures for 20+ federal programs/Medicaid

National Quality Partners

- Convenes stakeholders around critical health and healthcare topics
- Spurs action: recent examples include antibiotic stewardship, advanced illness care, shared decision making, and opioid stewardship

Measurement Science

- Convenes private- and public-sector leaders to reach consensus on complex issues in healthcare performance measurement
 - » Examples include HCBS, rural issues, telehealth, interoperability, attribution, risk adjustment for social risk factors, diagnostic accuracy, disparities.

Measure Incubator

 Facilitates efficient measure development and testing through collaboration and partnership

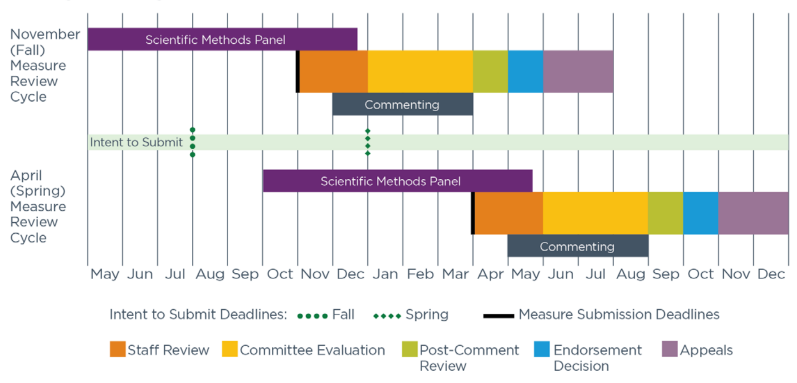
NQF Consensus Development Process (CDP) 6 Steps for Measure Endorsement

- Intent to Submit
- Call for Nominations
- Measure Evaluation
 - New structure/process
 - Newly formed NQF Scientific Methods Panel
 - Measure Evaluation Technical Report
- Public Commenting Period with Member Support
- Measure Endorsement
- Measure Appeals

Measure Review: Two Cycles Per Year

Consensus Development Process:

Two Cycles Every Contract Year



13

15 New Measure Review Topical Areas

	All Cause Admission/ Readmissions	Behavioral Health		All Cause	Ве
Cancer	Cardiovascular	Care Coordination	Infectious Disease	Admission/ Readmissions	Sub
Cost and Resource Use	Endocrine	Eyes, Ears, Nose and Throat Conditions	Palliative and End-of Life Care	Cardiovascular	Ef
Gastrointestinal	Genitourinary	Health and Well Being	Musculoskeletal	Neurology	Exp F
Neurology	Patient Safety	Pediatrics	Perinatal	Pediatrics	Per V
Person and Family- Centered Care	Pulmonary and Critical Care	Renal	Surgery	Primary Care and Chronic Illness	

All Cause Admission/ Readmissions	Behavioral Health & Substance Use	Cancer
Cardiovascular	Cost and Efficiency ^A	Geriatric and Palliative Care ^B
Neurology	Patient Experience & Function	Patient Safety ^c
Pediatrics	Perinatal and Women's Health	Prevention and Population Health ^D
Primary Care and Chronic Illness	Renal	Surgery

[□] Denotes expanded topic area

^A Cost & Efficiency will include efficiency-focused measures from other domains

^B Geriatric & Palliative Care includes pain-focused measures from other domains

^C Patient Safety will include acute infectious disease and critical measures

D Prevention and Population Health is formerly Health and Well Being

Role of the Standing Committee General Duties

- Act as a proxy for the NQF multistakeholder membership
- Serve 2-year or 3-year terms
- Work with NQF staff to achieve the goals of the project
- Evaluate candidate measures against the measure evaluation criteria
- Respond to comments submitted during the review period
- Respond to any directions from the CSAC

Role of the Standing Committee Measure Evaluation Duties

- All members evaluate ALL measures
- Evaluate measures against each criterion
 - Indicate the extent to which each criterion is met and rationale for the rating
- Make recommendations to the NQF membership for endorsement
- Oversee Cancer portfolio of measures
 - Promote alignment and harmonization
 - Identify gaps

Role of the Standing Committee Co-Chairs

- Co-facilitate Standing Committee (SC) meetings
- Work with NQF staff to achieve the goals of the project
- Assist NQF in anticipating questions and identifying additional information that may be useful to the SC
- Keep SC on track to meet goals of the project without hindering critical discussion/input
- Represent the SC at CSAC meetings
- Participate as an SC member

Role of NQF Staff

- NQF project staff works with SC to achieve the goals of the project and ensure adherence to the consensus development process:
 - Organize and staff SC meetings and conference calls
 - Guide the SC through the steps of the CDP and advise on NQF policy and procedures
 - Review measure submissions and prepare materials for Committee review
 - Draft and edit reports for SC review
 - Ensure communication among all project participants (including SC and measure developers)
 - Facilitate necessary communication and collaboration between different NQF projects

18

Role of NQF Staff Communication

- Respond to NQF member or public queries about the project
- Maintain documentation of project activities
- Post project information to NQF's website
- Work with measure developers to provide necessary information and communication for the SC to fairly and adequately evaluate measures for endorsement
- Publish final project report

Role of Methods Panel

- Scientific Methods Panel created to ensure higher-level and more consistent reviews of the scientific acceptability of measures
- The Methods Panel is charged with:
 - Conducting evaluation of complex measures for the Scientific Acceptability criterion, with a focus on reliability and validity analyses and results
 - Serve in advisory capacity to NQF on methodologic issues, including those related to measure testing, risk adjustment, and measurement approaches.
- The Methods Panel review will help inform the Standing Committee's endorsement decision. The Panel will not render endorsement recommendations.

20

Role of the Expert Reviewers

- In 2017, NQF executed a CDP redesign that resulted in restructuring and reducing the number of topical areas as well as a bi-annual measure review process.
- Given these changes, there is a need for diverse yet specific expertise to support longer and continuous engagement from standing committees.

Role of the Expert Reviewers

- The expert reviewer pool serves as an adjunct to NQF standing committees to ensure broad representation and provide technical expertise when needed.
- Expert reviewers will provide expertise as needed to review measures submitted for endorsement consideration by:
 - Replacing an inactive committee member;
 - Replacing a committee member whose term has ended; or
 - Providing expertise that is not currently represented on the committee.
- Expert reviewers may also:
 - Provide comments and feedback on measures throughout the measure review process
 - Participate in strategic discussions in the event no measures are submitted for endorsement consideration

NQF Consensus Development Process (CDP) Measure Evaluation

Complex Measures

- Outcome measures, including intermediate clinical outcomes
- Instrument-based measures (e.g., PRO-PMs)
- Cost/resource use measures
- Efficiency measures (those combining concepts of resource use and quality)
- Composite measures

Non-Complex Measures

- Process measures
- Structural measures
- Previously endorsed complex measures with no changes/updates to the specifications or testing

Questions?

Overview of NQF's Cancer Portfolio

Cancer Portfolio of Measures

- This portfolio contains measures related to Cancer conditions that can be used for accountability and public reporting for all populations and in all settings of care.
- NQF solicits new measures for possible endorsement.
- NQF currently has 29 endorsed measures within the area of cancer. Endorsed measures undergo periodic evaluation to maintain endorsement – "maintenance".

Cancer Portfolio of Measures

Screening			
NQF Number	Measure Title		
0508	Diagnostic Imaging: Inappropriate Use of "Probably Benign" Assessment Category in Screening Mammograms		

Diagnosis			
NQF Number	Measure Title		
0377	Hematology: Myelodysplastic Syndrome (MDS) and Acute Leukemias: Baseline Cytogenetic Testing Performed on Bone Marrow		
0379	Hematology: Chronic Lymphocytic Leukemia (CLL): Baseline Flow Cytometry		
0391	Breast Cancer Resection Pathology Reporting- pT category (primary tumor) and pN category (regional lymph nodes) with histologic grade		
0392	Colorectal Cancer Resection Pathology Reporting- pT category (primary tumor) and pN category (regional lymph nodes) with histologic grade		
1853	Radical Prostatectomy Pathology Reporting		
1854	Barrett's Esophagus		
1855	Quantitative HER2 evaluation by IHC uses the system recommended by the ASCO/CAP guidelines		

27

Cancer Portfolio of Measures (continued)

Treatment/Early Disease			
NQF Number	Measure Title		
0219	Post breast conservation surgery irradiation		
0220	Adjuvant hormonal therapy		
0378	Hematology: Myelodysplastic Syndrome (MDS): Documentation of Iron Stores in Patients Receiving Erythropoietin Therapy		
0380	Hematology: Multiple Myeloma: Treatment with Bisphosphonates		
0387*	Hormonal therapy for stage IC through IIIC, ER/PR positive breast cancer		
0559*	Combination chemotherapy is considered or administered within 4 months (120 days) of diagnosis for women under 70 with AJCC T1cN0M0, or Stage IB - III hormone receptor negative breast cancer		
1858*	Trastuzumab administered to patients with AJCC stage I (T1c) – III and human epidermal growth factor receptor 2 (HER2) positive breast cancer who receive adjuvant chemotherapy		

^{*}Measure included in more than one category

Cancer Portfolio of Measures (continued)

Treatment/Advanced Disease			
NQF Number	Measure Title		
0223	Adjuvant chemotherapy is recommended or administered within 4 months (120 days) of diagnosis to patients under the age of 80 with AJCC III (lymph node positive) colon cancer		
0385	Oncology: Chemotherapy for AJCC Stage III Colon Cancer Patients		
0387*	Hormonal therapy for stage IC through IIIC, ER/PR positive breast cancer		
0559*	Combination chemotherapy is considered or administered within 4 months (120 days) of diagnosis for women under 70 with AJCC T1cN0M0, or Stage IB - III hormone receptor negative breast cancer		
1822	External Beam Radiotherapy for Bone Metastases		
1858*	Trastuzumab administered to patients with AJCC stage I (T1c) – III and human epidermal growth factor receptor 2 (HER2) positive breast cancer who receive adjuvant chemotherapy		
1859	KRAS gene mutation testing performed for patients with metastatic colorectal cancer who receive anti-epidermal growth factor receptor monoclonal antibody therapy		
1860	Patients with metastatic colorectal cancer and KRAS gene mutation spared treatment with anti-epidermal growth factor receptor monoclonal antibodies		

^{*}Measure included in more than one category

Cancer Portfolio of Measures (continued)

Follow-up Care			
NQF Number	Measure Title		
0389	Prostate Cancer: Avoidance of Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients		
O390 Prostate Cancer: Adjuvant Hormonal Therapy for High Risk Prostate Cancer Patients			

Other			
NQF Number	Measure Title		
0381	Oncology: Treatment Summary Communication – Radiation Oncology		
0382	Oncology: Radiation Dose Limits to Normal Tissues		
0383	Oncology: Plan of Care for Pain – Medical Oncology and Radiation Oncology (paired with 0384)		
0384	Oncology: Pain Intensity Quantified – Medical Oncology and Radiation Oncology (paired with 0383)		
0386	Oncology: Cancer Stage Documented		
2930	Febrile Neutropenia Risk Assessment Prior to Chemotherapy		
2963	Prostate Cancer: Avoidance of Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients		

Activities and Timeline Fall 2017 Review Cycle

*All times ET

Activity	Date
Committee Orientation Web Meeting	Tuesday, February 6, 11:00am-1:00pm ET
(rescheduled)	
Committee Strategic Web Meeting	Monday, February 12, 12:00-2:00pm ET

Activities and Timeline Spring 2018 Review Cycle

*All times ET

Activity	Date
Measure Submission Deadline	April 16, 2018
Commenting & member support period on submitted measures opens	Monday, May 7, 2018
Measure Evaluation Web Meeting #1	Tuesday, July 10, 2018, 12-2pm ET
Measure Evaluation Web Meeting #2	Friday, July 13, 2018, 11am-1pm ET
Measure Evaluation Web Meeting #3	Monday, July 16, 2018, 1-3pm ET
Draft Report Posted for Public Comment	August 7-September 5, 2018
Post Draft Report Comment Call	Wednesday, September 26, 2018, 2-4pm ET
CSAC Review Period	October 19-November 8, 2018
Appeals Period	November 13-December 12, 2018

Questions?

Measure Evaluation Criteria Overview

NQF Measure Evaluation Criteria for Endorsement

NQF endorses measures for accountability applications (public reporting, payment programs, accreditation, etc.) as well as quality improvement.

- Standardized evaluation criteria
- Criteria have evolved over time in response to stakeholder feedback
- The quality measurement enterprise is constantly growing and evolving – greater experience, lessons learned, expanding demands for measures – the criteria evolve to reflect the ongoing needs of stakeholders.

Major Endorsement Criteria (page 28)

- Importance to measure and report: Goal is to measure those aspects with greatest potential of driving improvements; if not important, the other criteria are less meaningful (must-pass)
- Reliability and Validity-scientific acceptability of measure properties: Goal is to make valid conclusions about quality; if not reliable and valid, there is risk of improper interpretation (must-pass)
- Feasibility: Goal is to, ideally, cause as little burden as possible;
 if not feasible, consider alternative approaches
- Usability and Use: Goal is to use for decisions related to accountability and improvement; if not useful, probably do not care if feasible
- Comparison to related or competing measures

Criterion #1: Importance to Measure and Report (page 30-39)

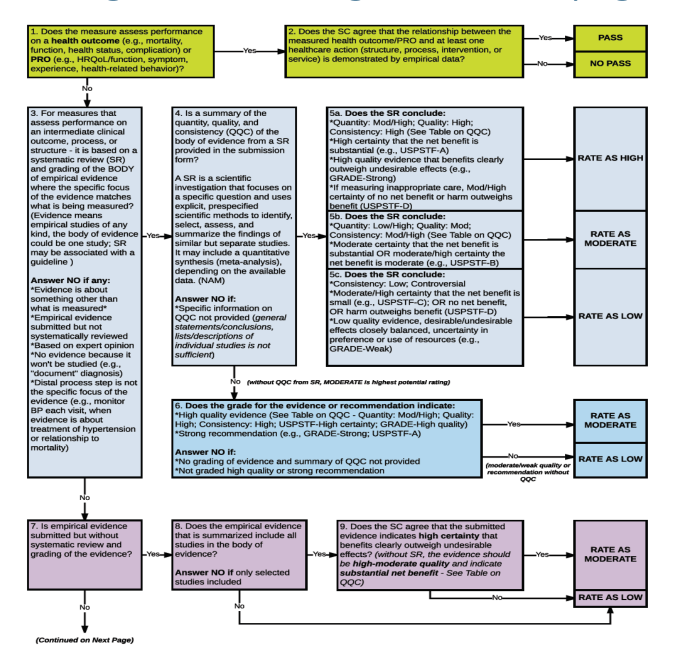
- 1. Importance to measure and report Extent to which the specific measure focus is evidence-based and important to making significant gains in healthcare quality where there is variation in or overall less-than-optimal performance.
 - 1a. Evidence: the measure focus is evidence-based
 - **1b.** Opportunity for Improvement: demonstration of quality problems and opportunity for improvement, i.e., data demonstrating considerable variation, or overall less-than-optimal performance, in the quality of care across providers; and/or disparities in care across population groups
 - 1c. Quality construct and rationale (composite measures only)

Subcriterion 1a: Evidence (page 31-37)

Outcome measures

- Empirical data demonstrate a relationship between the outcome and at least one healthcare structure, process, intervention, or service. If not available, wide variation in performance can be used as evidence, assuming the data are from a robust number of providers and results are not subject to systematic bias.
- Structure, process, intermediate outcome measures
 - The quantity, quality, and consistency of the body of evidence underlying the measure should demonstrate that the measure focuses on those aspects of care known to influence desired patient outcomes
 - » Empirical studies (expert opinion is not evidence)
 - » Systematic review and grading of evidence
 - Clinical Practice Guidelines variable in approach to evidence review
- For measures derived from patient (or family/parent/etc.) report
 - Evidence should demonstrate that the target population values the measured outcome, process, or structure and finds it meaningful.
 - Current requirements for structure and process measures also apply to patient-reported structure/process measures.

Rating Evidence: Algorithm #1 - page 34



Criterion #1: Importance to measure and report Criteria emphasis is different for new vs. maintenance measures

New measures	Maintenance measures
 Evidence – Quantity, quality, consistency (QQC) Established link for process measures with outcomes 	DECREASED EMPHASIS: Require measure developer to attest evidence is unchanged from last evaluation; Standing Committee to affirm no change in evidence IF changes in evidence, the Committee will evaluate as for new measures
 Gap – opportunity for improvement, variation, quality of care across providers 	INCREASED EMPHASIS: data on current performance, gap in care and variation

Criterion #2: Reliability and Validity—Scientific Acceptability of Measure Properties (page 39 -48)

Extent to which the measure, <u>as specified</u>, produces consistent (reliable) and credible (valid) results about the quality of healthcare delivery

2a. Reliability (must-pass)

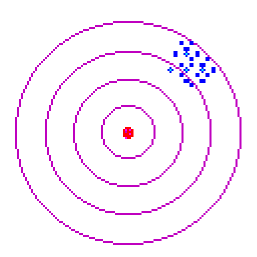
- 2a1. Precise specifications including exclusions
- 2a2. Reliability testing—data elements or measure score

2b. Validity (must-pass)

- 2b1. Validity testing—data elements or measure score
- 2b2. Justification of exclusions—relates to evidence
- 2b3. Risk adjustment—typically for outcome/cost/resource use
- 2b4. Identification of differences in performance
- 2b5. Comparability of data sources/methods
- 2b6. Missing data

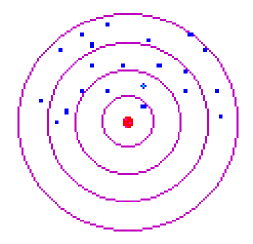
Reliability and Validity (page 40)

Assume the center of the target is the true score...



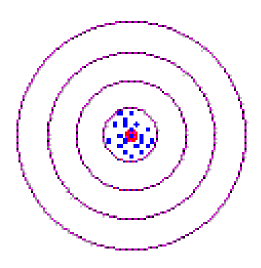
Reliable Not Valid

Consistent, but wrong



Neither Reliable Nor Valid

Inconsistent & wrong



Both Reliable And Valid

Consistent & correct

Evaluating Scientific Acceptability – Key Points (page 41)

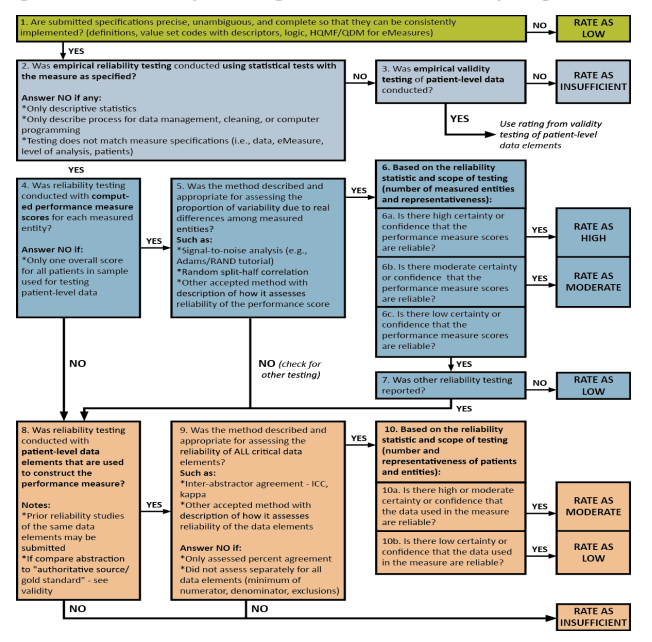
Empirical analysis to demonstrate the reliability and validity of the *measure as specified,* including analysis of issues that pose threats to the validity of conclusions about quality of care such as exclusions, risk adjustment/stratification for outcome and resource use measures, methods to identify differences in performance, and comparability of data sources/methods.

43

Reliability Testing Key points - page 42

- Reliability of the *measure score* refers to the proportion of variation in the performance scores due to systematic differences across the measured entities in relation to random variation or noise (i.e., the precision of the measure).
 - Example Statistical analysis of sources of variation in performance measure scores (signal-to-noise analysis)
- Reliability of the data elements refers to the repeatability/reproducibility
 of the data and uses patient-level data
 - Example –inter-rater reliability
- Consider whether testing used an appropriate method and included adequate representation of providers and patients and whether results are within acceptable norms
- Algorithm #2

Rating Reliability: Algorithm #2 - page 43



Validity testing (pages 44 - 49) Key points – page 47

Empirical testing

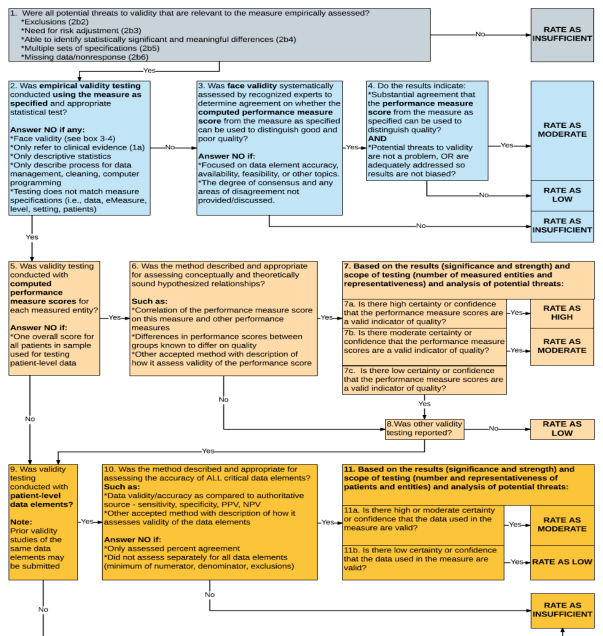
- Measure score assesses a hypothesized relationship of the measure results to some other concept; assesses the correctness of conclusions about quality
- Data element assesses the correctness of the data elements compared to a "gold standard"

Face validity

- Subjective determination by experts that the measure appears to reflect quality of care
 - » Empirical validity testing is expected at time of maintenance review; if not possible, justification is required.
 - » Requires systematic and transparent process, by identified experts, that explicitly addresses whether performance scores resulting from the measure as specified can be used to distinguish good from poor quality. The degree of consensus and any areas of disagreement must be provided/discussed.

46

Rating Validity: Algorithm #3 – page 48



Threats to Validity

- Conceptual
 - Measure focus is not a relevant outcome of healthcare or not strongly linked to a relevant outcome
- Unreliability
 - Generally, an unreliable measure cannot be valid
- Patients inappropriately excluded from measurement
- Differences in patient mix for outcome and resource use measures
- Measure scores that are generated with multiple data sources/methods
- Systematic missing or "incorrect" data (unintentional or intentional)

48

Criterion #2: Scientific Acceptability

N	ew measures	Maintenance measures
•	Measure specifications are precise with all information needed to implement the measure	NO DIFFERENCE: Require updated specifications
•	Reliability Validity (including risk adjustment)	DECREASED EMPHASIS : If prior testing adequate, no need for additional testing at maintenance with certain exceptions (e.g., change in data source, level of analysis, or setting)
		Must address the questions regarding use of social risk factors in risk-adjustment approach

Criterion #3: Feasibility (page 49) Key Points – page 50

Extent to which the required data are readily available, retrievable without undue burden, and can be implemented for performance measurement.

3a: Clinical data generated during care process

3b: Electronic sources

3c: Data collection strategy can be implemented

Criterion #4: Usability and Use (page 50) Key Points – page 51

Extent to which potential audiences (e.g., consumers, purchasers, providers, policymakers) are using or could use performance results for both accountability and performance improvement to achieve the goal of high-quality, efficient healthcare for individuals or populations.

Use (4a) Now must-pass for maintenance measures

4a1: Accountability and Transparency: Performance results are used in at least one accountability application within three years after initial endorsement and are publicly reported within six years after initial endorsement.

4a2: Feedback by those being measured or others: Those being measured have been given results and assistance in interpreting results; those being measured and others have been given opportunity for feedback; the feedback has been considered by developers.

Usability (4b)

4b1: Improvement: Progress toward achieving the goal of high-quality, efficient healthcare for individuals or populations is demonstrated.

4b2: Benefits outweigh the harms: The benefits of the performance measure in facilitating progress toward achieving high-quality, efficient healthcare for individuals or populations outweigh evidence of unintended negative consequences to individuals or populations (if such evidence exists).

51

Criteria #3-4: Feasibility and Usability and Use

New measures	Maintenance measures		
Feasibility			
Measure feasible, including eMeasure feasibility assessment	NO DIFFERENCE: Implementation issues may be more prominent		
Usability and Use			
Use: used in accountability applications and public reporting	INCREASED EMPHASIS: Much greater focus on measure use and		
Usability: impact and unintended consequences	usefulness, including both impact and unintended consequences		

Criterion #5: Related or Competing Measures (page 51-52)

If a measure meets the four criteria <u>and</u> there are endorsed/new related measures (same measure focus <u>or</u> same target population) or <u>competing</u> measures (both the same measure focus <u>and</u> same target population), the measures are compared to address harmonization and/or selection of the best measure.

- 5a. The measure specifications are harmonized with related measures OR the differences in specifications are justified.
- 5b. The measure is superior to competing measures (e.g., is a more valid or efficient way to measure) OR multiple measures are justified.

Updated guidance for measures that use ICD-10 coding: Fall 2017 and 2018

- Gap can be based on literature and/or data based on ICD-9 or ICD-10 coding
- Submit updated ICD-10 reliability testing if available; if not, testing based on ICD-9 coding will suffice
- Submit updated validity testing
 - Submit updated empirical validity testing on the ICD-10 specified measure, if available
 - OR face validity of the ICD-10 coding scheme plus face validity of the measure score as an indicator of quality
 - OR face validity of the ICD-10 coding scheme plus score-level empirical validity testing based on ICD-9 coding
 - OR face validity of the ICD-10 coding scheme plus data element level validity testing based on ICD-9 coding, with face validity of the measure score as an indicator of quality due at annual update

54

Evaluation Process

- Preliminary analysis (PA): To assist the Committee evaluation of each measure against the criteria, NQF staff and Methods Panel (if applicable) will prepare a PA of the measure submission and offer preliminary ratings for each criterion.
 - The PA will be used as a starting point for the Committee discussion and evaluation
 - Methods Panel will complete review of Scientific Acceptability criterion for complex measures
- Individual evaluation: Each Committee member conducts an in-depth evaluation on all measures (responses collected via SurveyMonkey
 - Each Committee member will be assigned a subset of measures for which they will serve as lead discussant in the evaluation meeting.

Evaluation Process

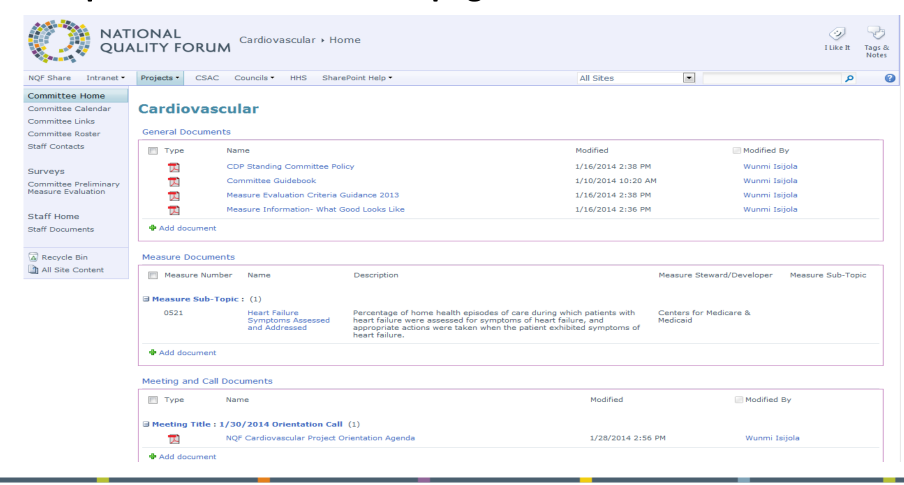
- Measure evaluation and recommendations at the inperson/web meeting: The entire Committee will discuss and rate each measure against the evaluation criteria and make recommendations for endorsement.
- Staff will prepare a draft report detailing the Committee's discussion and recommendations
 - This report will be released for a 30-day public and member comment period
- Post-comment call: The Committee will re-convene for a post-comment call to discuss comments submitted
- Final endorsement decision by the CSAC
- Appeals (if any)

Questions?

http://share.qualityforum.org/Projects/Cancer/SitePages/Home.aspx

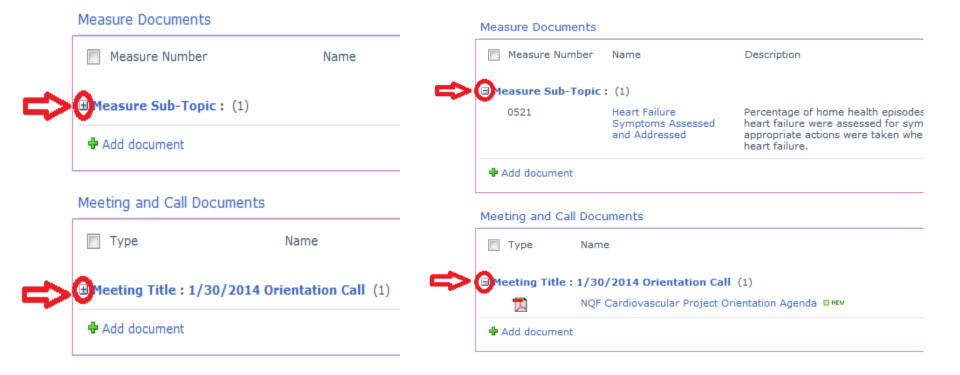
- Accessing SharePoint
- Standing Committee Policy
- Standing Committee Guidebook
- Measure Document Sets
- Meeting and Call Documents
- Committee Roster and Biographies
- Calendar of Meetings

Sample screen shot of homepage:



60

- Please keep in mind:
- + and signs :



Measure Worksheet and Measure Information

Measure Worksheet

- Preliminary analysis, including eMeasure Technical Review if needed, and preliminary ratings
- Member and public comments
- Information submitted by the developer
 - Evidence and testing attachments
 - Spreadsheets
 - Additional documents

Next Steps

Activities and Timeline Fall 2017 Review Cycle

*All times ET

Activity	Date
Committee Orientation Web Meeting	Tuesday, February 6, 11:00am-1:00pm ET
(rescheduled)	
Committee Strategic Web Meeting	Monday, February 12, 12:00-2:00pm ET

Activities and Timeline Spring 2018 Review Cycle

*All times ET

Activity	Date
Measure Submission Deadline	April 16, 2018
Commenting & member support period on submitted measures opens	Monday, May 7, 2018
Measure Evaluation Web Meeting #1	Tuesday, July 10, 2018, 12-2pm ET
Measure Evaluation Web Meeting #2	Friday, July 13, 2018, 11am-1pm ET
Measure Evaluation Web Meeting #3	Monday, July 16, 2018, 1-3pm ET
Draft Report Posted for Public Comment	August 7-September 5, 2018
Post Draft Report Comment Call	Wednesday, September 26, 2018, 2-4pm ET
CSAC Review Period	October 19-November 8, 2018
Appeals Period	November 13-December 12, 2018

Project Contact Info

- Email: cancerm@qualityforum.org
- NQF Phone: 202-783-1300
- Project page:
 http://www.qualityforum.org/Project Pages/cancer.aspx
- SharePoint site: <u>http://share.qualityforum.org/Projects/cancer/SitePages/Home.aspx</u>

Questions?

