NQF #0249 Hemodialysis Adequacy Clinical Performance Measure III: Hemodialysis Adequacy--HD Adequacy--
Minimum Delivered Hemodialysis Dose

NATIONAL QUALITY FORUM

Measure Submission and Evaluation Worksheet 5.0

This form contains the information submitted by measure developers/stewards, organized according to NQF’s measure evaluation criteria and process. The evaluation criteria, evaluation guidance documents, and a blank online submission form are available on the submitting standards web page.

<table>
<thead>
<tr>
<th>NQF #: 0249</th>
<th>NQF Project: Renal Endorsement Maintenance 2011</th>
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<tbody>
<tr>
<td>(for Endorsement Maintenance Review)</td>
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<tr>
<td>Original Endorsement Date: Nov 15, 2007</td>
<td>Most Recent Endorsement Date: Nov 15, 2007</td>
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**BRIEF MEASURE INFORMATION**

De.1 Measure Title: Hemodialysis Adequacy Clinical Performance Measure III: Hemodialysis Adequacy--HD Adequacy-- Minimum Delivered Hemodialysis Dose

Co.1.1 Measure Steward: Centers for Medicare & Medicaid Services

De.2 Brief Description of Measure: Percentage of all adult (>=18 years old) patients in the sample for analysis who have been on hemodialysis for 90 days or more and dialyzing thrice weekly whose average delivered dose of hemodialysis (calculated from the last measurements of the month using the UKM or Daugirdas II formula) was a spKt/V >= 1.2 during the study period.

2a1.1 Numerator Statement: Number of patients in denominator whose delivered dose of hemodialysis (calculated from the last measurements of the month using the UKM or Daugirdas II formula) was a spKt/V >= 1.2.

2a1.4 Denominator Statement: All adult (>= 18 years old) patients in the sample for analysis who have been on hemodialysis for 90 days or more and dialyzing thrice weekly.

2a1.8 Denominator Exclusions: Patients on HD less than 90 days; HD patients dialyzing <3 times per week or >3 times per week.

1.1 Measure Type: Outcome
2a1. 25-26 Data Source: Electronic Clinical Data
2a1.33 Level of Analysis: Facility

1.2-1.4 Is this measure paired with another measure? No

De.3 If included in a composite, please identify the composite measure (title and NQF number if endorsed):

**STAFF NOTES** (issues or questions regarding any criteria)

Comments on Conditions for Consideration:

Is the measure untested? Yes☐ No☐ If untested, explain how it meets criteria for consideration for time-limited endorsement:

1a. Specific national health goal/priority identified by DHHS or NPP addressed by the measure (check De.5): 5. Similar/related endorsed or submitted measures (check 5.1):

Other Criteria:

Staff Reviewer Name(s):

1. IMPACT, OPPORTUNITY, EVIDENCE - IMPORTANCE TO MEASURE AND REPORT

Importance to Measure and Report is a threshold criterion that must be met in order to recommend a measure for endorsement. All
three subcriteria must be met to pass this criterion. See [guidance on evidence](#).

**Measures must be judged to be important to measure and report in order to be evaluated against the remaining criteria.**

(evaluation criteria)

<table>
<thead>
<tr>
<th>1a. High Impact:</th>
<th>H</th>
<th>M</th>
<th>L</th>
<th>I</th>
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<tbody>
<tr>
<td><em>(The measure directly addresses a specific national health goal/priority identified by DHHS or NPP, or some other high impact aspect of healthcare.)</em></td>
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De.4 Subject/Topic Areas *(Check all the areas that apply):* Renal, Renal : End Stage Renal Disease (ESRD)

De.5 Cross Cutting Areas *(Check all the areas that apply):*

1a.1 Demonstrated High Impact Aspect of Healthcare: Affects large numbers, High resource use

1a.2 If “Other,” please describe:

1a.3 Summary of Evidence of High Impact *(Provide epidemiologic or resource use data):*

The dose of dialysis is used to estimate the ability of hemodialysis to clear the blood of accumulated toxins. In the adult population, outcome studies have shown an association between dose of hemodialysis in terms of small solute removal and clinical outcomes[1,2]. In addition, at least one prior study demonstrates that a change in dialysis dose is associated with a change in patient outcome [3]. Furthermore, studies demonstrate an association between dialysis adequacy as measured by Kt/V and outcomes [4,5,6]. Also, although higher dialysis dose is associated with improvement in clinical outcomes, analysis of CROWNWeb data from January 2010 indicate that only 66% of facilities had 70% or more of their patients receiving a dialysis dose of spKt/V of 1.2.

For this measure maintenance cycle, we propose that this measure remains in its current format. Since endorsement of this measure, published literature suggests there is insufficient evidence that compares methods of dialysis adequacy measurement, particularly measures that demonstrate superiority of alternative measures over spKt/V. It should also be noted that there have been no changes in the KDOQI Clinical Practice Guideline for Methods for Measuring and Expressing Hemodialysis Dose (CPG 2). Indeed, as stated in the KDOQI 2006 update, ‘The delivered Kt/V determined by single-pool urea kinetic modeling continues to be preferred as the most precise and accurate measure of dialysis.’ (p.12, KDOQI 2006 Update).

Currently, frequent hemodialysis (more than thrice weekly) is still rare, with approximately 1% of dialysis patients receiving this modality. As this population grows and the evidence base for alternative adequacy measurement methods grows, the use of stdKt/V, in particular should be evaluated by a Clinical Technical Expert Panel (CTEP), including a target measure because of the potential for a growing percentage of patients being dialyzed more than thrice weekly and where spKt/V is not comparable across treatment schedules.

Additional considerations for future expert review of the use of spKt/V measure relates to women and smaller patients. Recent studies that examine dialysis dosing in women and smaller patients should be considered [7,8]. In addition, because prior studies that evaluate the impact of hemodialysis dose on mortality have used spKt/V as the measure of hemodialysis adequacy, alternative methods of adequacy measurement should also be considered. Finally, recent clinical studies suggest the benefit of using online measurement methods for assessing ionic clearance, and these tools should be considered in the future [9].

Another point to be considered with this measure is the requirement for a 6-month duration of HD before inclusion into the measure. This is in contrast to a related measure which has received a time-limited endorsement, which requires only a 3 month duration of HD, but documentation that residual renal function (RRF) is less than 2ml/min/1.73m2 prior to inclusion. These may have RRF, and second, that measurement of this RRF is not necessarily standard practice. Because none of these contributing factors has changed, we propose to maintain the requirement for a 6-month duration of HD prior to inclusion into this measure.

1a.4 Citations for Evidence of High Impact cited in 1a.3:  

1b. Opportunity for Improvement:  [ ] H [ ] M [ ] L [ ] I [ ]
(There is a demonstrated performance gap - variability or overall less than optimal performance)

1b.1 Briefly explain the benefits (improvements in quality) envisioned by use of this measure:
Published studies indicate there is an association between low spKt/V and increased mortality. Furthermore, the 2006 KDOQI Hemodialysis Adequacy Guidelines indicate ‘minimally adequate dose of HD given 3 times per week to patients with Kr less than 2 mL/min/1.73 m2 should be an spKt/V of 1.2 per dialysis.’

1b.2 Summary of Data Demonstrating Performance Gap (Variation or overall less than optimal performance across providers):
[For Maintenance – Descriptive statistics for performance results for this measure - distribution of scores for measured entities by quartile/decile, mean, median, SD, min, max, etc.]
Analysis of CROWNWeb data from January 2010 indicate 66% of facilities had at least 70% of patients meeting the spKt/V >=1.2 dose requirement. Only 4% of facilities had 100% of patients meeting the requirements for this measure.

1b.3 Citations for Data on Performance Gap: [For Maintenance – Description of the data or sample for measure results reported in 1b.2 including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included] Performance gap analyses were performed using CROWNWeb data from January 2010. There were 3387 facilities reporting data for this measure, and a total of 293,694 patients in this reporting month. Mean number of patients per facility was 84 (SD=52).

1b.4 Summary of Data on Disparities by Population Group: [For Maintenance – Descriptive statistics for performance results for this measure by population group]
For each facility, the percent of patients by demographic group including sex, race, ethnicity, and age category, was calculated. Facilities were then divided into quintiles based on their percentage within each demographic category. Within each facility-level quintile, the average of each facility’s performance measure was calculated. The means were examined for trend across quintile. No disparities in performance were observed by race, sex, ethnicity, or age.
The range in percent of patients with spKt/V >= 1.2 across quintiles is presented below.
Population Group (Range):
Females (70.2%-72.5%)
Males (70.2%-72.5%)
Black (69.0%-73.8%)* highest performance in facilities with 69% or more Black patients
White (68.6%-74.1%)* highest performance in facilities with 0-27% White patients
Hispanic (67.7%-75.5%)* highest performance in facilities 19% or more Hispanic patients
Age (70.6%-73.6% by age group)

1b.5 Citations for Data on Disparities Cited in 1b.4: [For Maintenance – Description of the data or sample for measure results reported in 1b.4 including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included] CROWNWeb data from July 2009-December 2009 were analyzed. The number of facilities ranged from 3398-3453 and the total number of patients per month ranged from 263,743 - 290,713.

1c. Evidence (Measure focus is a health outcome OR meets the criteria for quantity, quality, consistency of the body of evidence.)
Is the measure focus a health outcome?  [ ] Yes  [ ] No  [ ] If not a health outcome, rate the body of evidence.
## NQF #0249 Hemodialysis Adequacy Clinical Performance Measure III: Hemodialysis Adequacy--HD Adequacy--Minimum Delivered Hemodialysis Dose

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Quality</th>
<th>Consistency</th>
<th>Does the measure pass subcriterion 1c?</th>
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<tbody>
<tr>
<td>M-H</td>
<td>M-H</td>
<td>M-H</td>
<td>Yes if additional research unlikely to change conclusion that benefits to patients outweigh harms; otherwise No</td>
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<tr>
<td>L</td>
<td>M-H</td>
<td>M</td>
<td>Yes if potential benefits to patients clearly outweigh potential harms; otherwise No</td>
</tr>
</tbody>
</table>

### Health outcome – rationale supports relationship to at least one healthcare structure, process, intervention, or service

#### 1c.1 Structure-Process-Outcome Relationship

**Briefly state the measure focus, e.g., health outcome, intermediate clinical outcome, process, structure; then identify the appropriate links, e.g., structure-process-health outcome; process-health outcome; intermediate clinical outcome-health outcome:**

*The measure focus is measurement of spKt/V >= 1.2.*

This process leads to improvement in mortality as follows:

*Measure spKt/V--> Assess value-->Impact on mortality.*

#### 1c.2-3 Type of Evidence

*Check all that apply:*

- Clinical Practice Guideline
- Selected individual studies (rather than entire body of evidence)

#### 1c.4 Directness of Evidence to the Specified Measure

**State the central topic, population, and outcomes addressed in the body of evidence and identify any differences from the measure focus and measure target population:**

*The body of evidence shows a relationship between low spKt/V and improved mortality and morbidity. This is a target measure for spKt/V below which a higher risk for adverse outcomes is observed. The evidence is directly related to this measure.*

#### 1c.5 Quantity of Studies in the Body of Evidence

**Total number of studies, not articles:** 11

#### 1c.6 Quality of Body of Evidence

**Summarize the certainty or confidence in the estimates of benefits and harms to patients across studies in the body of evidence resulting from study factors. Please address: a) study design/flaws; b) directness/indirectness of the evidence to this measure (e.g., interventions, comparisons, outcomes assessed, population included in the evidence); and c) imprecision/wide confidence intervals due to few patients or events:**

*Studies supporting the relationship between delivered dose of dialysis and clinical outcomes include a clinical trial and prospective and retrospective cohort studies. There was a well-designed randomized controlled clinical trial (the HEMO study), two were prospective studies and the remaining were retrospective cohort studies. The HEMO study with its randomized design and measurement of hard outcomes was given significant importance in defining the Hemodialysis Adequacy Guidelines for the KDOQI, thereby suggesting that the quality of at least this study was high. One of the prospective studies (ref 4) is based on the Dialysis Outcomes and Practice Patterns Study, which is an international prospective study of dialysis practices on patient outcomes. Although not a clinical trial, findings from the DOPPS have generally informed the formation of KDOQI clinical guidelines because the study population is large, nationally representative by design, with adequate longitudinal follow-up. Another study cited is based on the USRDS (ref 7) which included a national US random sample of prevalent hemodialysis patients. Finally, another study cited (ref 3) compares the association between dialysis adequacy and clinical outcomes with both the CMS and DOPPS datasets. Altogether, these suggest that the body of evidence for this measure is of generally acceptable quality.*

An overall grade was not assigned to the body of evidence. However, individual studies were graded in the KDOQI Guidelines based on applicability and methodological quality. Applicability was graded according to the population of interest. Three grades were defined including, (1) sample is representative of target population, or results are definitely applicable to the target population irrespective of study sample; (2) sample is representative of a relevant subgroup of the target population; and (3) sample is representative of a narrow subgroup of patients only, and may not be generalizable to other subgroups.

Methodological quality, or internal validity, referred to the design, conduct and reporting of the clinical study. A 3-level classification of study quality was devised: (1) least bias; results are valid; (2) susceptible to some bias, but not sufficient to invalidate the results;
and (3) significant bias that may invalidate the results. Of the studies referenced in 1c.15, six received the highest grade (1) for applicability, four received a grade of 2, and one received a 3. For methodological quality, three studies received the highest grade (1), one received a 2, and the remaining studies received a 3.

1c.7 Consistency of Results across Studies (Summarize the consistency of the magnitude and direction of the effect): Dialysis adequacy as calculated by spKt/V was utilized in several studies, and direction of effect (benefit of higher dialysis dose) tended to be consistent across studies. Comparison of magnitude of effect was not done because of differences in analytical methods. In the DOPPS study, the relative risk of mortality for spKt/V < 1.2 was 1.16 (Port FK, Pisoni RL, Bragg-Gresham JL, et al: DOPPS estimates of patient life years attributable to modifiable hemodialysis treatment practices in the United States. Blood Purif 22:175-180, 2004). Another analysis revealed that a 0.1 unit higher value of spKt/V had a 7.5% lower risk of mortality (8). In another cohort, patients who received an spKt/V of 0.7 had a 2.8 increased mortality risk RR as compared to patients who received an spKt/V of between 1.2-1.3 (9). Finally, in the HEMO study (1), patient mortality did not improve at higher target dialysis above that recommended by clinical guidelines. The HEMO study also used Urea Reduction Ratio, spKt/V and eKt/V in the design of the study.

1c.8 Net Benefit (Provide estimates of effect for benefit/outcome; identify harms addressed and estimates of effect; and net benefit - benefit over harms): Among the studies showing a significant improvement in mortality with increasing dose of spKt/V, relative risks (RR) were presented as spKt/V per increase of 1 unit and spKt/V per 0.1 unit, where spKt/V was analyzed as a continuous measure. The RR per 1 unit increase in spKt/V was 0.76 (95% CI: 0.64, 0.92; p=0.004) [3], and per 0.1 unit increase in spKt/V was 0.95; p<0.05 (no CI given) [8]. The HEMO trial found no significant difference in mortality among patients in the high dose group, with mean = 1.56 and SD=0.09, compared to the low dose group with mean=1.16 and SD=0.08 (RR=0.96; 95% CI: 0.84, 1.10) [1], thus supporting the current target spKt/V of 1.2. However, a subgroup analysis of the HEMO study [2] showed that survival rates in women randomized to the higher dose group were higher than women in the lower dose group (relative risk 0.81; p = 0.02) and this association persisted after adjusting for body size. In the remaining study, findings showed patients receiving the highest dialysis dose(spKt/V>2.4) compared to the standard dose group (spKt/V 1.2-1.3) had an increased risk of mortality (RR=2.5; p<0.05), although this may be suggestive of confounding by indication. No other significant associations between dose groups were found in this study [9]. All but one study showed a benefit for a minimum dose of dialysis when measured as spKt/V. Studies evaluating higher doses of dialysis adequacy did not demonstrate additional benefit at spKt/V doses higher than the current target of 1.2. The increase in mortality at the highest dialysis dose is thought to be due to confounding by indication and does not suggest that higher dialysis dose is associated with increased mortality.

1c.9 Grading of Strength/Quality of the Body of Evidence. Has the body of evidence been graded? No

1c.10 If body of evidence graded, identify the entity that graded the evidence including balance of representation and any disclosures regarding bias:

1c.11 System Used for Grading the Body of Evidence: Other

1c.12 If other, identify and describe the grading scale with definitions: The overall body of evidence was not graded, but individual studies were graded as indicated above.

1c.13 Grade Assigned to the Body of Evidence: An overall grade was not assigned, but individual studies were graded as above.

1c.14 Summary of Controversy/Contradictory Evidence: No controversial or contradictory evidence was found.

1c.15 Citations for Evidence other than Guidelines (Guidelines addressed below):

1c.16 **Quote verbatim, the specific guideline recommendation (Including guideline # and/or page #):**

Clinical Practice Guidelines for Hemodialysis Adequacy

**GUIDELINE 4. MINIMALLY ADEQUATE HEMODIALYSIS**

The minimally adequate dose of HD given 3 times per week to patients with Kr less than 2 mL/min/1.73 m2 should be an spKt/V (excluding RKF) of 1.2 per dialysis. For treatment times less than 5 hours, an alternative minimum dose is a URR of 65%. (A)

1c.17 **Clinical Practice Guideline Citation:** National Kidney Foundation. KDOQI Clinical Practice Guidelines and Clinical Practice Recommendations for Hemodialysis Adequacy, Update 2006.

1c.18 **National Guideline Clearinghouse or other URL:**


1c.19 **Grading of Strength of Guideline Recommendation.** Has the recommendation been graded? Yes

1c.20 **If guideline recommendation graded, identify the entity that graded the evidence including balance of representation and any disclosures regarding bias:** KDOQI members. No information on representation of disclosures regarding bias.

1c.21 **System Used for Grading the Strength of Guideline Recommendation:** Other

1c.22 **If other, identify and describe the grading scale with definitions:** The rating system defined in the KDOQI Guidelines was used to grade the strength of the Guideline recommendation. KDOQI defined grades as follows:

- **Grade A:** It is strongly recommended that clinicians routinely follow the guideline for eligible patients. There is strong evidence that the practice improves health outcomes.
- **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.
- **Grade CPR:** It is recommended that clinicians consider following the guideline for eligible patients. This recommendation is based on either weak evidence or on the opinions of the Work Group and reviewers that the practice might improve health outcomes.

1c.23 **Grade Assigned to the Recommendation:** A
1c.24 Rationale for Using this Guideline Over Others: No other guidelines are available.

Based on the NQF descriptions for rating the evidence, what was the developer's assessment of the quantity, quality, and consistency of the body of evidence?

1c.25 Quantity: High  
1c.26 Quality: High  
1c.27 Consistency: High

Was the threshold criterion, Importance to Measure and Report, met? 
(1a & 1b must be rated moderate or high and 1c yes)  Yes  
No

Provide rationale based on specific subcriteria:

For a new measure if the Committee votes NO, then STOP.
For a measure undergoing endorsement maintenance, if the Committee votes NO because of 1b. (no opportunity for improvement), it may be considered for continued endorsement and all criteria need to be evaluated.

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### 2. RELIABILITY & VALIDITY - 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)

Measure testing must demonstrate adequate reliability and validity in order to be recommended for endorsement. Testing may be conducted for data elements and/or the computed measure score. Testing information and results should be entered in the appropriate field. Supplemental materials may be referenced or attached in item 2.1. See guidance on measure testing.

S.1 Measure Web Page (In the future, NQF will require measure stewards to provide a URL link to a web page where current detailed specifications can be obtained). Do you have a web page where current detailed specifications for this measure can be obtained? Yes

S.2 If yes, provide web page URL: http://www.arborresearch.org/ESRD_QMS.aspx

2a. RELIABILITY. Precise Specifications and Reliability Testing:  
H  M  L  I

2a1. Precise Measure Specifications. (The measure specifications precise and unambiguous.)

2a1.1 Numerator Statement (Brief, narrative description of the measure focus or what is being measured about the target population, e.g., cases from the target population with the target process, condition, event, or outcome):  
Number of patients in denominator whose delivered dose of hemodialysis (calculated from the last measurements of the month using the UKM or Daugirdas II formula) was a spKt/V >= 1.2.

2a1.2 Numerator Time Window (The time period in which the target process, condition, event, or outcome is eligible for inclusion):  
The entire calendar month.

2a1.3 Numerator Details (All information required to identify and calculate the cases from the target population with the target process, condition, event, or outcome such as definitions, codes with descriptors, and/or specific data collection items/responses):  
The numerator will be determined by counting the patients in the denominator for whom “Kt/V Hemodialysis Method” is “Daugirdas II” OR “UKM” AND “Kt/V” is greater than or equal to 1.2.

2a1.4 Denominator Statement (Brief, narrative description of the target population being measured):  
All adult (>= 18 years old) patients in the sample for analysis who have been on hemodialysis for 90 days or more and dialyzing thrice weekly.

2a1.5 Target Population Category (Check all the populations for which the measure is specified and tested if any):  
Adult/Elderly Care

2a1.6 Denominator Time Window (The time period in which cases are eligible for inclusion):  
The entire calendar month.

2a1.7 Denominator Details (All information required to identify and calculate the target population/denominator such as definitions, codes with descriptors, and/or specific data collection items/responses):
The duration of hemodialysis treatment will be calculated as the difference between the first “Kt/V Collection Date” and “Date Regular Chronic Dialysis Began”. The denominator will include all in-center hemodialysis patients >=18 years old who have been on dialysis for 90 days or longer and “Sessions per Week” is equal to 3. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting month. In-center hemodialysis patients are defined as follows: “Admit Date” to the specified facility is prior or equal to the first day of the study period, AND the patient has not been discharged (“Discharge Date” is null or blank), OR “Discharge Date” from the facility is greater than or equal to the last day of the study period AND “Treatment Dialysis Broad Start Date” is prior or equal to the first day of the study period, AND “Dialysis Broad Type of Treatment” = ‘HD’, AND “Primary Dialysis Setting” = ‘Dialysis Facility/Center’ on the last day of the study period, AND “Date Regular Chronic Dialysis Began” is prior to the first day of the study period.

2a1.8 Denominator Exclusions (Brief narrative description of exclusions from the target population):
Patients on HD less than 90 days; HD patients dialyzing <3 times per week or >3 times per week.

2a1.9 Denominator Exclusion Details (All information required to identify and calculate exclusions from the denominator such as definitions, codes with descriptors, and/or specific data collection items/responses):
Exclusions to this measure include patients who are not receiving dialysis thrice weekly (“Sessions per Week” not equal to 3) and have not been on dialysis at least 90 days. The duration of hemodialysis treatment will be calculated as the difference between the first “Kt/V Collection Date” and “Date Regular Chronic Dialysis Began”.

2a1.10 Stratification Details/Variables (All information required to stratify the measure results including the stratification variables, codes with descriptors, definitions, and/or specific data collection items/responses):
No stratification for this measure.

2a1.11 Risk Adjustment Type (Select type. Provide specifications for risk stratification in 2a1.10 and for statistical model in 2a1.13): No risk adjustment or risk stratification

2a1.12 If “Other,” please describe:

2a1.13 Statistical Risk Model and Variables (Name the statistical method - e.g., logistic regression and list all the risk factor variables. Note - risk model development should be addressed in 2b4.):
Not applicable.

2a1.14-16 Detailed Risk Model Available at Web page URL (or attachment). Include coefficients, equations, codes with descriptors, definitions, and/or specific data collection items/responses. Attach documents only if they are not available on a webpage and keep attached file to 5 MB or less. NQF strongly prefers you make documents available at a Web page URL. Please supply login/password if needed:

2a1.17-18. Type of Score: Rate/proportion

2a1.19 Interpretation of Score (Classifies interpretation of score according to whether better quality is associated with a higher score, a lower score, a score falling within a defined interval, or a passing score): Better quality = Higher score

2a1.20 Calculation Algorithm/Measure Logic (Describe the calculation of the measure score as an ordered sequence of steps including identifying the target population; exclusions; cases meeting the target process, condition, event, or outcome; aggregating data; risk adjustment; etc.):
The duration of hemodialysis treatment will be calculated as the difference between the first “Kt/V Collection Date” and “Date Regular Chronic Dialysis Began”. The denominator will include all in-center hemodialysis patients >=18 years old who have been on dialysis for 90 days or longer and “Sessions per Week” is equal to 3. The patient’s age will be determined by subtracting the patient’s date of birth from the first day of the reporting month. In-center hemodialysis patients are defined as follows: “Admit Date” to the specified facility is prior or equal to the first day of the study period, AND the patient has not been discharged (“Discharge Date” is null or blank), OR “Discharge Date” from the facility is greater than or equal to the last day of the study period AND “Treatment Dialysis Broad Start Date” is prior or equal to the first day of the study period, AND “Dialysis Broad Type of Treatment” = ‘HD’, AND “Primary Dialysis Setting” = ‘Dialysis Facility/Center’ on the last day of the study period, AND “Date Regular Chronic Dialysis Began” is prior to the first day of the study period.
`HD`, AND “Primary Dialysis Setting” = “Dialysis Facility/Center” on the last day of the study period, AND “Date Regular Chronic Dialysis Began” is prior to the first day of the study period. The numerator will be determined by counting the patients in the denominator for whom “Kt/V Hemodialysis Method” is ‘Daugirdas II’ OR ‘UKM’ AND “Kt/V” is greater than or equal to 1.2.

2a1.21-23 **Calculation Algorithm/Measure Logic Diagram URL or attachment:**
Attachment
Appendix C CPM Calculation Flow charts_a 5.pdf

**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):
Not applicable.

**Data Source (Check all the sources for which the measure is specified and tested).** If other, please describe:
Electronic Clinical Data

**Data Source/Data Collection Instrument (Identify the specific data source/data collection instrument, e.g. name of database, clinical registry, collection instrument, etc.):** CROWNWeb

**Data Source/data Collection Instrument Reference Web Page URL or Attachment:** URL
http://www.projectcrownweb.org

2a1.30-32 **Data Dictionary/Code Table Web Page URL or Attachment:**
URL

2a1.33 **Level of Analysis** (Check the levels of analysis for which the measure is specified and tested): Facility

2a1.34-35 **Care Setting** (Check all the settings for which the measure is specified and tested): Dialysis Facility

2a2. **Reliability Testing.** (Reliability testing was conducted with appropriate method, scope, and adequate demonstration of reliability.)

2a2.1 **Data/Sample** (Description of the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included): CROWNWeb Phase II (with Test Batch Submissions). CROWNWeb is currently being released in phases, to allow the immediate collection of data from a limited number of facilities while providing future users an opportunity to complete the required steps to access the system. CMS moved into Phase II of its phase-in implementation process in July 2009. Data were collected from Phase II facilities and test batch submitters on a voluntary basis. These data are not a random national sample of facilities and hence results are not necessarily representative.

CROWNWeb data from July 2009-October 2010 were analyzed. The number of facilities per month ranged from 3415-3453. The total number of patients per month ranged from 263,743 - 330,187.

2a2.2 **Analytic Method** (Describe method of reliability testing & rationale): Reliability was assessed by calculating facility-level month-to-month correlations. Pearson correlation coefficients were calculated between the current performance month and previous month for reporting months July 2009 through October 2010.

2a2.3 **Testing Results** (Reliability statistics, assessment of adequacy in the context of norms for the test conducted): Month-to-month correlations ranged from 0.89 to 0.98, indicating very high month-to-month correlations and reliable data elements for this measure.

See Guidance for Definitions of Rating Scale: H=High; M= Moderate; L=Low; I =Insufficient; NA=Not Applicable
2b. VALIDITY. Validity, Testing, including all Threats to Validity:  

2b1.1 Describe how the measure specifications (measure focus, target population, and exclusions) are consistent with the evidence cited in support of the measure focus (criterion 1c) and identify any differences from the evidence:
The target population in the validity analysis was all ESRD patients on HD who are reported in CROWNWeb in 2009. The population and results from the validity analyses performed were consistent with the evidence provided. The validity analyses showed that relative to facilities with the highest performance scores, the Standardized Mortality Ratio (SMR) increased as performance scores decreased.

2b2. Validity Testing. (Validity testing was conducted with appropriate method, scope, and adequate demonstration of validity.)

2b2.1 Data/Sample (Description of the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included):
2009 CROWNWeb data (July - December) were used to calculate monthly performance scores, and the SMR was calculated using 2009 Medicare-paid dialysis claims and the Medical Evidence Form (Form CMS-2728). Documentation regarding the Medicare claims used to calculate the SMR is attached.

2b2.2 Analytic Method (Describe method of validity testing and rationale; if face validity, describe systematic assessment):
Validity was assessed using Poisson regression models to measure the association between facility level quintiles of performance scores and the 2009 SMR (methodology on SMR calculations is attached). Facility-level performance scores were divided into quintiles and the relative risk (RR) of mortality was calculated for each quintile. The highest quintile was used as the reference group. Thus, a RR>1.0 for the lower performance score quintiles would indicate a higher relative risk of mortality.

2b2.3 Testing Results (Statistical results, assessment of adequacy in the context of norms for the test conducted; if face validity, describe results of systematic assessment):
Quintiles of the performance score are as follows:
Q1: 0-65%
Q2: 65%-79%
Q3: 79%-87%
Q4: 88%-93%
Q5: 93%-100%
Results from the Poisson model indicated lower performance scores were significantly associated with a higher SMR (p<0.01). Relative risks (95% confidence intervals) of mortality for quintiles 1 through 4 were 1.11 (1.07,1.15), 1.13 (1.09, 1.17), 1.08 (1.05,1.12), 1.09 (1.05, 1.13) respectively.

2b3. Measure Exclusions. (Exclusions were supported by the clinical evidence in 1c or appropriately tested with results demonstrating the need to specify them.)

2b3.1 Data/Sample for analysis of exclusions (Description of the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included):
CROWNWeb data from July 2009-October 2010 were analyzed. The number of facilities per month ranged from 3415-3453. The total number of patients per month ranged from 263,743 - 330,187.

2b3.2 Analytic Method (Describe type of analysis and rationale for examining exclusions, including exclusion related to patient preference):
Exclusions for this measure included patients who are not on dialysis thrice weekly and patients who have been on hemodialysis for less than 6 months. The overall mean performance score for this measure was calculated three ways; 1) using data from all patients, 2) using patients with non-missing data, and 3) for patients with non-missing data who qualified as exclusions. Performance scores should improve with the omission of missing values. Furthermore, it is expected that the performance scores for the excluded patients would be lower compared to patients with non-missing data.

2b3.3 Results (Provide statistical results for analysis of exclusions, e.g., frequency, variability, sensitivity analyses):
The overall mean (SD) for all patients was 67% (32%). The mean was 83% (33%) after excluding missing data, and was slightly...
lower for the excluded population 80% (32%).

### 2b4. Risk Adjustment Strategy.
(For outcome measures, adjustment for differences in case mix (severity) across measured entities was appropriately tested with adequate results.)

#### 2b4.1 Data/Sample
(Description of the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included):
No risk adjustment is performed for this measure.

#### 2b4.2 Analytic Method
(Describe methods and rationale for development and testing of risk model or risk stratification including selection of factors/variables):
Not applicable.

#### 2b4.3 Testing Results
(Statistical risk model: Provide quantitative assessment of relative contribution of model risk factors; risk model performance metrics including cross-validation discrimination and calibration statistics, calibration curve and risk decile plot, and assessment of adequacy in the context of norms for risk models. Risk stratification: Provide quantitative assessment of relationship of risk factors to the outcome and differences in outcomes among the strata):
Not applicable.

#### 2b4.4 If outcome or resource use measure is not risk adjusted, provide rationale and analyses to justify lack of adjustment:
Disparities by population group were not observed (see results in Section 1b.4). Furthermore, there is no evidence suggesting this measure should be risk adjusted.

### 2b5. Identification of Meaningful Differences in Performance.
(The performance measure scores were appropriately analyzed and discriminated meaningful differences in quality.)

#### 2b5.1 Data/Sample
(Describe the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included):
Analyses were performed using CROWNWeb data from January 2010. There were 3400 facilities and a total of 293,694 patients in this reporting month. The mean number of patients per facility was 84 (SD=52).

#### 2b5.2 Analytic Method
(Describe methods and rationale to identify statistically significant and practically/meaningfully differences in performance):
Facility-level percent of patients with spKt/V>=1.2 was calculated as the number of patients within the facility on dialysis 6 months or more and dialyzing thrice weekly with spKt/V>=1.2, divided by the total number of patients within the facility. The distribution of facilities within measurement categories was examined.

#### 2b5.3 Results
(Provide measure performance results/scores, e.g., distribution by quartile, mean, median, SD, etc.; identification of statistically significant and meaningfully differences in performance):
Analysis of CROWNWeb data from January 2010 indicate 66% of facilities had at least 70% of patients meeting the spKt/V >=1.2 dose requirement. Only 4% of facilities had 100% of patients meeting the requirements for this measure.

### 2b6. Comparability of Multiple Data Sources/Methods.
(If specified for more than one data source, the various approaches result in comparable scores.)

#### 2b6.1 Data/Sample
(Describe the data or sample including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included):
Multiple data sources were not used.

#### 2b6.2 Analytic Method
(Describe methods and rationale for testing comparability of scores produced by the different data sources specified in the measure):
Not applicable.

#### 2b6.3 Testing Results
(Provide statistical results, e.g., correlation statistics, comparison of rankings; assessment of adequacy in the context of norms for the test conducted):
Not applicable.
2c. Disparities in Care:  

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<th>M</th>
<th>L</th>
<th>I</th>
<th>NA</th>
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(If applicable, the measure specifications allow identification of disparities.)

2c.1 If measure is stratified for disparities, provide stratified results (Scores by stratified categories/cohorts):  
This measure is not stratified.

2c.2 If disparities have been reported/identified (e.g., in 1b), but measure is not specified to detect disparities, please explain:
No disparities have been identified.

2.1-2.3 Supplemental Testing Methodology Information:
URL  

Steering Committee: Overall, was the criterion, Scientific Acceptability of Measure Properties, met?  
Reliability and Validity must be rated moderate or high  
Yes [ ] No [ ]

Provide rationale based on specific subcriteria:

If the Committee votes No, STOP

### 3. USABILITY

Extent to which intended audiences (e.g., consumers, purchasers, providers, policy makers) can understand the results of the measure and are likely to find them useful for decision making. (evaluation criteria)

C.1 Intended Purpose/Use (Check all the purposes and/or uses for which the measure is intended):  
Public Reporting, Quality Improvement (Internal to the specific organization), Quality Improvement with Benchmarking (external benchmarking to multiple organizations)

3.1 Current Use (Check all that apply; for any that are checked, provide the specific program information in the following questions):  
Public Reporting, Quality Improvement (Internal to the specific organization)

3a. Usefulness for Public Reporting:  

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<th>L</th>
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</table>

(The measure is meaningful, understandable and useful for public reporting.)

3a.1. Use in Public Reporting - disclosure of performance results to the public at large (If used in a public reporting program, provide name of program(s), locations, Web page URL(s)):  
If not publicly reported in a national or community program, state the reason AND plans to achieve public reporting, potential reporting programs or commitments, and timeline, e.g., within 3 years of endorsement:  
[For Maintenance – If not publicly reported, describe progress made toward achieving disclosure of performance results to the public at large and expected date for public reporting; provide rationale why continued endorsement should be considered.]

Quality measure results will be evaluated for future public reporting on Medicare’s Dialysis Facility Compare website and for Dialysis Facility Reports.

3a.2. Provide a rationale for why the measure performance results are meaningful, understandable, and useful for public reporting. If usefulness was demonstrated (e.g., focus group, cognitive testing), describe the data, method, and results:  
This measure has been reported in previous ESRD CPM Annual Reports. Healthcare providers and patients can easily understand the meaning of this measure. In addition, there is general acceptance by the Nephrology community of the use of spKt/V for the measurement of dialysis adequacy. Clinical practice guidelines further support this measure. The percent of patients with monthly adequacy measurements improved from 85% in 1998 to 94% in 2007.

3.2 Use for other Accountability Functions (payment, certification, accreditation). If used in a public accountability program, provide name of program(s), locations, Web page URL(s):  
The use of spKt/V to measure dialysis adequacy, with a target of 1.2, is currently in use for a CMS Quality Incentive Payment Demonstration Evaluation, a related project of the CMS Disease Management Demonstration Evaluation.  
### 3b. Usefulness for Quality Improvement: \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

(\text{The measure is meaningful, understandable and useful for quality improvement.})

### 3b.1. Use in QI. If used in quality improvement program, provide name of program(s), locations, Web page URL(s):

[For Maintenance – If not used for QI, indicate the reasons and describe progress toward using performance results for improvement]

The use of spKt/V to measure dialysis adequacy has been used in multiple quality improvement programs. An example is an initiative of ESRD Network 5 to improve performance for achieving spKt/V targets (http://www.esrdnet5.org/adequacyproj.asp). Similarly ESRD Network 18 includes spKt/V for its quality improvement initiatives (http://www.esrdnetwork18.org/pdfs/QI%20Tools%20-%20Forms/2010-2011%20Clinical%20Performance%20Goals-FINAL.pdf).

Also, in previous years, this measure was reported in ESRD CPM Annual Reports. The ESRD CPM Project was a national effort designed to assist dialysis providers to improve patient care and outcomes.

### 3b.2. Provide rationale for why the measure performance results are meaningful, understandable, and useful for quality improvement. If usefulness was demonstrated (e.g., QI initiative), describe the data, method and results:

See 3a.2.

Overall, to what extent was the criterion, Usability, met? \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

Provide rationale based on specific subcriteria:

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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: \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

4a.1-2 How are the data elements needed to compute measure scores generated? (Check all that apply).

Data used in the measure are:

Abstracted from a record by someone other than person obtaining original information (e.g., chart abstraction for quality measure or registry)

#### 4b. Electronic Sources: \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

4b.1 Are the data elements needed for the measure as specified available electronically (Elements that are needed to compute measure scores are in defined, computer-readable fields): ALL data elements in electronic health records (EHRs)

4b.2 If ALL data elements are not from electronic sources, specify a credible, near-term path to electronic capture, OR provide a rationale for using other than electronic sources:

#### 4c. Susceptibility to Inaccuracies, Errors, or Unintended Consequences: \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

4c.1 Identify susceptibility to inaccuracies, errors, or unintended consequences of the measurement identified during testing and/or operational use and strategies to prevent, minimize, or detect. If audited, provide results:

There are no potential barriers to retrieving data necessary for the measure, and there are no data availability issues.

#### 4d. Data Collection Strategy/Implementation: \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

4d.1 Please check if either of the following apply (regarding proprietary measures):

4d.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 and frequency of data collection, sampling, patient confidentiality, time and cost of data collection, other feasibility/implementation issues (e.g., fees for use of proprietary measures):

Since this measure has been collected for several years as part of the CPM project, facilities are familiar with the data required for this measure, and data are readily available. It is unlikely that data elements will be susceptible to inaccuracies, errors, or unintended consequences.

Overall, to what extent was the criterion, Feasibility, met? \[ \text{H} = \text{High}; \text{M} = \text{Moderate}; \text{L} = \text{Low}; \text{I} = \text{Insufficient}; \text{NA} = \text{Not Applicable} \]

Provide rationale based on specific subcriteria:
OVERALL SUITABILITY FOR ENDORSEMENT

<table>
<thead>
<tr>
<th>Does the measure meet all the NQF criteria for endorsement?</th>
<th>Yes</th>
<th>No</th>
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<tbody>
<tr>
<td>Rationale:</td>
<td></td>
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<tr>
<td>If the Committee votes No, STOP.</td>
<td></td>
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<tr>
<td>If the Committee votes Yes, the final recommendation is contingent on comparison to related and competing measures.</td>
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</table>

5. COMPARISON TO RELATED AND COMPETING MEASURES

If a measure meets the above criteria and there are endorsed or new related measures (either the same measure focus or the same target population) or competing measures (both the same measure focus and the same target population), the measures are compared to address harmonization and/or selection of the best measure before a final recommendation is made.

5.1 If there are related measures (either same measure focus or target population) or competing measures (both the same measure focus and same target population), list the NQF # and title of all related and/or competing measures:

- 0247 : Hemodialysis Adequacy Clinical Performance Measure I: Hemodialysis Adequacy- Monthly measurement of delivered dose
- 0248 : Hemodialysis Adequacy Clinical Performance Measure II: Method of Measurement of Delivered Hemodialysis Dose
- 0323 : Hemodialysis Adequacy: Solute
- 1418 : Frequency of Adequacy Measurement for Pediatric Hemodialysis Patients
- 1421 : Method of Adequacy Measurement for Pediatric Hemodialysis Patients
- 1423 : Minimum spKt/V for Pediatric Hemodialysis Patients

5a. Harmonization

5a.1 If this measure has EITHER the same measure focus OR the same target population as NQF-endorsed measure(s): Are the measure specifications completely harmonized?

5a.2 If the measure specifications are not completely harmonized, identify the differences, rationale, and impact on interpretability and data collection burden:

5b. Competing Measure(s)

5b.1 If this measure has both the same measure focus and the same target population as NQF-endorsed measure(s): Describe why this measure is superior to competing measures (e.g., a more valid or efficient way to measure quality); OR provide a rationale for the additive value of endorsing an additional measure. (Provide analyses when possible):

CONTACT INFORMATION

Co.1 Measure Steward (Intellectual Property Owner): Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-01-02, Baltimore, Maryland, 21244-1850

Co.2 Point of Contact: Edward Q., Garcia III, MHS, Health Policy Analyst, MMSNQF@hsag.com, 410-786-6738

Co.3 Measure Developer if different from Measure Steward: Arbor Research/UM-KECC, 340 E. Huron St, Suite 300, Ann Arbor, Michigan, 48104

Co.4 Point of Contact: Claudia, Dahlerus, Claudia.Dahlerus@ArborResearch.org, 734-665-4108

Co.5 Submitter: Thomas, Dudley, Thomas.Dudley@cms.hhs.gov, 410-786-1442, Centers for Medicare & 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. Describe the members’ role in measure development.

Clinical and data technical expert panels (TEPs) were held in September and October 2006, respectively. Since 2006, no TEPs have been held for adult hemodialysis adequacy measures.

**Measure Developer/Steward Updates and Ongoing Maintenance**

Ad.2 If adapted, provide title of original measure, NQF # if endorsed, and measure steward. Briefly describe the reasons for adapting the original measure and any work with the original measure steward:

No changes to this measure are requested.

Ad.3 Year the measure was first released: 2007

Ad.4 Month and Year of most recent revision: 11, 2007

Ad.5 What is your frequency for review/update of this measure? Every 3 years

Ad.6 When is the next scheduled review/update for this measure? 06, 2013

Ad.7 Copyright statement:

Ad.8 Disclaimers:

Ad.9 Additional Information/Comments: On 11/8/11, the denominator specifications changed from requiring patients to be on dialysis for 6 months or more to 90 days or more. Changes were made to the following items in the Specifications section: 2a1.4, 2a1.7, 2a1.8, 2a1.9, and 2a1.20.

Additionally, on 11/8/11, modifications were made the following items in the Importance section: 1c6, 1c7, 1c11, 1c21, 1c22.

**Date of Submission (MM/DD/YY):** 06/23/2011
Hemodialysis Adequacy
CPM III: Minimum Delivered Hemodialysis Dose
Numerator: Number of patients in the denominator whose delivered dose of hemodialysis (calculated from last measurements of the month using the UKM or Daugirdas II formula) was a spKt/V > 1.2 during the study period.
Denominator: All adult HD patients (≥ 18 years old) in the sample for analysis who have been ESRD for 6 months or more and dialyzing three times per week.
Exclusion: Pediatric patients, peritoneal dialysis patients, acute HD, transient dialysis patients (<30 days in this center), kidney transplant patients, patients with ESRD less than 90 days, patients with RRF > 2 ml/min/1.73m² (if measured in the last 3 months),

Start Date of Birth

Date of Birth

Calculate age: studydate - DOB

<18

18+

Calculate # days ESRD: studydate - date of first ESRD

< 6 months

≥ 6 month

Freq. of HD

3X/Wk.

spKt/V

Calculate Measure:
spKt/V ≥ 1.2 = Yes
spKt/V < 1.2 = No

Excluded due to missing/invalid data

Exclude for failing to meet inclusion criteria