This form contains the measure information submitted by stewards. Blank fields indicate no information was provided. Attachments also may have been submitted and are provided to reviewers. The subcriteria and most of the footnotes from the evaluation criteria are provided in Word comments within the form and will appear if your cursor is over the highlighted area. Hyperlinks to the evaluation criteria and ratings are provided in each section.

TAP/Workgroup (if utilized): Complete all yellow highlighted areas of the form. Evaluate the extent to which each subcriterion is met. Based on your evaluation, summarize the strengths and weaknesses in each section.

Note: If there is no TAP or workgroup, the SC also evaluates the subcriteria (yellow highlighted areas).

Steering Committee: Complete all pink highlighted areas of the form. Review the workgroup/TAP assessment of the subcriteria, noting any areas of disagreement; then evaluate the extent to which each major criterion is met; and finally, indicate your recommendation for the endorsement. Provide the rationale for your ratings.

Evaluation ratings of the extent to which the criteria are met
C = Completely (unquestionably demonstrated to meet the criterion)
P = Partially (demonstrated to partially meet the criterion)
M = Minimally (addressed BUT demonstrated to only minimally meet the criterion)
N = Not at all (NOT addressed; OR incorrectly addressed; OR demonstrated to NOT meet the criterion)
NA = Not applicable (only an option for a few subcriteria as indicated)

(for NQF staff use) NQF Review #: 1457  NQF Project: End Stage Renal Disease

<table>
<thead>
<tr>
<th>MEASURE DESCRIPTIVE INFORMATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>De.1 Measure Title: Access-Related Bacteremia (rate)</td>
</tr>
<tr>
<td>De.2 Brief description of measure: Overall access-related bacteremia: Six-month rolling average rate of access-related bacteremia with IV antibiotic therapy, among adult chronic hemodialysis (HD) patients (Express as: rate per 1000 HD patient days)</td>
</tr>
<tr>
<td>Specific access types: Six-month rolling average rate of fistula/graft/catheter-related bacteremia with IV antibiotic therapy, among adult chronic hemodialysis (HD) patients using a fistula/graft/catheter for HD access (Express as: rate per 1000 fistula/graft/catheter patient days)</td>
</tr>
<tr>
<td>1.1-2 Type of Measure: Process</td>
</tr>
<tr>
<td>De.3 If included in a composite or paired with another measure, please identify composite or paired measure</td>
</tr>
<tr>
<td>De.4 National Priority Partners Priority Area: Population health</td>
</tr>
<tr>
<td>De.5 IOM Quality Domain: Safety</td>
</tr>
<tr>
<td>De.6 Consumer Care Need: Living with illness</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CONDITIONS FOR CONSIDERATION BY NQF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Four conditions must be met before proposed measures may be considered and evaluated for suitability as voluntary consensus standards:</td>
</tr>
<tr>
<td>A. The measure is in the public domain or an intellectual property (measure steward agreement) is signed. Public domain only applies to governmental organizations. All non-government organizations must sign a measure steward agreement even if measures are made publicly and freely available.</td>
</tr>
<tr>
<td>A.1 Do you attest that the measure steward holds intellectual property rights to the measure and the right to use aspects of the measure owned by another entity (e.g., risk model, code set)? Yes</td>
</tr>
<tr>
<td>A.2 Indicate if Proprietary Measure (as defined in measure steward agreement);</td>
</tr>
<tr>
<td>A.3 Measure Steward Agreement: Government entity and in the public domain - no agreement necessary</td>
</tr>
<tr>
<td>A.4 Measure Steward Agreement attached:</td>
</tr>
</tbody>
</table>

Rating: C=Completely; P=Partially; M=Minimally; N=Not at all; NA=Not applicable
B. The measure owner/steward verifies there is an identified responsible entity and process to maintain and update the measure on a schedule that is commensurate with the rate of clinical innovation, but at least every 3 years. Yes, information provided in contact section

C. The intended use of the measure includes both public reporting and quality improvement.

►Purpose: Public reporting, Internal quality improvement

D. The requested measure submission information is complete. Generally, measures should be fully developed and tested so that all the evaluation criteria have been addressed and information needed to evaluate the measure is provided. Measures that have not been tested are only potentially eligible for a time-limited endorsement and in that case, measure owners must verify that testing will be completed within 12 months of endorsement.

D.1 Testing: No, testing will be completed within 12 months

D.2 Have NQF-endorsed measures been reviewed to identify if there are similar or related measures? Yes

(for NQF staff use) Have all conditions for consideration been met? Met

Staff Notes to Steward (If submission returned):

Staff Notes to Reviewers (Issues or questions regarding any criteria):

Staff Reviewer Name(s):

TAP/Workgroup Reviewer Name:

Steering Committee Reviewer Name:

1. IMPORTANCE TO MEASURE AND REPORT

Extent to which the specific measure focus is important to making significant gains in health care quality (safety, timeliness, effectiveness, efficiency, equity, patient-centeredness) and improving health outcomes for a specific high impact aspect of healthcare where there is variation in or overall poor performance. Measures must be judged to be important to measure and report in order to be evaluated against the remaining criteria (evaluation criteria)

1a. High Impact

(for NQF staff use) Specific NPP goal:

1a.1 Demonstrated High Impact Aspect of Healthcare: Affects large numbers, Leading cause of morbidity/mortality, Severity of illness, Patient/societal consequences of poor quality

1a.2

1a.3 Summary of Evidence of High Impact: The Clinical Technical Expert Panel (C-TEP) felt it was important to have a measure that would be less subject to interpretation and based upon a specific, definitive, and standard measure of infection diagnosis. Thus, this measure was proposed to base some of the calculations in the hemodialysis vascular access-related measure group only upon those cases in which the blood culture is positive for an infection. This more specific measure of bacteremia will provide meaningful comparisons over time within and between dialysis units. Furthermore, infections resulting in bacteremia often represent more severe infections with greater potential for major adverse outcomes than seen in non-bacteremic infections and therefore are another important reason for specific monitoring of this important subset of infections.

In addition, the Clinical Technical Expert Panel (C-TEP) felt it was very important to determine rates of infection associated with different types of vascular access used for HD particularly since prior studies have shown much higher rates of access-related infection for central venous catheters versus native arteriovenous fistulae or prosthetic grafts. Dialysis access-related infection, particularly for catheters, has been shown to be associated with high mortality and morbidity rates, and high costs to the health care system. Reducing dialysis access-related infection rates are expected to have a high impact on reducing

Comment [KP1]: 1a. The measure focus addresses:

• a specific national health goal/priority identified by NQF’s National Priorities Partners; OR

• a demonstrated high impact aspect of healthcare (e.g., affects large numbers, leading cause of morbidity/mortality, high resource use (current and/or future), severity of illness, and patient/societal consequences of poor quality).
health care costs, and moreover, improve patient survival and patient quality of life by decreasing the occurrence of life-threatening sepsis events which are one of the possible consequences of a dialysis access-related infection. Use of various insertion/exit site disinfection procedures and various anti-microbial lock solutions in the care of catheters along with other vascular access-related infection control practices have led to substantially reduced rates of access-related infection in numerous studies [1-45]. Routinely monitoring access-related infection rates by vascular access type will provide important feedback to dialysis facilities, health policy makers, and infection-control experts regarding the effectiveness of ongoing infection control practices and impact of future changes in practice upon these types of infection rates.

The overall proposed scheme for monitoring dialysis access related infection in hemodialysis patients is described as follows:

Serious infections lead to higher hospitalization rates and poorer survival which both lead to high healthcare costs. There are three surrogate measures of serious infection: 1a) IV Antibiotic Therapy which is a surrogate for “suspected” serious infection, 1b) positive blood cultures or bacteremia, and 1c) clinical confirmation of infection. Methods of monitoring the rate of serious infection due to HD access practice include measuring the rate of 2a) clinically confirmed serious infections and 2b) serious infections with bacteremia by access type: AV fistulae, AV grafts and catheters.

1a.4 Citations for Evidence of High Impact:
<table>
<thead>
<tr>
<th></th>
<th>Authors</th>
<th>Reference</th>
</tr>
</thead>
</table>
Citations for data on disparities by population group:

N/A

1. Opportunity for Improvement

1b. Benefits (improvements in quality) envisioned by use of this measure:

Infection is known to be the second leading cause of mortality among dialysis patients, and is associated with high costs and high morbidity. However, monitoring infection rates across dialysis facilities has been lacking. By measuring catheter-related bacteremia, dialysis facilities and quality improvement organizations will be able to more accurately characterize the nature and severity of infections on a national level and implement quality improvement programs for reducing infection rates which are expected to result in improved survival, quality of life, and reduced morbidity and health care costs for dialysis patients.

1b. Summary of data demonstrating performance gap (variation or overall poor performance) across providers:

Preliminary analyses of Centers for Medicare & Medicaid Services (CMS) claims data show large variation in access-related infection across United States (US) dialysis facilities.

1b.3 Citations for data on performance gap:


1b.4 Summary of Data on disparities by population group:

N/A

1b.5 Citations for data on disparities:

N/A

1c. Outcome or Evidence to Support Measure Focus

1c.1 Relationship to Outcomes (For non-outcome measures, briefly describe the relationship to desired outcome. For outcomes, describe why it is relevant to the target population):

Measuring new IV antibiotic therapy is a surrogate for suspected serious infection, such that measurement of the frequency of new IV antibiotic therapy will be used to help facilities monitor this indicator of serious infection and target ways to prevent and reduce infection which is the desired outcome. The C-TEP felt it was important to have a measure that would be less subject to interpretation and based upon a specific, definitive, and standard measure of infection diagnosis. This, measure was proposed to base some of the calculations in the hemodialysis vascular access-related measure group upon those cases in which the blood culture is positive for an infection. This more specific measure of bacteremia will provide meaningful comparisons over time within and between dialysis units. Furthermore, infections resulting in bacteremia often represent more severe infections with greater potential for major adverse outcomes than seen in non-bacteremic infections and therefore are another important reason for specific monitoring of this important subset of infections. In addition, routinely monitoring infection rates will provide important feedback to dialysis facilities, health policy makers, and infection-control experts regarding the effectiveness of ongoing infection control practices and impact of future changes in practice upon these types of infection rates.

1c.2 Evidence that an association exists between the measure of resource use and level of performance with respect to one or more of the following:

- Intermediate outcome
- Process
- Patient experience
- System efficiency
- Balanced scorecard
- Health/avoidance of harm or cost/benefit.

Comment [k3]: All citations and references presented in this measure are maintained internally and presented to the C-TEP for review and approval. Data sources are cited in the text where appropriate and a complete list of references is provided in the reference section.

Comment [k5]: 4 Clinical care processes typically include multiple steps: assess → identify problem/potential problem → choose/plan intervention (with patient input) → provide intervention → evaluate impact on health status. If the measure focus is one step in such a multi-step process, the step with the greatest effect on the desired outcome should be selected as the focus of measurement. For example, although assessment of immunization status and recommending immunization are necessary steps, they are not sufficient to achieve the desired impact on health status - patients must be vaccinated to achieve immunity. This does not preclude consideration of measures of preventive screening interventions where there is a strong link with desired outcomes (e.g., NQF #1457: screen for hearing loss).
Furthermore, routinely monitoring access-related infection rates by vascular access type will provide important feedback to dialysis facilities, health policy makers, and infection-control experts regarding the effectiveness of ongoing infection control practices and impact of future changes in practice upon these types of infection rates.

The proposed scheme described above provides an overview of the overall proposed scheme for monitoring dialysis access-related infection in HD patients, with this particular measure contributing to elements 1b, 2a and 2b in this overall schema.

1c.2-3. Type of Evidence: Cohort study, Observational study, Evidence-based guideline, Randomized controlled trial, Meta-analysis

1c.4 Summary of Evidence (as described in the criteria; for outcomes, summarize any evidence that healthcare services/care processes influence the outcome):
A large body of literature exists showing strong associations between central venous catheter use in HD patients with poorer survival and greater morbidity [1-40]. Recent studies have shown a nearly 20% higher hazard of mortality for every 20% higher facility % catheter use [2]. The prevalence of numerous patient comorbidity indicators was similar in facilities with higher versus lower catheter use. Lower mortality has been observed with reduction in catheter use in facility- and patient-level access use studies [7, 10, 13, 40, 41]. Furthermore, much of the 30-40% higher case-mix adjusted mortality rate for US HD patients compared to those in several European countries appears to be explained by differences in vascular access use between these two regions [2]. Rates of access-related infection, including sepsisemia, have been shown to be substantially higher for patients dialyzing with a central venous catheter versus an arteriovenous fistula or graft [2, 5, 9, 14, 19, 28, 34, 36, 42, 43]. Access-related sepsisemia is strongly associated with poor survival, high rates of hospitalization, and high treatment costs (> $25,000 per episode) [9, 15, 18-20, 27, 44-48]. Numerous clinical trials have demonstrated large variability in access-related infection rates among facilities treating HD patients, while demonstrating large reductions in access-related infection rates through quality improvement programs focused on using certain anti-microbial lock solutions and/or other access-related infection control regimens [38, 49-91]. These trials provide strong evidence that access-related infection rates are modifiable with the possibility to reduce high rates of access-related infection to substantially lower levels. Several HD guideline committees and health care agencies have developed recommendations for either catheter use and/or access-related infection rates [92-96].

1c.5 Rating of strength/quality of evidence (also provide narrative description of the rating and by whom):

The evidence pertinent to this area for quality measure monitoring is of high quantity, moderate quality, and of moderate to high consistency based on a review of the literature and overviews of this subject area during guideline development by National Kidney Foundation (NKF) Kidney Disease Outcome Quality Initiative (KDOQI) and Centers for Disease Control (CDC) guideline committees. The magnitude and certainty of net benefit are expected to be moderate to high with low to no risks to patients in facilities reporting these data for purposes of quality measurement/monitoring.

1c.6 Method for rating evidence: United States Preventive Services Task Force (USPSTF) and Grading of Recommendations, Assessment, Development and Evaluation (GRADE)

1c.7 Summary of Controversy/Contradictory Evidence: Some suspected serious infections may be treated only with oral antibiotics and these will not be accounted for. This limitation is perceived to be a relatively minor exclusion in view of current practice, and has been accepted to limit the data collection to intravenous antibiotic therapy which is indicated to be much more reliable, more uniform, and less burdensome than data collection that would include oral antibiotic therapy. Furthermore, clinical confirmation of whether a suspected infection was confirmed and whether the confirmed infection was vascular access-related is expected to vary across physicians with some degree of subjectivity thus resulting in some variability in findings due to differences in interpretation of patient symptoms and laboratory findings. In addition, when some, but not all, blood cultures are indicated to be positive for an infection, variation in concluding whether blood cultures were positive for an infection is recognized as well.

26) Moist, L.M., et al., Increased hemodialysis catheter use in Canada and associated mortality risk:
54) Allon, M., Prophylaxis against dialysis catheter-related bacteremia with a novel antimicrobial lock


1c.9 Quote the Specific guideline recommendation (including guideline number and/or page number):

1. Kidney Disease Outcomes Quality Initiative (KDOQI) Clinical Practice Guidelines for Vascular Access (2006) 8.3.3.1 Catheter complications/performance should be as follows: Tunneled catheter-related infection less than 10% at 3 months and less than 50% at 1 year. (B)


1. Surveillance
   A. Conduct surveillance in ICUs and other patient populations to determine CRBSI rates, monitor trends in those rates, and assist in identifying lapses in infection control practices (3,12,16,247-250). Category IA
   B. Express ICU data as the number of catheter-associated BSIs per 1,000 catheter-days for both adults and children and stratify by birth weight categories for neonatal ICUs to facilitate comparisons with national data in comparable patient populations and healthcare settings (3,12,16,247-250). Category IB
   C. Investigate events leading to unexpected life-threatening or fatal outcomes. This includes any process variation for which a recurrence would likely present an adverse outcome (13).

   Guideline 7.16- HD: Vascular access- All HD units should collect and audit data on the form of vascular access in use in incident and prevalent haemodialysis patients and the rates of bacteraemia per 1000 patient days using central venous catheters, arterio-venous grafts and arterio-venous fistulae.

1c.10 Clinical Practice Guideline Citation: 1) Kidney Disease: Improving Global Outcomes (KDIGO) Clinical Practice Guidelines for Vascular Access (2006)
1c.11 National Guideline Clearinghouse or other URL: http://www.qualitymeasures.ahrq.gov

1c.12 Rating of strength of recommendation (also provide narrative description of the rating and by whom):
The certainty of net benefit is moderate to high, and the magnitude of the net benefit is expected to be moderate to substantial yielding a USPSTF Grade B level of recommendation. This is consistent with strength of recommendations from the following: (1) National Kidney Foundation KDOQI guideline (2006) 8.3.3.1 (shown above): Rates the strength of this guideline recommendation as Grade B: It is recommended that clinicians routinely follow the guideline for eligible patients. There is moderately strong evidence that the practice improves health outcomes. (2) CDC for Surveillance (guidelines shown above): Rates the evidence for this guideline as Category IA. Strongly recommended for implementation and strongly supported by well-designed experimental, clinical, or epidemiologic studies.

1c.13 Method for rating strength of recommendation (If different from USPSTF system, also describe rating and how it relates to USPSTF):
USPSTF

1c.14 Rationale for using this guideline over others:
N/A

TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for Importance to Measure and Report?

Steering Committee: Was the threshold criterion, Importance to Measure and Report, met?
Rationale: 1

2. SCIENTIFIC ACCEPTABILITY OF MEASURE PROPERTIES

Extent to which the measure, as specified, produces consistent (reliable) and credible (valid) results about the quality of care when implemented. (evaluation criteria)

2a. MEASURE SPECIFICATIONS

S.1 Do you have a web page where current detailed measure specifications can be obtained?
S.2 If yes, provide web page URL:

2a. Precisely Specified

2a.1 Numerator Statement (Brief, text description of the numerator - what is being measured about the target population, e.g. target condition, event, or outcome):
Overall access-related bacteremia: Number of months that hemodialysis (HD) patients initiated a new IV antibiotic therapy for a newly suspected infection during the six-month period ending with the current reporting month, and for which the infection was related to the HD access, and blood cultures were consistent with bacteremia.

Specific access types: Number of months that HD patients initiated a new IV antibiotic therapy for a newly suspected infection during the six-month period ending with the current reporting month, and for which the infection was related to the fistula/graft/catheter used as HD access, and blood cultures were consistent with bacteremia.

2a.2 Numerator Time Window (The time period in which cases are eligible for inclusion in the numerator): Six months ending with the current reporting month. (for all access types)

2a.3 Numerator Details (All information required to collect/calculate the numerator, including all codes,
logic, and definitions):  
Vascular access-related bacteremia:  
A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the vascular catheter used as HD access (RQMT_1315), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

Specific access types:  
A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the arteriovenous fistula used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the arteriovenous graft used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the catheter used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

2a.4 Denominator Statement (Brief, text description of the denominator - target population being measured):  
Overall access-related bacteremia: All adult (18+) chronic maintenance HD patient days during the six-month period ending with the current reporting month.

Specific access types: All adult (18+) chronic maintenance HD fistula/graft/catheter days during the six-month period ending with the current reporting month.

2a.5 Target population gender: Female, Male

2a.6 Target population age range: Adults 18 years or older.

2a.7 Denominator Time Window (The time period in which cases are eligible for inclusion in the denominator):  
Six months ending with the current reporting month. (for all access types)

2a.8 Denominator Details (All information required to collect/calculate the denominator - the target population being measured - including all codes, logic, and definitions):  
Vascular access-related bacteremia:  
HD days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period and on chronic HD at the facility. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not
The number of HD days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria.

Specific access types:

Fistula days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a arteriovenous fistula at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is an arteriovenous fistula and the date access type changed to an arteriovenous fistula is on or before the last day of the reporting period.

The number of fistula days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.

Graft days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a arteriovenous graft at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is an arteriovenous graft and the date access type changed to an arteriovenous graft is on or before the last day of the reporting period.

The number of graft days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.

Catheter days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a catheter at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is a catheter and the date access type changed to a catheter is on or before the last day of the reporting period.

The number of catheter days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.

**2a.9 Denominator Exclusions (Brief text description of exclusions from the target population):** HD patients < 18 yrs old.

**Comment [k9]:** 11 Risk factors that influence outcomes should not be specified as exclusions.
12 Patient preference is not a clinical exception to eligibility and can be influenced by provider interventions.
2a.10 Denominator Exclusion Details (All information required to collect exclusions to the denominator, including all codes, logic, and definitions):
None

2a.11 Stratification Details/Variables (All information required to stratify the measure including the stratification variables, all codes, logic, and definitions):
This measure can be stratified by vascular access type as noted in the numerator and denominator statements.

2a.12-13 Risk Adjustment Type: No risk adjustment necessary

2a.14 Risk Adjustment Methodology/Variables (List risk adjustment variables and describe conceptual models, statistical models, or other aspects of model or method):
N/A

2a.15-17 Detailed risk model available Web page URL or attachment:

2a.18-19 Type of Score: Rate/proportion
2a.20 Interpretation of Score: Better quality = Lower score

2a.21 Calculation Algorithm (Describe the calculation of the measure as a flowchart or series of steps):
Vascular access-related bacteremia:
HD days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period and on chronic HD at the facility. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred.

The number of HD days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria.

A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the vascular access (RQMT_1315), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

Specific access types:
Fistula days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a arteriovenous fistula at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is an arteriovenous fistula and the date access type changed to an arteriovenous fistula is on or before the last day of the reporting period.

The number of fistula days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.
A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the arteriovenous fistula used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

Graft days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a arteriovenous graft at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is an arteriovenous graft and the date access type changed to an arteriovenous graft is on or before the last day of the reporting period.

The number of graft days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.

A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the arteriovenous graft used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

Catheter days are included in the denominator from a patient who is >= 18 years old at the start of the six-month reporting period, on chronic HD at the facility, and access type for dialysis is a catheter at any time during the six-month reporting period. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting period. The patient will be considered on chronic dialysis if the date of initiating regular chronic dialysis is prior to or equal to the last day of the six-month reporting period. The patient will be considered to be on HD if HD treatment start date is on or before the last day of the six-month reporting period and the patient was receiving HD during the six-month reporting period. A patient is considered to be treated in a facility if the admit date is on or before the last day of the reporting period and the discharge date is on or after the first day of the period or discharge has not occurred. A patient will be considered to be using a catheter if access type for HD any time during the six-month reporting period is a catheter and the date access type changed to a catheter is on or before the last day of the reporting period.

The number of catheter days in the denominator is calculated by summing the number of days during the reporting period that a patient meets the above inclusion criteria and was using a catheter during the six-month reporting period.

A month is included in the numerator if a patient in the denominator had been prescribed an IV antibiotic (RQMT_1319 and RQMT_1323) during that month for a newly suspected infection which was clinically confirmed (RQMT_1312) and related to the catheter used as HD access (RQMT_1315 and RQMT_1314), the date that the patient was prescribed IV antibiotic therapy (RQMT_1534) falls within the parameters of the reporting period, this date occurred when the patient was considered to be a chronic HD patient, and blood cultures for this confirmed infection were consistent with bacteremia (RQMT_1462 and RQMT_1317).

2a.22 Describe the method for discriminating performance (e.g., significance testing):
The performance of the facility will be compared to State, Network and National performance. Calculation
of the facility-level measure will be performed by: (a) summing the numerator values for each reporting period-eligible facility patient to obtain a facility-level numerator sum, (b) summing the denominator values for each reporting period-eligible facility patient to obtain a facility-level denominator sum, and (c) dividing the facility-level numerator sum by the facility-level denominator and multiply the result by 1000 to obtain the number of infections per 1000 HD/fistula/graft/catheter days.

2a.23 Sampling (Survey) Methodology If measure is based on a sample (or survey), provide instructions for obtaining the sample, conducting the survey and guidance on minimum sample size (response rate): N/A

2a.24 Data Source (Check the source(s) for which the measure is specified and tested) 
Electronic clinical data

2a.25 Data source/data collection instrument (identify the specific data source/data collection instrument, e.g. name of database, clinical registry, collection instrument, etc.): CROWNWeb


2a.32-35 Level of Measurement/Analysis (Check the level(s) for which the measure is specified and tested) 
Facility/Agency

2a.36-37 Care Settings (Check the setting(s) for which the measure is specified and tested) 
Dialysis Facility

2a.38-41 Clinical Services (Healthcare services being measured, check all that apply) 
Dialysis

<table>
<thead>
<tr>
<th>TESTING/ANALYSIS</th>
</tr>
</thead>
<tbody>
<tr>
<td>2b. Reliability testing</td>
</tr>
<tr>
<td>2b.1 Data/sample (description of data/sample and size): N/A</td>
</tr>
<tr>
<td>2b.2 Analytic Method (type of reliability &amp; rationale, method for testing): N/A</td>
</tr>
<tr>
<td>2b.3 Testing Results (reliability statistics, assessment of adequacy in the context of norms for the test conducted): N/A</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>2c. Validity testing</th>
</tr>
</thead>
<tbody>
<tr>
<td>2c.1 Data/sample (description of data/sample and size): N/A</td>
</tr>
<tr>
<td>2c.2 Analytic Method (type of validity &amp; rationale, method for testing): Face validity is the only validity assessed, therefore testing is not applicable.</td>
</tr>
<tr>
<td>2c.3 Testing Results (statistical results, assessment of adequacy in the context of norms for the test conducted): N/A</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>2d. Exclusions Justified</th>
</tr>
</thead>
<tbody>
<tr>
<td>2d.1 Summary of Evidence supporting exclusion(s): Measures are currently limited to HD patients since a separate expert panel will be convened in the future</td>
</tr>
</tbody>
</table>

2d.2 Exclusions Justified

| Comment [KP10]: 2b. Reliability testing demonstrates the measure results are repeatable, producing the same results a high proportion of the time when assessed in the same population in the same time period. |
| Comment [KP11]: 2d. Exclusions Justified the specific topic; ability of measure scores to predict scores on some other related valid measure; content validity for multi-item scales/test-retest for survey items. Reliability testing may address the data items or final measure score. |
| Comment [KP12]: 2c. Validity testing demonstrates that the measure reflects the quality of care provided, adequately distinguishing good and poor quality. If face validity is the only validity addressed, it is systematically assessed. |
| Comment [KP13]: 2b. Reliability testing include, but are not limited to: inter-rater/abstractor or intra-rater/abstractor studies; internal consistency for multi-item scales; test-retest for survey items. Reliability testing may address the data items or final measure score. |
| Comment [KP14]: 2d. Clinically necessary measure exclusions are identified and must be: supported by evidence of sufficient frequency of occurrence so that results are distorted without the exclusion; AND clinically appropriate exception (e.g., contraindication) to eligibility for the measure focus; AND precisely defined and specified: –If there is substantial variability in exclusions across providers, the measure is specified so that exclusions are computable and the effect on the measure is transparent (i.e., impact clearly delineated, such as number of cases excluded, exclusion rates by type of exclusion); if patient preference (e.g., informed decision-making) is a basis for exclusion, there must be evidence that it strongly impacts performance on the measure and the measure must be specified so that the information about patient preference and the effect on the measure is transparent (e.g., numerator category) ... [2] |
| Comment [KP15]: 2d. Exclusions Justified: Measures are currently limited to HD patients since a separate expert panel will be convened in the future |

Rating: C=Completely; P=Partially; M=Minimally; N=Not at all; NA=Not applicable
### 2e. Risk Adjustment for Outcomes/Resource Use Measures

<table>
<thead>
<tr>
<th>NQF #1457</th>
<th>2e</th>
<th>C</th>
<th>P</th>
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<th>N</th>
<th>NA</th>
</tr>
</thead>
</table>

#### 2e.1 Data/sample (description of data/sample and size): N/A

**2e.2 Analytic Method (type of risk adjustment, analysis, & rationale):**

* N/A

**2e.3 Testing Results (risk model performance metrics):**

* N/A

**2e.4 If outcome or resource use measure is not risk adjusted, provide rationale:** There are no compelling reasons to risk adjust measure.

### 2f. Identification of Meaningful Differences in Performance

<table>
<thead>
<tr>
<th>NQF #1457</th>
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<th>NA</th>
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</thead>
</table>

#### 2f.1 Data/sample from Testing or Current Use (description of data/sample and size): N/A

#### 2f.2 Methods to identify statistically significant and practically/meaningfully differences in performance (type of analysis & rationale):

* N/A

#### 2f.3 Provide Measure Scores from Testing or Current Use (description of scores, e.g., distribution by quartile, mean, median, SD, etc.; identification of statistically significant and meaningfully differences in performance):

* N/A

### 2g. Comparability of Multiple Data Sources/Methods

<table>
<thead>
<tr>
<th>NQF #1457</th>
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<th>P</th>
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<th>N</th>
<th>NA</th>
</tr>
</thead>
</table>

#### 2g.1 Data/sample (description of data/sample and size): N/A

#### 2g.2 Analytic Method (type of analysis & rationale):

* N/A

#### 2g.3 Testing Results (e.g., correlation statistics, comparison of rankings):

* N/A

### 2h. Disparities in Care

<table>
<thead>
<tr>
<th>NQF #1457</th>
<th>2h</th>
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</table>

#### 2h.1 If measure is stratified, provide stratified results (scores by stratified categories/cohorts): N/A

#### 2h.2 If disparities have been reported/identified, but measure is not specified to detect disparities, provide follow-up plans:

* N/A

**TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for Scientific Acceptability of Measure Properties?**

* 2
### 3a. Meaningful, Understandable, and Useful Information

**3a.1 Current Use:** Testing not yet completed


**3a.3 If used in other programs/initiatives** (if used in quality improvement or other programs/initiatives, name of initiative(s), locations, Web page URL(s). If not used for QI, state the plans to achieve use for QI within 3 years):


**3a.4 Data/sample (description of data/sample and size):** 32 dialysis facilities provided HD reported adverse events related to infection to the CDC in 2006. These facilities submitted data on 28,047 patient-months.

**3a.5 Methods (e.g., focus group, survey, QI project):** Staff from the participating dialysis facilities monitored and reported vascular access type, new IV antimicrobial starts and positive blood cultures for patients and entered data monthly into NHSN’s reporting tool. The data were accumulated from all centers and analyzed at CDC. The definition of an access-associated bloodstream infection was a microorganism identified in a blood culture where the infection source was the vascular access site. A bloodstream infection was defined as a positive blood culture report, regardless of the infection source, and included access-associated bloodstream infections. The definition of vascular access infection was either a local access infection or an access-associated bloodstream infection.

**3a.6 Results (qualitative and/or quantitative results and conclusions):** The pooled mean rates of IV antibiotic starts among patients with arteriovenous fistulas, grafts, permanent and temporary central venous catheters were 1.8, 2.4, 6.4, and 25.4 per 100 patient-months, respectively. For bloodstream infection, the pooled mean rates were 0.5, 0.9, 4.2, and 27.1 per 100 patient-months and for access-related bloodstream infection, the pooled means were 0.2, 0.4, 3.1, and 17.8 in these groups. For vascular access infection, the pooled rates were 0.4, 0.9, 4.8, and 22.9 per 100 patient-months respectively.

### 3b. Harmonization

**3b.1 NQF # and Title of similar or related measures:**

(for NQF staff use) Notes on similar/related endorsed or submitted measures:

**3b.2 Are the measure specifications harmonized? If not, why?**
### 3c. Distinctive or Additive Value

3c.1 Describe the distinctive, improved, or additive value this measure provides to existing NQF-endorsed measures:

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</thead>
<tbody>
<tr>
<td>3c</td>
<td>C</td>
<td>P</td>
<td>M</td>
<td>N</td>
<td>NA</td>
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</table>

5.1 If this measure is similar to measure(s) already endorsed by NQF (i.e., on the same topic and the same target population), Describe why it is a more valid or efficient way to measure quality: N/A

### TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for Usability?

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### 4. FEASIBILITY

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

#### 4a. Data Generated as a Byproduct of Care Processes

4a.1-2 How are the data elements that are needed to compute measure scores generated?

Data generated as byproduct of care processes during care delivery (Data are generated and used by healthcare personnel during the provision of care, e.g., blood pressure, lab value, medical condition)

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<tr>
<td>4a</td>
<td>C</td>
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</table>

#### 4b. Electronic Sources

4b.1 Are all the data elements available electronically? (Elements that are needed to compute measure scores are in defined, computer-readable fields, e.g., electronic health record, electronic claims)

- Yes

4b.2 If not, specify the near-term path to achieve electronic capture by most providers.

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#### 4c. Exclusions

4c.1 Do the specified exclusions require additional data sources beyond what is required for the numerator and denominator specifications?

- No

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</table>

4c.2 If yes, provide justification.

#### 4d. Susceptibility to Inaccuracies, Errors, or Unintended Consequences

4d.1 Identify susceptibility to inaccuracies, errors, or unintended consequences of the measure and describe how these potential problems could be audited. If audited, provide results.

Facilities may not be aware of IV antibiotics prescribed if patients are hospitalized. Claims data may help with auditing of this. This measure requires physician input of whether infection was access-related which will have a degree of subjectivity.

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#### 4e. Data Collection Strategy/Implementation

4e.1 Describe what you have learned/modified as a result of testing and/or operational use of the measure regarding data collection, availability of data/missing data, timing/frequency of data collection, patient confidentiality, time/cost of data collection, other feasibility/implementation issues:

Data elements were reviewed and input was received by a data technical expert panel which includes representatives from many types of US dialysis facilities. The proposed measures are based on feedback.

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<tr>
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from this group regarding feasibility of data collection.

4e.2 Costs to implement the measure (costs of data collection, fees associated with proprietary measures): N/A

4e.3 Evidence for costs: N/A

4e.4 Business case documentation: Infection, particularly for those resulting in bacteremia in association with catheter use in dialysis patients, has been shown to be associated with high costs to the health care system. Reducing infection rates are expected to have a high impact on reducing health care costs.

TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for Feasibility?

<table>
<thead>
<tr>
<th>Rationale</th>
<th>4</th>
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</table>

Steering Committee: Overall, to what extent was the criterion, Feasibility, met?

Rationale:

<table>
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RECOMMENDATION

(for NQF staff use) Check if measure is untested and only eligible for time-limited endorsement.

Steering Committee: Do you recommend for endorsement?

Comments:

<table>
<thead>
<tr>
<th>Y</th>
<th>N</th>
<th>A</th>
</tr>
</thead>
</table>

CONTACT INFORMATION

Co.1 Measure Steward (Intellectual Property Owner)

Co.1 Organization
Centers for Medicare and Medicaid Services, 7500 Security Boulevard, Baltimore, Maryland, 21244

Co.2 Point of Contact
Thomas, Dudley, Thomas.Dudley@cms.hhs.gov, 410-786-1442-

Measure Developer if different from Measure Steward

Co.3 Organization
Arbor Research/UM-KECC, 315 W. Huron, Suite 360, Ann Arbor, Michigan, 48103

Co.4 Point of Contact
Adrienne, Janney, adrienne.janney@arborresearch.org, 734-665-4108-

Co.5 Submitter if different from Measure Steward POC
Thomas, Dudley, Thomas.Dudley@cms.hhs.gov, 410-786-1442-, Centers for Medicare and Medicaid Services

Co.6 Additional organizations that sponsored/participated in measure development

ADDITIONAL INFORMATION

Workgroup/Expert Panel involved in measure development

Ad.1 Provide a list of sponsoring organizations and workgroup/panel members’ names and organizations.

Dr. Michael Allon, expert panel chair (University of Alabama at Birmingham)
Ms. Lesley Dinwiddie (Nephrology Nurse Consulting, Nurse Consultant)
Dr. Eduardo Lacson (Fresenius Medical Care)
Dr. Derrick Latos (Nephrology Associates, Inc., Forum of ESRD Networks)
Dr. Charmaine Lok (Toronto General Research Institute, Toronto General Hospital)
Dr. Ted Steinman (Beth Israel Hospital, Harvard Medical School)  
Dr. Daniel Weiner (Tufts Medical Center)  
Ms. Raynel Wilson (ESRD Network 9 & ESRD Network 10, The Renal Network, Inc.)  
Dr. Ronald Pisoni, moderator for contractor (Arbor Research Collaborative for Health)  
Ms. Natalie Lueth, analyst for contractor (University of Michigan KECC)  

| Ad.2 | If adapted, provide name of original measure: |
| Ad.3-5 | If adapted, provide original specifications URL or attachment |

**Measure Developer/Steward Updates and Ongoing Maintenance**  
Ad.6 Year the measure was first released:  
Ad.7 Month and Year of most recent revision:  
Ad.8 What is your frequency for review/update of this measure? Three years  
Ad.9 When is the next scheduled review/update for this measure? 2013  

| Ad.10 | Copyright statement/disclaimers: |
| Ad.11 -13 | Additional Information web page URL or attachment: |

**Date of Submission (MM/DD/YY):** 12/21/2010
4. Clinical care processes typically include multiple steps: assess → identify problem/potential problem → choose/plan intervention (with patient input) → provide intervention → evaluate impact on health status. If the measure focus is one step in such a multi-step process, the step with the greatest effect on the desired outcome should be selected as the focus of measurement. For example, although assessment of immunization status and recommending immunization are necessary steps, they are not sufficient to achieve the desired impact on health status - patients must be vaccinated to achieve immunity. This does not preclude consideration of measures of preventive screening interventions where there is a strong link with desired outcomes (e.g., mammography) or measures for multiple care processes that affect a single outcome.

2d. Clinically necessary measure exclusions are identified and must be:
• supported by evidence of sufficient frequency of occurrence so that results are distorted without the exclusion; AND
• a clinically appropriate exception (e.g., contraindication) to eligibility for the measure focus; AND
• precisely defined and specified:
  – if there is substantial variability in exclusions across providers, the measure is specified so that exclusions are computable and the effect on the measure is transparent (i.e., impact clearly delineated, such as number of cases excluded, exclusion rates by type of exclusion);
if patient preference (e.g., informed decision-making) is a basis for exclusion, there must be evidence that it strongly impacts performance on the measure and the measure must be specified so that the information about patient preference and the effect on the measure is transparent (e.g., numerator category computed separately, denominator exclusion category computed separately).