



Measure Information

This document contains the information submitted by measure developers/stewards, but is organized according to NQF's measure evaluation criteria and process. The item numbers refer to those in the submission form but may be in a slightly different order here. In general, the item numbers also reference the related criteria (e.g., item 1b.1 relates to sub criterion 1b).

Brief Measure Information

NQF #: 3754

Corresponding Measures:

Measure Title: Risk Standardized Mortality Ratio for Late-Stage Chronic Kidney Disease (CKD) and End Stage Renal Disease (ESRD)

Measure Steward: Centers for Medicare & Medicaid Services

sp.02. Brief Description of Measure:

The Risk Standardized Mortality Ratio for Late-Stage CKD and ESRD (hereafter the "CKD and ESRD Mortality Measure") is an outcome measure to assess how well providers prevent mortality among patients with stage 4 or 5 CKD or ESRD. This measure assesses nephrology practices (sometimes referred to as "providers" in this submission) who care for adult Medicare Fee-for-Service (FFS) beneficiaries with late-stage CKD and ESRD.

The CKD and ESRD Mortality Measure originated as a re-specification of the National Quality Forum (NQF)-endorsed Standardized Mortality Ratio for Dialysis Facilities Measure (NQF #0369), and is being submitted to NQF as a new measure due to the substantive nature of the changes. While the Standardized Mortality Ratio for Dialysis Facilities Measure assessed dialysis facilities and only included patients with ESRD, the CKD and ESRD Mortality Measure expanded the measure cohort and to additionally include patients with stage 4 or 5 CKD, and expands the measured entities to include nephrology practices more broadly. The risk model has been changed to accommodate the updated setting and cohort.

1b.01. Developer Rationale:

sp.12. Numerator Statement: The measure outcome is all-cause mortality within the measurement year. Mortality is defined as death for any reason within the measurement period for patients age 19 and older with Stage 4 CKD, Stage 5 CKD, or ESRD at risk during the measurement period. Hospice enrollment is a censoring event and mortality after enrollment is not counted in the outcome.

sp.14. Denominator Statement:

The cohort includes Medicare Fee-For-Service beneficiaries (patients) who are 19 years and older with Stage 4 CKD, Stage 5 CKD, or ESRD and who are being treated by a nephrology practice. Patients are not included if they are enrolled in Medicare hospice, or have had a kidney transplant within the past 12 months.

The measure uses a time-to-event methodology in which the denominator represents the person-time in which eligible patients are at risk for the outcome. The observed patient-level denominator is the amount of eligible

person-time for which the patient is at risk for the outcome. The observed provider-level denominator is the total person-time of eligible patients attributed to the provider.

sp.16. Denominator Exclusions: The measure excludes patients with metastatic and advanced cancers, defined as specific cancer-related ICD-10 codes from an inpatient encounter. A full list of codes is available in the attached data dictionary.

Measure Type: Outcome

sp.28. Data Source:

Claims

Other (specify)

Patient Enrollment data including the hospice enrollment, ESRD/dialysis, and vital statistics

sp.07. Level of Analysis:

Clinician: Group/Practice

IF Endorsement Maintenance – Original Endorsement Date:

Most Recent Endorsement Date:

IF this measure is included in a composite, NQF Composite#/title:

IF this measure is paired/grouped, NQF#/title:

sp.03. IF PAIRED/GROUPED, what is the reason this measure must be reported with other measures to appropriately interpret results?:

1. Importance to Measure and Report

Extent to which the specific measure focus is evidence-based, important to making significant gains in healthcare quality, and improving health outcomes for a specific high-priority (high-impact) aspect of healthcare where there is variation in or overall less-than-optimal performance. Measures must be judged to meet all sub criteria to pass this criterion and be evaluated against the remaining criteria

Please separate added or updated information from the most recent measure evaluation within each question response in the Importance to Measure and Report: Evidence section. For example:

Current Submission:

Updated evidence information here.

Previous (Year) Submission:

Evidence from the previous submission here.

1a.01. Provide a logic model.

Briefly describe the steps between the healthcare structures and processes (e.g., interventions, or services) and the patient's health outcome(s). The relationships in the diagram should be easily understood by general, non-technical audiences. Indicate the structure, process or outcome being measured.

[Response Begins]

[Response Ends]

1a.02. Provide evidence that the target population values the measured outcome, process, or structure and finds it meaningful.

Describe how and from whom input was obtained.

[Response Begins]

[Response Ends]

1a.03. Provide empirical data demonstrating the relationship between the outcome (or PRO) and at least one healthcare structure, process, intervention, or service.

[Response Begins]

[Response Ends]

1b.01. Briefly explain the rationale for this measure.

Explain how the measure will improve the quality of care, and list the benefits or improvements in quality envisioned by use of this measure.

[Response Begins]

[Response Ends]

1b.02. Provide performance scores on the measure as specified (current and over time) at the specified level of analysis.

Include mean, std dev, min, max, interquartile range, and scores by decile. Describe the data source including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities include. This information also will be used to address the sub-criterion on improvement (4b) under Usability and Use.

[Response Begins]

[Response Ends]

1b.03. If no or limited performance data on the measure as specified is reported above, then provide a summary of data from the literature that indicates opportunity for improvement or overall less than optimal performance on the specific focus of measurement. Include citations.

[Response Begins]

[Response Ends]

1b.04. Provide disparities data from the measure as specified (current and over time) by population group, e.g., by race/ethnicity, gender, age, insurance status, socioeconomic status, and/or disability.

Describe the data source including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included. Include mean, std dev, min, max, interquartile range, and scores by decile. For measures that show high levels of performance, i.e., “topped out”, disparities data may demonstrate an opportunity for improvement/gap in care for certain sub-populations. This information also will be used to address the sub-criterion on improvement (4b) under Usability and Use.

[Response Begins]

[Response Ends]

1b.05. If no or limited data on disparities from the measure as specified is reported above, then provide a summary of data from the literature that addresses disparities in care on the specific focus of measurement. Include citations. Not necessary if performance data provided in above.

[Response Begins]

[Response Ends]

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. Measures must be judged to meet the sub criteria for both reliability and validity to pass this criterion and be evaluated against the remaining criteria.

sp.01. Provide the measure title.

Measure titles should be concise yet convey who and what is being measured (see [What Good Looks Like](#)).

[Response Begins]

Risk Standardized Mortality Ratio for Late-Stage Chronic Kidney Disease (CKD) and End Stage Renal Disease (ESRD)

[Response Ends]

sp.02. Provide a brief description of the measure.

Including type of score, measure focus, target population, timeframe, (e.g., Percentage of adult patients aged 18-75 years receiving one or more HbA1c tests per year).

[Response Begins]

The Risk Standardized Mortality Ratio for Late-Stage CKD and ESRD (hereafter the “CKD and ESRD Mortality Measure”) is an outcome measure to assess how well providers prevent mortality among patients with stage 4 or 5 CKD or ESRD. This measure assesses nephrology practices (sometimes referred to as "providers" in this submission) who care for adult Medicare Fee-for-Service (FFS) beneficiaries with late-stage CKD and ESRD.

The CKD and ESRD Mortality Measure originated as a re-specification of the National Quality Forum (NQF)-endorsed Standardized Mortality Ratio for Dialysis Facilities Measure (NQF #0369), and is being submitted to NQF as a new measure due to the substantive nature of the changes. While the Standardized Mortality Ratio for Dialysis Facilities Measure assessed dialysis facilities and only included patients with ESRD, the CKD and ESRD Mortality Measure expanded the measure cohort and to additionally include patients with stage 4 or 5 CKD, and expands the measured entities to include nephrology practices more broadly. The risk model has been changed to accommodate the updated setting and cohort.

[Response Ends]

sp.04. Check all the clinical condition/topic areas that apply to your measure, below.

Please refrain from selecting the following answer option(s). We are in the process of phasing out these answer options and request that you instead select one of the other answer options as they apply to your measure.

Please do not select:

- Surgery: General

[Response Begins]

Renal: Chronic Kidney Disease (CKD)

Renal: End Stage Renal Disease (ESRD)

[Response Ends]

sp.05. Check all the non-condition specific measure domain areas that apply to your measure, below.

[Response Begins]

Access to Care

Care Coordination

Health and Functional Status

[Response Ends]

sp.06. Select one or more target population categories.

Select only those target populations which can be stratified in the reporting of the measure's result.

Please refrain from selecting the following answer option(s). We are in the process of phasing out these answer options and request that you instead select one of the other answer options as they apply to your measure.

Please do not select:

- *Populations at Risk: Populations at Risk*

[Response Begins]

Adults (Age >= 18)

Populations at Risk: Populations at Risk

[Response Ends]

sp.07. Select the levels of analysis that apply to your measure.

Check ONLY the levels of analysis for which the measure is SPECIFIED and TESTED.

Please refrain from selecting the following answer option(s). We are in the process of phasing out these answer options and request that you instead select one of the other answer options as they apply to your measure.

Please do not select:

- *Clinician: Clinician*
- *Population: Population*

[Response Begins]

Clinician: Group/Practice

[Response Ends]

sp.08. Indicate the care settings that apply to your measure.

Check ONLY the settings for which the measure is SPECIFIED and TESTED.

[Response Begins]

Ambulatory Care

[Response Ends]

sp.09. Provide a URL link to a web page specific for this measure that contains current detailed specifications including code lists, risk model details, and supplemental materials.

Do not enter a URL linking to a home page or to general information. If no URL is available, indicate "none available".

[Response Begins]

none available

[Response Ends]

sp.12. Attach the data dictionary, code table, or value sets (and risk model codes and coefficients when applicable). Excel formats (.xlsx or .csv) are preferred.

Attach an excel or csv file; if this poses an issue, [contact staff](#). Provide descriptors for any codes. Use one file with multiple worksheets, if needed.

[Response Begins]

Available in attached Excel or csv file

[Response Ends]

Attachment: 3754_DataDictionary_MortalityCKDESRD_01052023.xlsx

For the question below: state the outcome being measured. Calculation of the risk-adjusted outcome should be described in sp.22.

sp.13. State the numerator.

Brief, narrative description of the measure focus or what is being measured about the target population, i.e., cases from the target population with the target process, condition, event, or outcome).

DO NOT include the rationale for the measure.

[Response Begins]

The measure outcome is all-cause mortality within the measurement year. Mortality is defined as death for any reason within the measurement period for patients age 19 and older with Stage 4 CKD, Stage 5 CKD, or ESRD at risk during the measurement period. Hospice enrollment is a censoring event and mortality after enrollment is not counted in the outcome.

[Response Ends]

For the question below: describe how the observed outcome is identified/counted. Calculation of the risk-adjusted outcome should be described in sp.22.

sp.14. Provide details needed to calculate the numerator.

All information required to identify and calculate the cases from the target population with the target process, condition, event, or outcome such as definitions, time period for data collection, specific data collection items/responses, code/value sets.

Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at sp.11.

[Response Begins]

The observed outcome at the patient level is death of a patient from any cause during the measurement period (provided the patient is eligible for the cohort and considered “at risk” for the outcome, as discussed below in Sp.15 “State the denominator”). The observed outcome at the provider level is the total number of deaths among “at risk” patients included the measure who are attributed to that provider.

We identify deaths for Medicare FFS beneficiaries who are 19 years and older in the Medicare Enrollment Database (EDB). We use the date of death in the enrollment database (EDB) which is derived from the Social Security Administration and has been verified.

There is one censoring event that is not counted toward the mortality outcome, discussed in greater detail in sp.16 (denominator details). Patients who enroll in Medicare hospice are considered eligible for the outcome up to the point of hospice enrollment, at which point they are no longer “at risk” and contribute no further person-time (that is, mortality post-enrollment is not counted toward the outcome).

[Response Ends]

For the question below: state the target population for the outcome. Calculation of the risk-adjusted outcome should be described in sp.22.

sp.15. State the denominator.

Brief, narrative description of the target population being measured.

[Response Begins]

The cohort includes Medicare Fee-For-Service beneficiaries (patients) who are 19 years and older with Stage 4 CKD, Stage 5 CKD, or ESRD and who are being treated by a nephrology practice. Patients are not included if they are enrolled in Medicare hospice, or have had a kidney transplant within the past 12 months.

The measure uses a time-to-event methodology in which the denominator represents the person-time in which eligible patients are at risk for the outcome. The observed patient-level denominator is the amount of eligible person-time for which the patient is at risk for the outcome. The observed provider-level denominator is the total person-time of eligible patients attributed to the provider.

[Response Ends]

For the question below: describe how the target population is identified. Calculation of the risk-adjusted outcome should be described in sp.22.

sp.16. Provide details needed to calculate the denominator.

All information required to identify and calculate the target population/denominator such as definitions, time period for data collection, specific data collection items/responses, code/value sets.

Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at sp.11.

[Response Begins]

This measure includes Medicare FFS patients:

1. With stage 4 or 5 CKD or ESRD during the measurement period, and
2. Who are age 18 or older in the year prior to the measurement period (that is, age 19 or older in the measurement period), and
3. With continuous enrollment in Medicare FFS Parts A and B for one full year prior to the performance year as well as the full performance year, or up to the date at which they are censored (due to hospice enrollment) or die in the performance year, and
4. Attributed to a nephrology practice (definition described below).

Stage 4 and 5 CKD patients are defined as those with at least one occurrence of International Classification of Diseases, 10th Revision (ICD-10) code N18.4 “Chronic kidney disease, Stage 4 (Severe)” or N18.5 “Chronic kidney disease, Stage 5” in at least one claim during the performance year. Patients with ESRD are defined as those enrolled in Medicare ESRD or ESRD-Dialysis coverage.

This measure does not include:

1. Patients who are enrolled in the Medicare hospice program, or
2. Patients who have had a kidney transplant within the 12 months prior to becoming otherwise eligible for the cohort.

Hospice patients are identified from the Medicare Enrollment Database. Kidney transplants are identified as a patient with one of the codes in Table sp.16:1 below.

Code	Description of Code
OTY00Z0	Transplantation of Right Kidney, Allogeneic, Open Approach
OTY00Z1	Transplantation of Right Kidney, Syngeneic, Open Approach
OTY00Z2	Transplantation of Right Kidney, Zooplastic, Open Approach
OTY10Z0	Transplantation of Left Kidney, Allogeneic, Open Approach
OTY10Z1	Transplantation of Left Kidney, Syngeneic, Open Approach
OTY10Z2	Transplantation of Left Kidney, Zooplastic, Open Approach
50360-50365	Kidney transplant
50380	Kidney transplant
S2065	Kidney transplant

Table sp.16:1. Transplant Codes, from Agency for Healthcare Research and Quality Clinical Classifications Software (AHRQ CCS) 105

Patient attribution: Nephrology practices responsible for patient care are defined as those having at least two encounters with that patient, providing specialty services (with specialty code 39) during the performance year.

Specifically, we first identified all the nephrology practices that provided any nephrology specialty services (with specialty code 39) during the performance year to a given patient. Eligible patient visits were defined as those with those nephrology practices by specific Healthcare Common Procedure Coding System (HCPCS) codes prescribed by the Kidney Care Choices Model, listed in Tables sp.16:2-3. These tables include eligible Evaluation and Management Coding (E&M) services and/or received monthly capitation payments (MCP) for ESRD/dialysis services. If a patient visited multiple practices that provide specialty care, the patient is attributed, 1) to the practice that provided highest number of E&M claims or MCP claims to the patient; or if there is a tie, 2) to the practice that billed the most for those services; or there is still a tie, 3) the practice who provided the most recent service; or if there is still a tie, 4) a randomly selected practice. This approach aligns with the CMS Kidney Care Choices Model strategy to identify nephrology practices¹.

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Service	CPT /HCPCS Codes
Office/Outpatient Visit E/M	99201-99205 99211-99215
Prolonged E/M	99354-99355
Transitional Care Management Services	99495-99496
Advance Care Planning	99497-99498
Welcome to Medicare and Annual Wellness Visits	G0402, G0438, G0439
Chronic Care Management Services	99490

Table sp.16:2. E&M HCPCS Codes Identifying Providers Who Delivered Nephrology Specialty Services

Service	CPT Codes
ESRD related services monthly, for patients 12-19	90957-90959
ESRD related services monthly, for patients 20 years of age and older	90960-90962
End-Stage Renal Disease Services	90965, 90966

Table sp.16:3. MCP HCPCS Codes Identifying Providers Who Delivered Nephrology Specialty Services

Denominator calculation (time at risk for included cohort): The raw denominator is the sum of at-risk time for the cohort. The at-risk time from each patient is calculated from when the patient becomes eligible for the cohort in the measurement period (detailed below) until the earliest time of either: death; enrollment in Medicare hospice; or end of the measurement period.

As noted above, patients must have 12 months of claims data prior to the performance period to be eligible. Patients are included in the cohort once they both 1) have Stage 4 or 5 CKD or ESRD enrollment and 2) are attributed to a nephrology practice in the performance year. A patient becomes eligible for the measure and their “at risk” period begins once both of these conditions are met within the performance year.

The measure considers the length of time a patient is eligible (at risk) for the outcome. As noted above, patients must have 12 months of claims data prior to the performance period and at least one Stage 4 or 5 CKD claim or ESRD enrollment during the measurement year to be eligible. An eligible patient will enter the cohort and begin contributing at-risk time once they are both attributed to a nephrology practice and have a confirmed diagnosis of Stage 4 or 5 CKD or ESRD enrollment, either:

- The beginning of the performance year (January 1) if the patient had both Stage 4 or 5 CKD or ESRD and had two encounters with their attributed nephrology practice in the previous year, or
- During the performance year on the earliest date at which the patient is both 1) attributed to a nephrology practice (as described above) and 2) has a Stage 4 or 5 CKD diagnosis or ESRD enrollment

An included patient will leave the cohort (stop contributing at-risk time) on the first date of the following:

- Date of death (outcome event)
- Date of enrollment in Medicare hospice
 - Censoring event: not counted in the outcome; no longer eligible for the outcome for the remainder of the measurement year.
- End of the measurement year

The total person-time contributed by a patient is the span between the date of entering the cohort and the date of leaving the cohort.

Note: Because the first three months of chronic dialysis entail a high risk of mortality, if a patient dies within that period, the outcome is attributed to the provider with the plurality of pre-dialysis E&M services.

References:

1. Request for Applications (RFA): Kidney Care Choices (KCC) Model. 2019. Centers for Medicare & Medicaid Services (CMS), Center for Medicare and Medicaid Innovation (CMMI). <https://innovation.cms.gov/files/x/kcc-rfa.pdf>

[Response Ends]

sp.17. Describe the denominator exclusions.

Brief narrative description of exclusions from the target population.

[Response Begins]

The measure excludes patients with metastatic and advanced cancers, defined as specific cancer-related ICD-10 codes from an inpatient encounter. A full list of codes is available in the attached data dictionary.

[Response Ends]

sp.18. Provide details needed to calculate the denominator exclusions.

All information required to identify and calculate exclusions from the denominator such as definitions, time period for data collection, specific data collection items/responses, code/value sets – Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at sp.11.

[Response Begins]

Patients with metastatic and advanced cancers are excluded from the measure. We identify these patients using ICD-10 codes from inpatient claims from the year prior to the measurement year. Specific ICD-10 codes are derived from the following CMS Condition Categories: CC8, CC10, CC12, CC177, CC178; a full list of codes is available in the attached data dictionary.

Rationale: The outcome for these patients is likely more influenced by cancer treatment than care associated with their chronic kidney disease or nephrologist. Additionally, many patients in this population may be too ill for dialysis and have a high risk of mortality separately from their CKD or ESRD condition; thus, it is not appropriate to attribute outcomes for these patients to their nephrologists' quality of care.

[Response Ends]

sp.19. Provide all information required to stratify the measure results, if necessary.

Include the stratification variables, definitions, specific data collection items/responses, code/value sets, and the risk-model covariates and coefficients for the clinically-adjusted version of the measure when appropriate. Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format in the Data Dictionary field.

[Response Begins]

This measure is not currently stratified.

[Response Ends]

sp.20. Is this measure adjusted for socioeconomic status (SES)?

[Response Begins]

No

[Response Ends]

sp.21. Select the risk adjustment type.

Select type. Provide specifications for risk stratification and/or risk models in the Scientific Acceptability section.

[Response Begins]

Statistical risk model with risk factors (specify number of risk factors)

[Statistical risk model with risk factors (specify number of risk factors) Please Explain]

71 risk factors

[Response Ends]

sp.22. Select the most relevant type of score.

Attachment: If available, please provide a sample report.

[Response Begins]

Ratio

[Response Ends]

sp.23. Select the appropriate interpretation of the measure score.

Classifies interpretation of score according to whether better quality or resource use is associated with a higher score, a lower score, a score falling within a defined interval, or a passing score

[Response Begins]

Better quality = Lower score

[Response Ends]

sp.24. Diagram or describe the calculation of the measure score as an ordered sequence of steps.

Identify the target population; exclusions; cases meeting the target process, condition, event, or outcome; time period of data, aggregating data; risk adjustment; etc.

[Response Begins]

Calculation algorithm:

First, identify the cohort by those patients meeting all inclusion criteria:

- Patient is enrolled in Medicare FFS Parts A and B for one full year prior to the performance year, as well as the full performance year or until the date of outcome (death) or censoring (due to hospice enrollment) in the performance year,
- Patient is at least 18 years old at the start of the year prior to the performance year (that is, at least 19 years old at the start of the performance year),
- Patient has Stage 4 or 5 CKD or ESRD:
 - At least one occurrence of ICD-10 code N18.4 (CKD, Stage 4) or N18.5 (CKD, Stage 5) in at least one claim during the performance year, or

- Enrolled in either ESRD or ESRD for Dialysis Medicare coverage for at least one day in the measurement period,
- Patient is not already enrolled in hospice, and
- If patient had a prior kidney transplant, at least one year has passed post-transplant.

Second, apply exclusion to the cohort:

- Metastatic and advanced cancers, defined as specific cancer-related ICD-10 codes from an inpatient encounter. Patients are excluded if coded with advanced or metastatic cancer within one year prior to the earliest date of: being attributed to a nephrology practice; being diagnosed with stage 4 or 5 CKD; or enrollment in ESRD or ESRD-dialysis.

Third, attribute patients to nephrology practices as detailed above in section sp.16, denominator details.

Fourth, calculate the outcome. The measure uses a time-to-event outcome, which incorporates not only whether mortality occurred, but also the total “at risk” time for each patient in the cohort during the performance year. A Cox proportional hazard frailty model is used to calculate the measure score. The start time for each patient is either the beginning of the performance year if there is at least one Stage 4 or 5 CKD claim or ESRD enrollment period observed in the prior year, or else when the patient becomes eligible for the cohort in the performance year. The end time for each patient is the first of the date of death, date of hospice enrollment, or the end of the performance year. The precise methodology for cohort eligibility detailed in sp.16 “Details needed to calculate the denominator.”

Measure score calculation method for time-to-event outcome:

Assume that the hazard function of an event for patient i serviced by provider j , with a vector of risk factors X_{ij} is defined as a frailty model under the proportional hazard framework:

$$h_{ij}(t_{ij}) = w_j h_0(t_{ij}) \exp(X_{ij}\beta)$$

where the w_j is the frailty for each provider j (that is, the provider-level hazard effect).

So, for the patient ij , define the predicted probability of progression (that is, predicted for a patient with the same clinical risk factors with that specific nephrologist) at time t as cumulative hazard at the time t_{ij}

$$P_{ij} = H_{ij}(t_{ij}) = \int_0^{t_{ij}} w_j h_0(t) \exp(X_{ij}\beta) dt = w_j \exp(X_{ij}\beta) \int_0^{t_{ij}} h_0(t) dt = w_j \exp(X_{ij}\beta) H_0(t_{ij})$$

Correspondingly, we define the expected probability of progression (that is, expected for a patient with the same clinical risk factors with a nephrologist of median quality) by setting $w_j = 1$ as:

$$E_{ij} = \exp(X_{ij}\beta) H_0(t_{ij})$$

The risk-standardized ratio (RSR) in a frailty model for provider j will simply be the frailty estimate w_j for provider j since

$$RSR_j = \frac{\text{predicted number of events}}{\text{expected number of events}} = \frac{\sum_{i=1}^{n_j} P_{ij}}{\sum_{i=1}^{n_j} E_{ij}} = \frac{w_j \sum_{i=1}^{n_j} \exp(X_{ij}\beta) H_0(t_{ij})}{\sum_{i=1}^{n_j} \exp(X_{ij}\beta) H_0(t_{ij})} = w_j$$

where n_j is the number of patients seeing provider j .

The frailty estimate (that is, the ratio of predicted to expected mortality hazard) is distributed according to a lognormal distribution, $\log(w_j) \sim N(0, \theta)$, where $\text{median}(w_j) = 1$. The 95% confidence interval for $RSMR_j$ (the frailty) will be a direct output from estimation software. For a given nephrology practice, an RSMR of precisely 1 indicates median performance; an RSMR greater than 1 indicates a higher risk of death (and therefore worse performance) than expected while an RSMR less than 1 indicates lower risk (better performance).

There is also an option to convert the measure score from a ratio to a rate. A Risk Standardized Incidence Rate (RSIR) may be calculated as $RSIR_j = RSMR_j * IR$, where the constant IR is the national incidence rate per 100 patient-years (calculated as 100 times the total number of deaths divided by the total patient-years). A nephrology practice with an RSMR of 1 (median performance) would have an RSIR equal to IR (the overall national rate); similarly an RSIR greater than IR indicates a higher rate of death (worse performance) and RSIR less than IR

indicates a lower rate of death (better performance). As RSMR and RSIR are directly proportional, the choice of score does not affect providers' relative performance, only the interpretation of the numeric scores. We have elected to report the RSMR (ratio) here as the centering of the distribution around 1 more clearly highlights providers' relative performance.

[Response Ends]

sp.27. If measure testing is based on a sample, provide instructions for obtaining the sample and guidance on minimum sample size.

Examples of samples used for testing:

- *Testing may be conducted on a sample of the accountable entities (e.g., hospital, physician). The analytic unit specified for the particular measure (e.g., physician, hospital, home health agency) determines the sampling strategy for scientific acceptability testing.*
- *The sample should represent the variety of entities whose performance will be measured. The [2010 Measure Testing Task Force](#) recognized that the samples used for reliability and validity testing often have limited generalizability because measured entities volunteer to participate. Ideally, however, all types of entities whose performance will be measured should be included in reliability and validity testing.*
- *The sample should include adequate numbers of units of measurement and adequate numbers of patients to answer the specific reliability or validity question with the chosen statistical method.*
- *When possible, units of measurement and patients within units should be randomly selected.*

[Response Begins]

Not applicable. The measure is not based on a sample or survey.

[Response Ends]

sp.30. Select only the data sources for which the measure is specified.

[Response Begins]

Claims

Other (specify)

[Other (specify) Please Explain]

Patient Enrollment data including the hospice enrollment, ESRD/dialysis, and vital statistics

[Response Ends]

sp.31. Identify the specific data source or data collection instrument.

For example, provide the name of the database, clinical registry, collection instrument, etc., and describe how data are collected.

[Response Begins]

Data sources for the measure:

Medicare Part A inpatient and Part B outpatient claims: This data source contains claims data for FFS inpatient and outpatient services including: Medicare inpatient hospital care, outpatient hospital services, as well as inpatient and outpatient physician claims for the 12 months prior to eligibility for the cohort.

Medicare Enrollment Database (EDB): This database contains Medicare beneficiary demographic, benefit/coverage, and vital status information. This data source was used to obtain information on several inclusion/exclusion indicators such as Medicare enrollment in ESRD and ESRD-Dialysis, hospice, and vital status. These data have previously been shown to accurately reflect patient vital status¹.

Reference:

1. Fleming C., Fisher ES, Chang CH, Bubolz D, Malenda J. Studying outcomes and hospital utilization in the elderly: The advantages of a merged data base for Medicare and Veterans Affairs Hospitals. Medical Care. 1992; 30(5): 377-91.

[Response Ends]

sp.32. Provide the data collection instrument.

[Response Begins]

No data collection instrument provided

[Response Ends]

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 fields in the Scientific Acceptability sections of the Measure Submission Form.

- Measures must be tested for all the data sources and levels of analyses that are specified. If there is more than one set of data specifications or more than one level of analysis, contact NQF staff about how to present all the testing information in one form.
- All required sections must be completed.
- For composites with outcome and resource use measures, Questions 2b.23-2b.37 (Risk Adjustment) also must be completed.
- If specified for multiple data sources/sets of specifications (e.g., claims and EHRs), Questions 2b.11-2b.13 also must be completed.
- An appendix for supplemental materials may be submitted (see Question 1 in the Additional section), but there is no guarantee it will be reviewed.
- Contact NQF staff with any questions. Check for resources at the [Submitting Standards webpage](#).
- For information on the most updated guidance on how to address social risk factors variables and testing in this form refer to the release notes for the [2021 Measure Evaluation Criteria and Guidance](#).

Note: The information provided in this form is intended to aid the Standing Committee and other stakeholders in understanding to what degree the testing results for this measure meet NQF's evaluation criteria for testing.

2a. Reliability testing demonstrates the measure data elements are repeatable, producing the same results a high proportion of the time when assessed in the same population in the same time period and/or that the measure score is precise. For instrument-based measures (including PRO-PMs) and composite performance measures, reliability should be demonstrated for the computed performance score.

2b1. Validity testing demonstrates that the measure data elements are correct and/or the measure score correctly reflects the quality of care provided, adequately identifying differences in quality. For instrument based measures (including PRO-PMs) and composite performance measures, validity should be demonstrated for the computed performance score.

2b2. Exclusions are supported by the clinical evidence and are of sufficient frequency to warrant inclusion in the specifications of the measure;

AND

If patient preference (e.g., informed decision-making) is a basis for exclusion, there must be evidence that the exclusion impacts performance on the measure; in such cases, 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).

2b3. For outcome measures and other measures when indicated (e.g., resource use):

- an evidence-based risk-adjustment strategy (e.g., risk models, risk stratification) is specified; is based on patient factors (including clinical and social risk factors) that influence the measured outcome and are present at start of care; 14,15 and has demonstrated adequate discrimination and calibration
- rationale/data support no risk adjustment/ stratification.

2b4. Data analysis of computed measure scores demonstrates that methods for scoring and analysis of the specified measure allow for identification of statistically significant and practically/clinically meaningful 16 differences in performance;

OR

there is evidence of overall less-than-optimal performance.

2b5. If multiple data sources/methods are specified, there is demonstration they produce comparable results.

2b6. Analyses identify the extent and distribution of missing data (or nonresponse) and demonstrate that performance results are not biased due to systematic missing data (or differences between responders and non-responders) and how the specified handling of missing data minimizes bias.

2c. For composite performance measures, empirical analyses support the composite construction approach and demonstrate that:

2c1. the component measures fit the quality construct and add value to the overall composite while achieving the related objective of parsimony to the extent possible; and

2c2. the aggregation and weighting rules are consistent with the quality construct and rationale while achieving the related objective of simplicity to the extent possible.

(if not conducted or results not adequate, justification must be submitted and accepted)

Definitions

Reliability testing applies to both the data elements and computed measure score. Examples of reliability testing for data elements 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 of the measure score addresses precision of measurement (e.g., signal-to-noise).

Validity testing applies to both the data elements and computed measure score. Validity testing of data elements typically analyzes agreement with another authoritative source of the same information. Examples of validity testing of the measure score include, but are not limited to: testing hypotheses that the measures scores indicate quality of care, e.g., measure scores are different for groups known to have differences in quality assessed by another valid quality measure or method; correlation of measure scores with another valid indicator of quality for the specific topic; or relationship to conceptually related measures (e.g., scores on process measures to scores on outcome measures). Face validity of the measure score as a quality indicator may be adequate if accomplished through a systematic and transparent process, by identified experts, and explicitly addresses whether performance

scores resulting from the measure as specified can be used to distinguish good from poor quality. The degree of consensus and any areas of disagreement must be provided/discussed.

Examples of evidence that an exclusion distorts measure results include, but are not limited to: frequency of occurrence, variability of exclusions across providers, and sensitivity analyses with and without the exclusion.

Patient preference is not a clinical exception to eligibility and can be influenced by provider interventions.

Risk factors that influence outcomes should not be specified as exclusions.

With large enough sample sizes, small differences that are statistically significant may or may not be practically or clinically meaningful. The substantive question may be, for example, whether a statistically significant difference of one percentage point in the percentage of patients who received smoking cessation counseling (e.g., 74 percent v. 75 percent) is clinically meaningful; or whether a statistically significant difference of \$25 in cost for an episode of care (e.g., \$5,000 v. \$5,025) is practically meaningful. Measures with overall less-than-optimal performance may not demonstrate much variability across providers.

Please separate added or updated information from the most recent measure evaluation within each question response in the Scientific Acceptability sections. For example:

Current Submission:

Updated testing information here.

Previous (Year) Submission:

Testing from the previous submission here.

2a.01. Select only the data sources for which the measure is tested.

[Response Begins]

Claims

Other (specify)

[Other (specify) Please Explain]

Beneficiary Enrollment data including the hospice enrollment, ESRD/dialysis enrollment and vital statistics

[Response Ends]

2a.02. If an existing dataset was used, identify the specific dataset.

The dataset used for testing must be consistent with the measure specifications for target population and healthcare entities being measured; e.g., Medicare Part A claims, Medicaid claims, other commercial insurance, nursing home MDS, home health OASIS, clinical registry).

[Response Begins]

Multiple datasets were used for these analyses. See section 2a.07 for additional details.

[Response Ends]

2a.03. Provide the dates of the data used in testing.

Use the following format: "MM-DD-YYYY - MM-DD-YYYY"

[Response Begins]

Dates of data vary by dataset. Please see section 2a.07 for details.

[Response Ends]

2a.04. Select the levels of analysis for which the measure is tested.

Testing must be provided for all the levels specified and intended for measure implementation, e.g., individual clinician, hospital, health plan.

Please refrain from selecting the following answer option(s). We are in the process of phasing out these answer options and request that you instead select one of the other answer options as they apply to your measure.

Please do not select:

- Clinician: Clinician
- Population: Population

[Response Begins]

Clinician: Group/Practice

[Response Ends]

2a.05. List the measured entities included in the testing and analysis (by level of analysis and data source).

Identify the number and descriptive characteristics of measured entities included in the analysis (e.g., size, location, type); if a sample was used, describe how entities were selected for inclusion in the sample.

[Response Begins]

In testing, measured entities are any clinician groups billing for nephrology services to Medicare FFS patients 18 years or older, grouped by taxpayer identification number (TIN) and identified through Medicare FFS administrative claims data. There were 3,009 measured entities.

[Response Ends]

2a.06. Identify the number and descriptive characteristics of patients included in the analysis (e.g., age, sex, race, diagnosis), separated by level of analysis and data source; if a sample was used, describe how patients were selected for inclusion in the sample.

If there is a minimum case count used for testing, that minimum must be reflected in the specifications.

[Response Begins]

The number of patients varied by dataset; see 2a.07 for details.

[Response Ends]

2a.07. If there are differences in the data or sample used for different aspects of testing (e.g., reliability, validity, exclusions, risk adjustment), identify how the data or sample are different for each aspect of testing.

[Response Begins]

#3754 Risk Standardized Mortality Ratio for Late-Stage Chronic Kidney Disease (CKD) and End Stage Renal Disease (ESRD) , Submission Last Updated: Jan 09, 2023

Dataset	Applicable Section	Description of Dataset
Mortality Development Dataset (Medicare Fee-For-Service Administrative Claims Data)	Section 2a.09 Reliability Testing Section 2b.01 Validity Testing Section 2b.05 Meaningful Differences Section 2b.30 Risk Adjustment/Stratification Section 2b.16 Testing of Measure Exclusions 2b.20 Statistical Risk Model Discrimination Statistics 2b.28 Statistical Risk Model Calibration Statistics	Dates of data: January 1, 2017 – December 31, 2018 Number of patients in the dataset: 758,162 See table 2a.07:2 below for patient characteristics. Number of measured entities (nephrology practices): 3,009
Mortality EHR Dataset	Section 2b.01 Validity Testing	Dates of data: January 1, 2013, through December 31, 2019; representing performance years 2014-2018 (2013 used for patient history, and 2019 for outcome runout) Number of patients in the dataset: 2,860 Number of patient visits in the dataset: 5,658
Master Beneficiary Summary File (MBSF)	Section 2b.30: Risk adjustment/Stratification for Outcome or Resource Use Measures	Dates of Data: July 2016 – June 2019 We used dual-eligible status (for Medicare and Medicaid) derived from the MBSF to study the association between the measure outcome and dual-eligible status.
The American Community Survey (ACS)	Section 2b.30: Risk adjustment/Stratification for Outcome or Resource Use Measures	Dates of Data: 2013-2017 We used the AHRQ SES index score derived from the American Community Survey (2013-2017) to study the association between the outcome and social risk factors. The AHRQ SES index score is based on patient 9-digit zip code level of residence and incorporates 7 census variables found in the American Community Survey.
US Department of Agriculture Economic Research Service: 2013 Rural-Urban Continuum Codes	Section 2b.30: Risk adjustment/Stratification for Outcome or Resource Use Measures	Dates of Data: Collected in 2010 census and 2006-2010 American Community Survey; used for urban/rural social risk factor analysis

Table 2a.07:1. Dataset Descriptions

Total Patients	Number of Patients	Percentage of Patients
Age in the pre-measure year (2017)	*	*

Total Patients	Number of Patients	Percentage of Patients
Mean (SD)	70.14	13.8
Minimum, Maximum	18	109
P1, P99	31	94
Q1, Q3	63	80
Q2 (IQR)	72	17
Gender	*	*
Male	391,058	51.6
Female	367,104	48.4
Race	*	*
Non-Black	588,426	77.6
Black	169,736	22.4
Dual in 2018	*	*
No	565,854	74.6
Yes	192,302	25.4

Table 2a.07:2. Patient Characteristics, Mortality Development Dataset (N=758,162)

*Intentionally left blank

Data for the EHR dataset was derived from a single health system and included deidentified, retrospective demographics (age, sex, gender, race), creatinine and eGFR values, and claims history (comorbidities) for all patients with any outpatient visit from 2013-2019 with Stage 4 or 5 CKD diagnosis or eGFR lab value under 30. Minor data cleaning was applied, including: encounters on the same day were combined as one; patients who only had one encounter were removed; and for encounters where patients only had creatinine, eGFR was calculated using the CKD-EPI 2009 equation¹.

Reference:

1. Equation can be accessed at: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2763564/>

[Response Ends]

2a.08. List the social risk factors that were available and analyzed.

For example, patient-reported data (e.g., income, education, language), proxy variables when social risk data are not collected from each patient (e.g. census tract), or patient community characteristics (e.g. percent vacant housing, crime rate) which do not have to be a proxy for patient-level data.

[Response Begins]

We selected SES variables to analyze after reviewing the literature, developing our conceptual model, and examining available national data sources. The causal pathways for SES variable selection are described below in Section 2b.23. The SES variables used for analysis were:

- **Dual-eligible status:** Dual-eligible status (i.e., enrolled in both Medicare and Medicaid) patient-level data is obtained from the CMS Master Beneficiary Summary File (MBSF).
 - Following guidance from ASPE^{1,2}, NQF³, and a body of literature demonstrating differential health care and health outcomes among dual-eligible patients, we identified dual eligibility as a key variable. We recognize that Medicare-Medicaid dual eligibility has limitations as a proxy for

patients' income or assets because it does not provide a range of results and is only a dichotomous outcome.

- **AHRQ-validated SES index score** (summarizing the information from the following seven variables): percentage of people in the labor force who are unemployed, percentage of people living below poverty level, median household income, median value of owner-occupied dwellings, percentage of people ≥25 years of age with less than a 12th grade education, percentage of people ≥25 years of age completing ≥4 years of college, and percentage of households that average ≥1 people per room.
 - We selected the AHRQ SES index score because it is a well-validated variable that describes the average SES of people living in defined geographic areas⁴. Its value as a proxy for patient-level information is dependent on having the most granular-level data with respect to communities that patients live in. We used the percentage of patients with an AHRQ SES index score equal to or below 43 to define the lowest quartile of the AHRQ SES Index.
- **Race (Black).** Data source: Medicare enrollment database.
 - We used the Medicare enrollment database to identify the patient-level race variable (Black) that we used in these analyses. The Black variable has been shown to be reliable for use in this dataset⁵.
- **Urbanicity.** Data Source: US Department of Agriculture Economic Research Service: 2013 Rural-Urban Continuum Codes
 - The Rural-Urban Continuum Codes assign each county in the US to one of nine subgroups. They distinguish metropolitan counties by the population size of their metro area, and nonmetropolitan counties by degree of urbanization and adjacency to a metro area. The most recent Rural-Urban Continuum Codes are based on data from the 2010 decennial census and the 2006-10 American Community Survey⁶.

References:

1. Department of Health and Human Services, Office of the Assistant Secretary of Planning and Evaluation (ASPE). Report to Congress: Social Risk factors and Performance Under Medicare's Value-based Payment Programs. 2016; <https://aspe.hhs.gov/pdf-report/report-congress-social-risk-factors-and-performance-under-medicare-value-based-purchasing-programs>. Accessed November 10, 2019.
2. Department of Health and Human Services, Office of the Assistant Secretary of Planning and Evaluation (ASPE). Second Report to Congress: Social Risk Factors and Performance in Medicare's Value-based Purchasing Programs. 2020; <https://aspe.hhs.gov/reports/second-report-congress-social-risk-medicare-value-based-purchasing-programs> Accessed July 2, 2020.
3. National Quality Forum. Driving Measurable Health Improvements Together Developing and Testing Risk Adjustment Models for Social and Functional Status-Related Risk within Healthcare Performance Measurement August 2021. Accessed at: <https://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=96087>
4. Bonito A, Bann C, Eicheldinger C, Carpenter L. Creation of new race-ethnicity codes and socioeconomic status (SES) indicators for Medicare beneficiaries. Final Report, Sub-Task. 2008;2.
5. Waldo DR. Accuracy and Bias of Race/Ethnicity Codes in the Medicare Enrollment Database. Health Care Financing Review. 2004;26(2)
6. USDA ERS. Rural-Urban Continuum Codes. 2020; <https://www.ers.usda.gov/data-products/rural-urban-continuum-codes.aspx>. Accessed December 29, 2022

[Response Ends]

Note: If accuracy/correctness (validity) of data elements was empirically tested, separate reliability testing of data elements is not required – in 2a.09 check patient or encounter-level data; in 2a.010 enter “see validity testing section of data elements”; and enter “N/A” for 2a.11 and 2a.12.

2a.09. Select the level of reliability testing conducted.

Choose one or both levels.

[Response Begins]

Accountable Entity Level (e.g., signal-to-noise analysis)

[Response Ends]

2a.10. For each level of reliability testing checked above, describe the method of reliability testing and what it tests.

Describe the steps—do not just name a method; what type of error does it test; what statistical analysis was used.

[Response Begins]

We provide the signal-to-noise reliability statistic among all providers and those with 25 or more cases, showing the mean, standard deviation, and median, quartiles, minimum and maximum. We are using 25 as an example minimum case count.

We used the formula for signal-to-noise reliability presented by Adams et al. to calculate individual clinician-level and TIN-level reliability scores¹. To estimate the overall signal and noise, we first calculated the ICC for the Model Participant, j , using the estimates of between-entity variance τ^2 and the formula for intraclass correlation coefficient (ICC) presented by Shrout and Fleiss². Specifically, the signal-to-noise reliability score for Model Participant, j , R_j is calculated as:

$$R_j = \frac{n_j * ICC}{1 + (n_j - 1) * ICC}$$

while

$$ICC = \frac{\tau^2}{\tau^2 + \pi^2/6\gamma^2}$$

n_j is the number of patients for the nephrologist j , τ^2 is the between agency variance in a Weibull model with lognormal frailty that used to approximate the Cox model with lognormal frailty specified above and represent the signal, and $\frac{\pi^2}{6\gamma^2}$ represents the noise and γ is the shape parameter of the Weibull distribution.

R_j ranges from 0 to 1.0. The higher the score, the higher the reliability. Also, we can see that the reliability of agency measure score will vary depending on the number of patient encounters. Entities with higher volume will tend to have more reliable scores, while those with lower volume will tend to have less reliable scores.

References:

1. Adams J, Mehrota, A, Thoman J, McGlynn, E. (2010). Physician cost profiling – reliability and risk of misclassification. NEJM, 362(11): 1014-1021.
2. Shrout PE, Fleiss JL. Intraclass correlations: uses in assessing rater reliability. Psychological bulletin. 1979;86(2):420. 35. Landis JR, Koch GG. The measurement of observer agreement for categorical data. Biometrics. 1977:159-174.

[Response Ends]

2a.11. For each level of reliability testing checked above, what were the statistical results from reliability testing?

For example, provide the percent agreement and kappa for the critical data elements, or distribution of reliability statistics from a signal-to-noise analysis. For score-level reliability testing, when using a signal-to-noise analysis, more than just one overall statistic should be reported (i.e., to demonstrate variation in reliability across providers). If a particular method yields only one statistic, this should be explained. In addition, reporting of results stratified by sample size is preferred (pg. 18, [NQF Measure Evaluation Criteria](#)).

[Response Begins]

Description	Number of Providers	Mean (SD)	Median (IQR)	Minimum – Maximum
Among All Nephrology Practices	3009	0.623 (0.286)	0.703 (0.430 – 0.867)	0.021 - 0.990
Among Nephrology Practices with at least 25 cases	2403	0.742 (0.173)	0.783 (0.608 – 0.891)	0.344 - 0.990

Table 2a.11. Signal-To-Noise Reliability Statistics Among All Nephrology Practices and Those With At Least 25 Cases, Mortality Development Dataset

[Response Ends]

2a.12. Interpret the results, in terms of how they demonstrate reliability.

(In other words, what do the results mean and what are the norms for the test conducted?)

[Response Begins]

Among all nephrology practices (including those with small case counts), the median signal-to-noise reliability was 0.703, indicating at least half of the providers have a reliability above 0.7. Among those with at least 25 cases, the median signal-to-noise reliability was 0.783. These results demonstrate reliability sufficient for a publicly reported quality measure.

[Response Ends]

2b.01. Select the level of validity testing that was conducted.

[Response Begins]

Empirical validity testing

[Response Ends]

2b.02. For each level of testing checked above, describe the method of validity testing and what it tests.

Describe the steps—do not just name a method; what was tested, e.g., accuracy of data elements compared to authoritative source, relationship to another measure as expected; what statistical analysis was used.

[Response Begins]

Data Element Validity

We validated the accuracy of those patients with Stage 4 or 5 CKD or ESRD (cohort) and the outcome of mortality.

Data element validation was completed for the variable ICD-10 codes N18.4 and N18.5, in the **Mortality EHR Dataset**, further explained Section 2a.07 above. To establish data element validity, we sought to determine the percent agreement comparing patients with at least one outpatient encounter in a calendar year with a diagnosis code for Stage 4 CKD (N18.4) or Stage 5 CKD (N18.5) and the presence of a confirmatory lab value. Since the measure uses claims data to identify patients with Stage 4 and Stage 5 kidney disease, we sought to confirm this with EHR data that measures eGFR, since we heard feedback related to concerns about the ability to accurately identify patients with CKD in claims data. Clinically, CKD is defined by the Estimated Glomerular Filtration Rate (eGFR) kidney function biomarker, which measures how efficiently the kidneys filter waste from blood. Stage 4 CKD is defined as eGFR between 15-29 mL per minute per 1.72 square meters. Stage 5 CKD is defined as eGFR under 15 mL per minute per 1.72 square meters. For encounters that did not have an eGFR in the EHR, a creatinine result was converted into an eGFR¹.

Agreement was defined as: beginning with an outpatient encounter with Stage 4 CKD (ICD-10 code N18.4) or Stage 5 CKD (ICD-10 code N18.5), there was either a) within the same encounter a lab value for eGFR between 15-19; or b) encounter within 180 days prior (or 30-days forward) with a lab value for eGFR between 15-29.

We additionally assessed the validity of using ESRD or ESRD-Dialysis enrollment as the indicator of ESRD status for the cohort by analyzing alignment between ESRD enrollment and documentation of dialysis facility billing codes. Among beneficiaries in 2018, we compared the overlap in ESRD enrollment (which requires completion of CMS Form 2728 documenting evidence of ESRD and start date) to 1) occurrence of dialysis facility claim (Type of Bill [TOB] 72) and 2) TOB 72 claim plus Monthly Capitation Payment (MCP) HCPCS billing codes for ESRD (table sp.16:3).

The outcome of mortality we consider valid as the Medicare Enrollment Database is used for vital statistic information. These data have previously been shown to accurately reflect patient vital status².

Empirical Measure Validity

For empirical validity, we first searched the literature, CMS Measures Inventory Tool, and NQF Quality Positioning System for related measures that are publicly reported, have clinical justification for comparison, and have some overlap in providers with the test dataset for this measure to use for comparative purposes for establishing empiric validity.) As explained further below, we did not find an appropriate or applicable measure.

We then sought another empiric approach to assessing validity. As noted in section 5 (Related and Competing measures), the CKD and ESRD Mortality Measure originated as a re-specification of the validated and endorsed NQF #0369 “Standardized Mortality Ratio for Dialysis Facilities,” a risk-adjusted mortality measure for ESRD patients undergoing dialysis. The CKD and ESRD Mortality Measure includes an expanded cohort of patients (specifically including patients with stage 4 and 5 CKD) in the setting of nephrology practices (rather than dialysis centers) and therefore a modified risk model, but otherwise bears key conceptual and methodological similarities to NQF#0369 (which was most recently re-endorsed in the 2020 Spring cycle). Therefore, to validate the re-specification we computed subgroup risk-decile plots comparing observed to expected outcomes among ESRD patients (similar to the NQF#0369 cohort) and among CKD stage 4 and 5 patients (the expanded group) to assess a) if observed and predicted mortality is less overall among CKD patients than ESRD as would be expected; b) that risk can be differentiated well among both groups in the current measure; and c) that the model performs well in both the “original” ESRD cohort as well as the clinically distinct CKD cohort when all are included.

References:

1. Equation can be accessed at: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2763564/>
2. Fleming C., Fisher ES, Chang CH, Bubolz D, Malenda J. Studying outcomes and hospital utilization in the elderly: The advantages of a merged data base for Medicare and Veterans Affairs Hospitals. Medical Care. 1992; 30(5): 377-91.

[Response Ends]

2b.03. Provide the statistical results from validity testing.

Examples may include correlations or t-test results.

[Response Begins]

Data Element Validity

The observed match rate was high among non-missing eGFRs in both datasets; a majority of patients with a Stage 4 or 5 CKD diagnosis have documented lab values supporting that diagnosis:

- The match rate for patient-visits with a Stage 4 CKD diagnosis and a confirmatory eGFR laboratory value was 88.1%.
- The match rate for patient-visits with a Stage 5 CKD diagnosis and a confirmatory eGFR laboratory value was 90.1%.

In 2018, we found 367,637 total patients enrolled in Medicare ESRD. We found 312,324 total patients with TOB72 claims, of whom 310,264 were also enrolled in ESRD. Examining the overlap shows that 57,373 patients (15.6%) with ESRD enrollment did not have TOB72, while only 2,060 (0.67%) of patients with TOB72 claims were not enrolled in ESRD.

Among patients with TOB72 claims we found that 302,740 also had MCP ESRD service codes, of whom 301,298 were also ESRD-enrolled. Examining the overlap shows that 66,339 patients (18.0%) with ESRD enrollment did not have TOB72 ESRD claims, while only 1,442 (0.48%) of patients with TOB72 ESRD claims were not enrolled in ESRD.

Empirical Validity

We initially identified three electronic clinical quality measures (eQMs) from the Merit-based Incentive Payment System (MIPS) with some potential relationship to renal care: “Diabetes: Medical Attention for Nephropathy” (a process measure), “Diabetes: Hemoglobin A1c (HbA1c) Poor Control (>9%),” and “Controlling High Blood Pressure” (both intermediate outcome measures). Our search did not identify any other potentially comparable measures that would be feasible to test. Upon further consideration of these measures, we determined they were not suitable validity comparators for the CKD and ESRD Mortality Measure for the conceptual and empirical limitations discussed below.

Conceptually, while they are important aspects of care for patients, **the Attention for Nephropathy and A1c Poor Control measures are not necessarily clinically relevant measures of a nephrologist’s quality of care.** Both measures reflect a cohort of diabetic patients; while many CKD and ESRD patients may have comorbid diabetes, treatment of diabetes usually falls outside the direct scope of their nephrology specialty care. Attention for Nephropathy is a process measure that primarily evaluates screening for renal disease among Diabetic patients, which is less relevant in the care of late-stage CKD patients. A1c Poor Control is largely addressed by physicians other than nephrologists (such as general practitioners and endocrinologists); nephrologists are typically not directly involved in A1c management for their patients. Hypertension control may be more commonly addressed directly by nephrologists but still present critical empirical concerns as documented below.

The goal for empiric validation is to find a clinically relevant quality measure that assesses the same providers for a similar signal of care quality. Unfortunately, empirically, the utility of analyses for MIPS measures are limited as scores for both the comparison measures and the CKD and ESRD Mortality Measure are aggregated at the taxpayer identification number (TIN) level which describes groups of clinicians. **While the CKD and ESRD Mortality Measure only includes nephrology care within a TIN, the comparison measures include data across all individual providers within the TIN, of which nephrology may be only a small part.** Of 3,009 TINs receiving a CKD and ESRD Mortality Measure score in the 2018 testing dataset, only 437 (15%) also reported scores in A1c poor control and Controlling High Blood Pressure for MIPS in the same period; even fewer (344, 11%) reported Nephropathy Attention scores. Among the 437 with Controlling High Blood Pressure scores, the median TIN included 23 distinct specialties with only 2.8% of all claims falling under nephrology; this means that for most TINs the MIPS measure score is driven by non-nephrology care and thus would not provide a helpful comparison for empiric validity. By comparison among the 2,252 TINs for which we could calculate the CKD and ESRD Mortality Measure scores but for which we had no blood pressure control scores, the median TIN included just 2 unique specialties with 81.0% of claims falling under

nephrology. These TINs are more relevant for nephrology care as a large majority of claims are for nephrologists, but unfortunately do not have corresponding MIPS scores for comparison. We found similar issues for the A1c Poor Control measure.

Comparison measure:	N (%)	Median nephrology claim % (IQR)*	Median # specialties (IQR)
With BP control	437 (14.5%)	2.8 (0.7, 68.9)	23 (4, 40)
Without BP control	2,252 (85.5%)	81.0 (2.7, 100)	2 (1, 14)
With A1c control	437 (14.5%)	2.4 (0.7, 54.5)	25 (4, 41)
Without A1c control	2,252 (85.5%)	82.6 (2.7, 100)	2 (1, 13)

Table 2b.03. Comparison of TINs with CKD and ESRD Mortality Measure scores, with vs. without blood pressure (BP) measure scores and with vs. without A1c control measure scores

*Nephrology claim percent is defined as the number of claim lines with “Nephrology” as the listed specialty out of the total number of claim lines for the TIN.

In summary we did not find publicly available scores that were both clinical and conceptually relevant to nephrology providers' care and available at the clinician group level for groups with a substantial presence of nephrologists included in our measure.

To provide some assurance that the measure performs well in reference to an independent standard, we have tested the calibration of the patient-level risk model for the patients newly added to the previously NQF-endorsed dialysis facility mortality measure. Using a patient-level logistic regression model to obtain predicted death risk for subgroups of ESRD patients and for stage 4/5 CKD patients (using the same model coefficients from the entire combined cohort), we compared to the observed mortality across deciles of the predicted values as shown in Figures 2b.03:1-2. The overall mortality risk was 11.0% among the ESRD subgroup and 6.0% among the Stage 4/5 CKD subgroup. The bottom-decile predicted risk was 2.7% for ESRD and 1.7% for CKD; the top-decile predicted risk was 30.1% for ESRD and 17.5% for CKD.

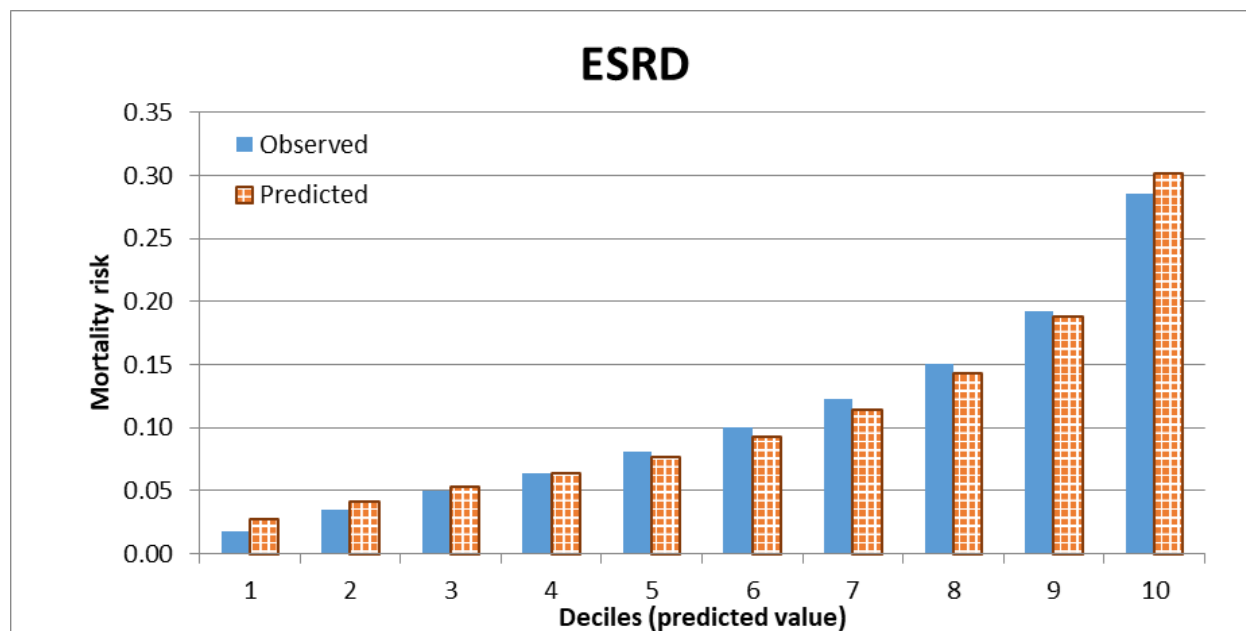


Figure 2b.03:1. Observed vs. predicted risk of death by deciles of predicted value, ESRD subgroup (n=297,787)

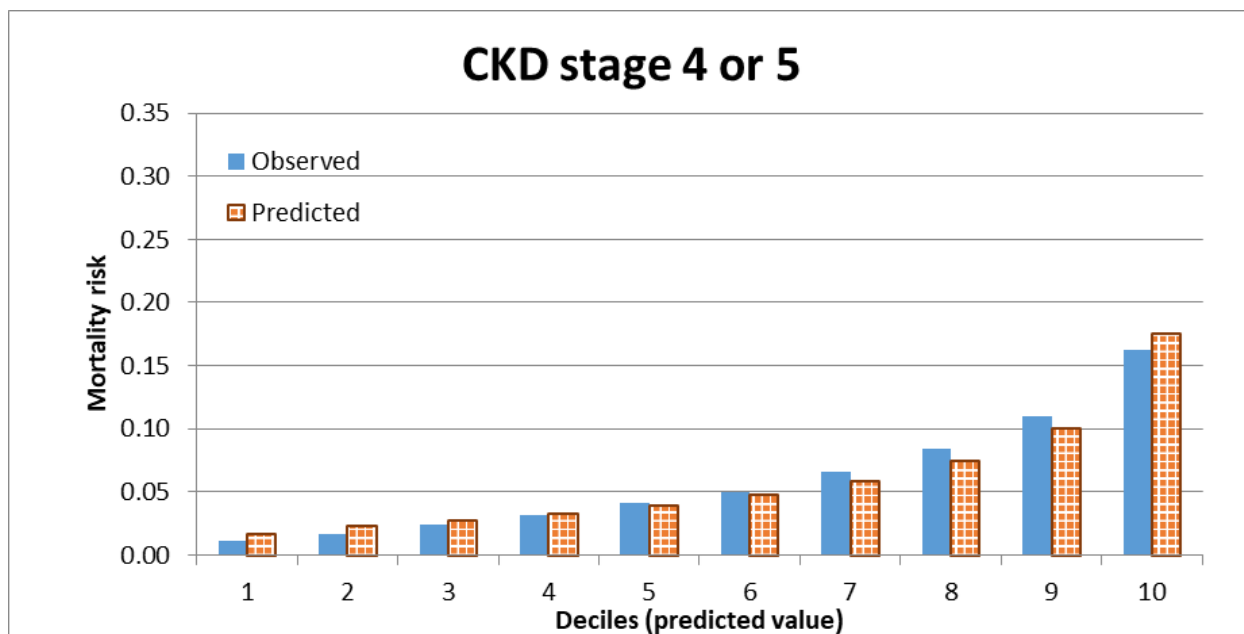


Figure 2b.03:2. Observed vs. predicted risk of death by deciles of predicted value, CKD subgroup (n=460,375)

[Response Ends]

2b.04. Provide your interpretation of the results in terms of demonstrating validity. (i.e., what do the results mean and what are the norms for the test conducted?)

[Response Begins]

Data Element Validity

The match rates were high for both Stage 4 CKD and Stage 5 CKD, indicating that the claims data is valid to be used in cohort determination for the measure.

Because ESRD enrollment requires documented evidence of ESRD (submitted in Form 2728), and ESRD enrollment is required in order to bill Medicare for regular dialysis treatment, the face validity of ESRD enrollment as an indicator of ESRD status (with a clear start date) is high. This is corroborated by comparison to two other potential claims-based means of identifying ESRD status; ESRD enrollment includes nearly all beneficiaries with ESRD billing claims. We conclude that ESRD enrollment is the most comprehensive and reliable record for ESRD/dialysis status available using administrative data sources.

Empirical Validity

The validity of the CKD and ESRD Mortality Measure is supported by the following:

- Mortality, in patients with CKD and ESRD, has inherent face validity as a quality measure; we have also provided evidence of data element validity to demonstrate capture of patients with CKD and ESRD in claims.
- The CKD and ESRD Mortality Measure is a re-specification of the Standardized Mortality Ratio for Dialysis Facilities (NQF#0369) measure that has been deemed valid and is currently NQF-endorsed.
- Our internal validity results show that as expected, the subgroup of patients with CKD have overall lower mortality compared to patients with ESRD.
- Our model validation results show good calibration for both CKD and ESRD patients, which supports the expanded cohort of the CKD and ESRD Mortality Measure as a respecification of NQF#0369.

The subgroup analysis revealed a few important findings about the measure methodology that support its use in an expanded context from the original NQF#0369 measure. First, the risk adjustment model has a clear predictive ability to differentiate outcomes in clinically distinct subgroups, ranging from 2.7%-30.1% from the bottom to top deciles in the ESRD subgroup and 1.7%-17.5% in the CKD subgroup. Second, the overall mortality is much lower among the CKD subgroup (6.0%, compared to 11.0% among the ESRD subgroup), which aligns with the expectation that ESRD patients are generally at higher risk for death. Finally, the predicted and observed deaths align closely within each subgroup, demonstrating that the common risk model is well-calibrated both in the original NQF#0369 cohort of ESRD patients and in the expanded cohort of stage 4/5 CKD patients.

[Response Ends]

2b.05. Describe the method for determining if statistically significant and clinically/practically meaningful differences in performance measure scores among the measured entities can be identified.

Describe the steps—do not just name a method; what statistical analysis was used? Do not just repeat the information provided in Importance to Measure and Report: Gap in Care/Disparities.

[Response Begins]

Examination of provider-level results include measure scores for all nephrologists and those with at least 25 patients, along with statistics summarizing their distribution. We are using 25 as an example minimum case count, which aligns with CMS publicly reported outcome measures.

[Response Ends]

2b.06. Describe the statistical results from testing the ability to identify statistically significant and/or clinically/practically meaningful differences in performance measure scores across measured entities.

Examples may include number and percentage of entities with scores that were statistically significantly different from mean or some benchmark, different from expected; how was meaningful difference defined.

[Response Begins]

As discussed in Section sp.24 (calculation of measure score), RSMR is a ratio measure with a score of 1 indicating median performance for a given case mix, a score less than 1 indicating lower mortality (better performance) than expected, and a score greater than 1 indicating greater mortality than expected.

Statistics	All Nephrology Practices (N=3,009)	Nephrology Practices with 25 + Patients (N=2,403)
RSMR: Mean (SD)	1.006 (0.108)	1.005 (0.118)
RSMR: Median (IQR)	0.997 (0.944 -1.056)	0.994 (0.928- 1.068)
RSMR: Range (min-max)	0.672 - 1.676	0.672 - 1.676
Volume: Mean (SD)	252.0 (388.7)	313.1 (413.3)
Volume: Median (IQR)	113 (36 - 310)	172 (74 - 391)
Volume: min - max	1 - 4940	25 - 4940

Table 2b.06. Measure Performance Statistics (RSMR) for All Providers and Providers with 25 or More Patients, Mortality Development Dataset

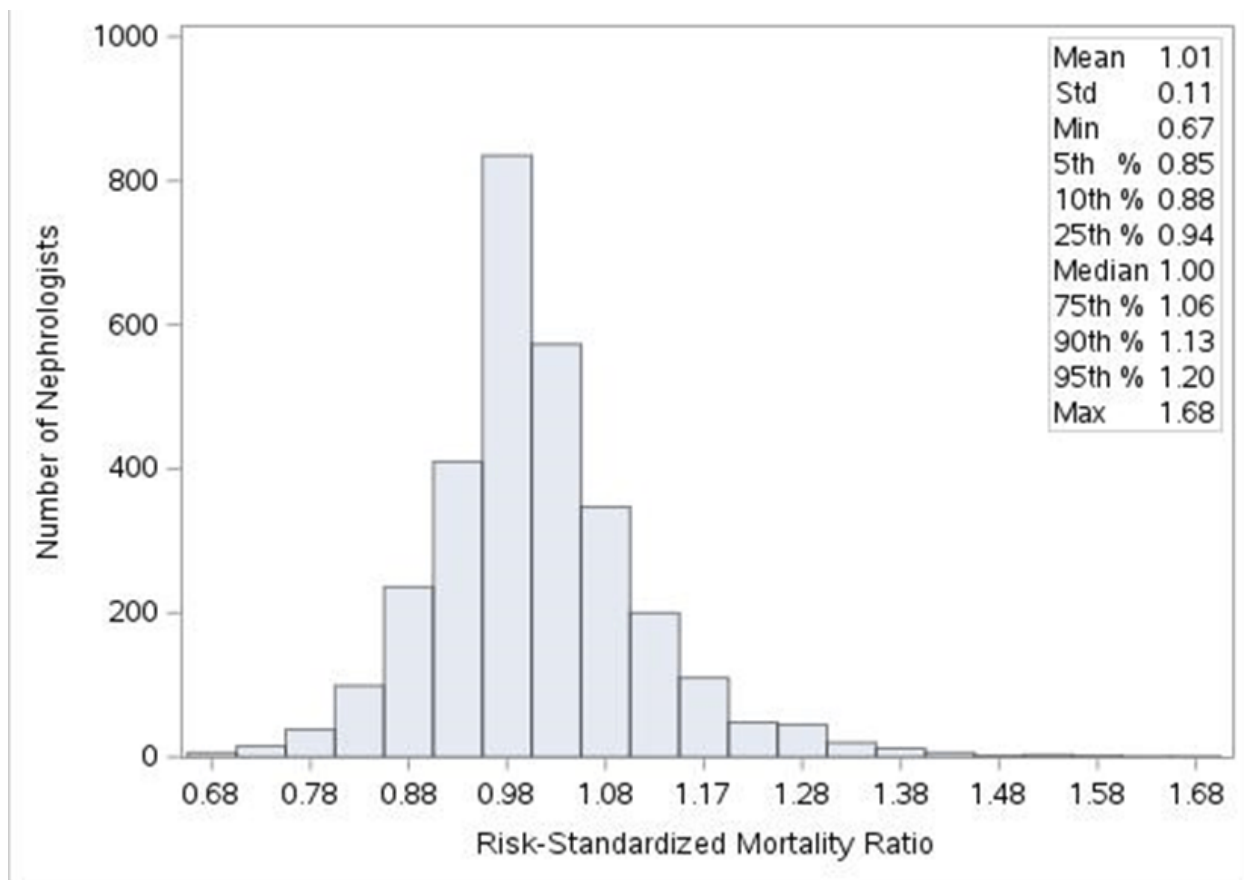


Figure 2b.06:1. Distribution of RSMR, all nephrology practices (n=3009)

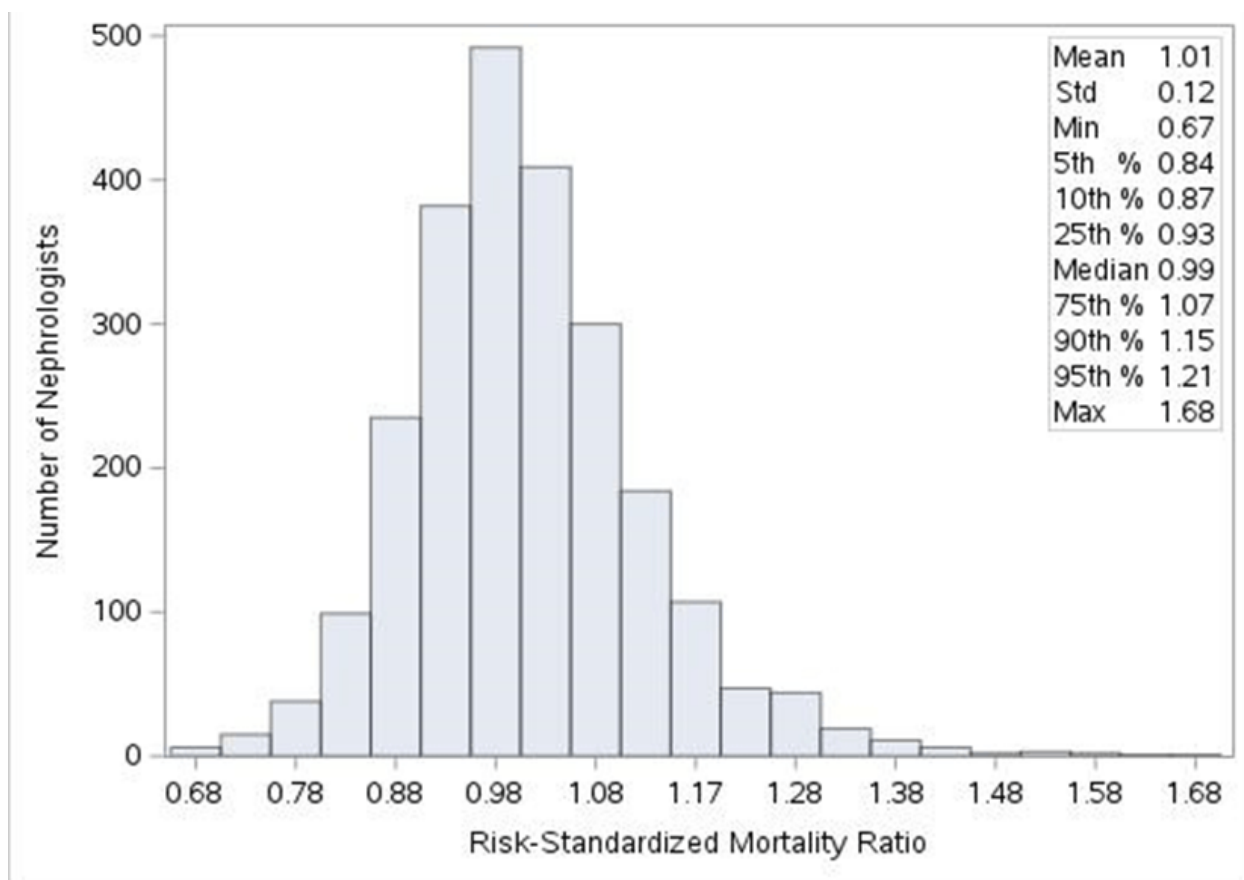


Figure 2b.06:2. Distribution of RSMR, nephrology practices with 25+ patients (n=2403)

[Response Ends]

2b.07. Provide your interpretation of the results in terms of demonstrating the ability to identify statistically significant and/or clinically/practically meaningful differences in performance across measured entities.

In other words, what do the results mean in terms of statistical and meaningful differences?

[Response Begins]

As shown by these distributions of the performance score in Section 2b.06, there was a substantial variation in performance between measure entities after accounting for clinical risk. The range of 0.68 to 1.65 across providers with 25 or more patients (a 2.4-fold increase in mortality hazard between the best and worst-quality nephrologists after accounting for case mix) indicates a large gap that can be explained by a meaningful difference in performance. There is also a fairly broad distribution between the extreme values as illustrated in the histogram. At top practices, there are fewer deaths on a risk-adjusted basis. This shows that not all entities will score the same, and that entities can be distinguished from each other in terms of quality.

[Response Ends]

2b.08. Describe the method of testing conducted to identify the extent and distribution of missing data (or non-response) and demonstrate that performance results are not biased due to systematic missing data (or differences between responders and non-responders). Include how the specified handling of missing data minimizes bias.

Describe the steps—do not just name a method; what statistical analysis was used.

[Response Begins]

The measure used claims-based data for development and testing. All administratively coded diagnoses were available for measurement. Lack of a claim is treated as not having the corresponding diagnosis or procedure.

[Response Ends]

2b.09. Provide the overall frequency of missing data, the distribution of missing data across providers, and the results from testing related to missing data.

For example, provide results of sensitivity analysis of the effect of various rules for missing data/non-response. If no empirical sensitivity analysis was conducted, identify the approaches for handling missing data that were considered and benefits and drawbacks of each).

[Response Begins]

The measure used claims-based data for development and testing. All administratively coded diagnoses were available for measurement.

[Response Ends]

2b.10. Provide your interpretation of the results, in terms of demonstrating that performance results are not biased due to systematic missing data (or differences between responders and non-responders), and how the specified handling of missing data minimizes bias.

In other words, what do the results mean in terms of supporting the selected approach for missing data and what are the norms for the test conducted; if no empirical analysis was conducted, justify the selected approach for missing data.

[Response Begins]

The measure used claims-based data for development and testing. All administratively coded diagnoses were available for measurement. The data on patient deaths were obtained from the Medicare Enrollment Database; these data have previously been shown to accurately reflect patient vital status¹. As discussed in section 2b (Validity testing) the validation of the claims-based methodology against EHR data demonstrates that the measure satisfactorily identifies the appropriate patients.

Reference:

1. Fleming C., Fisher ES, Chang CH, Bubolz D, Malenda J. Studying outcomes and hospital utilization in the elderly: The advantages of a merged data base for Medicare and Veterans Affairs Hospitals. Medical Care. 1992; 30(5): 377-91.

[Response Ends]

Note: This item is directed to measures that are risk-adjusted (with or without social risk factors) OR to measures with more than one set of specifications/instructions (e.g., one set of specifications for how to identify and compute the measure from medical record abstraction and a different set of specifications for claims or eCQMs). It does not apply to measures that use more than one source of data in one set of specifications/instructions (e.g., claims data to identify the denominator and medical record abstraction for the numerator). Comparability is not required when comparing performance scores with and without social risk factors in the risk adjustment model.

However, if comparability is not demonstrated for measures with more than one set of specifications/instructions, the different specifications (e.g., for medical records vs. claims) should be submitted as separate measures.

2b.11. Indicate whether there is more than one set of specifications for this measure.

[Response Begins]

No, there is only one set of specifications for this measure

[Response Ends]

2b.12. Describe the method of testing conducted to compare performance scores for the same entities across the different data sources/specifications.

Describe the steps—do not just name a method. Indicate what statistical analysis was used.

[Response Begins]

[Response Ends]

2b.13. Provide the statistical results from testing comparability of performance scores for the same entities when using different data sources/specifications.

Examples may include correlation, and/or rank order.

[Response Begins]

[Response Ends]

2b.14. Provide your interpretation of the results in terms of the differences in performance measure scores for the same entities across the different data sources/specifications.

In other words, what do the results mean and what are the norms for the test conducted.

[Response Begins]

[Response Ends]

2b.15. Indicate whether the measure uses exclusions.

[Response Begins]

Yes, the measure uses exclusions.

[Response Ends]

2b.16. Describe the method of testing exclusions and what was tested.

Describe the steps—do not just name a method; what was tested, e.g., whether exclusions affect overall performance scores; what statistical analysis was used?

[Response Begins]

All exclusions were determined by careful clinical review and have been made based on clinically relevant decisions. The prevalence of exclusions in the Mortality Development Dataset are reported below.

[Response Ends]

2b.17. Provide the statistical results from testing exclusions.

Include overall number and percentage of individuals excluded, frequency distribution of exclusions across measured entities, and impact on performance measure scores.

[Response Begins]

There were 7,156 patients excluded due to having metastatic cancer within one year prior to their Stage 4 or Stage 5 CKD diagnosis. The final study cohort was 758,162, so this represents 0.94% of the final cohort.

[Response Ends]

2b.18. Provide your interpretation of the results, in terms of demonstrating that exclusions are needed to prevent unfair distortion of performance results.

In other words, the value outweighs the burden of increased data collection and analysis. Note: If patient preference is an exclusion, the measure must be specified so that the effect on the performance score is transparent, e.g., scores with and without exclusion.

[Response Begins]

The measure excludes patients with metastatic and advanced cancers, since the outcome is not a reliable signal of care quality among these patients. Many patients in this population may be too ill for dialysis and have a high risk of mortality regardless of the quality of their CKD or ESRD care; thus, we find it inappropriate to attribute outcomes for these patients to their nephrologists' quality of care. These exclusions are minimal in practice but are important for measure validity given the outcome.

[Response Ends]

2b.19. Check all methods used to address risk factors.

[Response Begins]

Statistical risk model with risk factors (specify number of risk factors)

[Statistical risk model with risk factors (specify number of risk factors) Please Explain]

71

[Response Ends]

2b.20. If using statistical risk models, provide detailed risk model specifications, including the risk model method, risk factors, risk factor data sources, coefficients, equations, codes with descriptors, and definitions.

[Response Begins]

The goal of risk adjustment is to account for differences among nephrology practices in patient demographic and clinical characteristics. The measure incorporates risk adjustment to account for factors that are associated with the outcome, vary across providers, and are unrelated to quality of care so that measure scores reflect differences in care quality. Accounting for case-mix differences is important because it recognizes that some providers care for older, sicker patients who have higher mortality rates. Through the risk-adjustment modeling, a higher expected outcome rate is set for providers who care for patients with certain risk factors. We identified potential candidate risk factors using a focused literature search, clinical experts' input, and empirical analysis. We used logistic regression with a binary outcome to select risk variables for final models.

We considered age and medical history (comorbidities/frailty) as candidate variables.

- Comorbidities for inclusion in risk adjustment were identified through inpatient and outpatient administrative claims during the twelve months prior to entering the cohort.
- We used publicly available CMS condition categories (CMS-CCs) to group ICD-10 diagnosis codes into CMS-CCs, and selected comorbidities based on clinical relevance and statistical significance.
- We aligned with other CMS outcome measures by using the Yale-Modified FY20 v24 CC Map that contains 197 CMS-CCs.

Next,

- We examined all condition categories (CMS-CCs).
- Examined frequencies and bivariate associations with outcome (including odds ratios) of all CMS-CCs.
- CMS-CCs that were not statistically significant were removed, unless deemed clinically relevant to the outcome by expert nephrologists (ex: diabetic-related CC). Statical significance was defined by having a p-value less than 0.05 (14 CC removed).
- CMS-CCs with low frequency (<1% of cohort), were grouped into one variable, except for CC1 HIV/AIDS and CC51 Dementia with Complications (48 CC grouped).
- CC132 Kidney Transplant Status was split into two: CC132Z ICD-10-CM codes beginning with 'Z' (codes indicating general aftercare or status); and CC132T ICD-10-CM codes beginning with 'T' (codes indicative of a kidney failure or complication).

This process resulted in 117 candidate risk variables

Final Risk Variable Selection

We selected the final set of risk variables using bootstrap methods using logistic regression from the candidate variables:

- 500 random samples were generated with replacement.
- For each of the 500 samples, a logistic regression model (binary outcome) was selected by using backward selection approach.
- All variables significant at $p < 0.0001$ were retained in each final bootstrap risk model. For each variable, we note its % retained in the 500 bootstrap models.
- We then selected all variables that were retained in the model which are above 70% threshold (cut-off). The threshold for cut-off was based on clinical and statistical evaluation to have a clinically meaningful, statistically robust, and parsimonious risk model.
- Low frequency CC variable was removed. This group was very heterogeneous; removing aligns with many other measures that excluded prior to bootstrap results.
- Six additional CCs were included that were below the 70% cut-off, for face validity, per our expert nephrologists (Diabetes with Chronic Complications [CC18], Diabetes without Complication [CC19], Unstable Angina and Other Acute Ischemic Heart Disease [CC87], Dialysis Status [CC134], Acute Renal Failure [CC135], Unspecified Renal Failure [CC 140])
- Proteinuria identified by ICD-10 code (R80.9) was included as a risk variable. We included the Proteinuria code as a separate variable based on input from nephrologists regarding its clinical relevance and importance for face validity.
- We then examined the impact of several variables interacting with others, and the impact of including ESRD coverage as a risk variable.
- ESRD coverage was added as a risk variable.

There are 71 final risk variables. We evaluated the performance of the model in the Cox model with the selected risk factors.

[Response Ends]

2b.21. If an outcome or resource use measure is not risk-adjusted or stratified, provide rationale and analyses to demonstrate that controlling for differences in patient characteristics (i.e., case mix) is not needed to achieve fair comparisons across measured entities.

[Response Begins]

Not applicable; the measure is risk-adjusted.

[Response Ends]

2b.22. Select all applicable resources and methods used to develop the conceptual model of how social risk impacts this outcome.

[Response Begins]

Published literature

Internal data analysis

[Response Ends]

2b.23. Describe the conceptual and statistical methods and criteria used to test and select patient-level risk factors (e.g., clinical factors, social risk factors) used in the statistical risk model or for stratification by risk.

Please be sure to address the following: potential factors identified in the literature and/or expert panel; regression analysis; statistical significance of $p < 0.10$ or other statistical tests; correlation of x or higher. Patient factors should be present at the start of care, if applicable. Also discuss any “ordering” of risk factor inclusion; note whether social risk factors are added after all clinical factors. Discuss any considerations regarding data sources (e.g., availability, specificity).

[Response Begins]

Methods for identifying clinical risk factors are detailed in Section 2b.20.

A patient’s risk of mortality is likely influenced by social risk factors (SRFs). Kidney care providers have the ability to address these SRFs and mitigate the impact on mortality. We consider whether to adjust for SRF using a comprehensive approach that evaluates the following:

1. Conceptual influence of SRFs on measure outcome (and provider role)
2. Feasibility of utilizing meaningful SRFs in available data
3. Empiric testing of SRFs for inclusion in the measure risk models

Updated NQF guidance emphasizes that developers should share the conceptual model that was used to guide empiric testing and decisions around inclusion of social risk factors within the measure’s risk model [1]. Conceptual models should illustrate the pathway between the social and/or functional status-related risk factors, patient clinical factors, quality of care, and the measured healthcare outcome. Our conceptual model for CKD and ESRD mortality is shown in Figure 2b.23. For this CKD and ESRD Mortality Measure, the conceptual relationship, or potential causal pathways by which these possible social risk factors influence the risk of mortality are varied and complex. Some social risk factors may, for instance, influence the patient’s ability to manage self-care such as following dietary recommendations. However, the best quality care should reduce mortality for all patient groups, especially if tailored to a particular patient’s situation and preferences. Therefore, the conceptual rationale for risk-adjustment is limited.

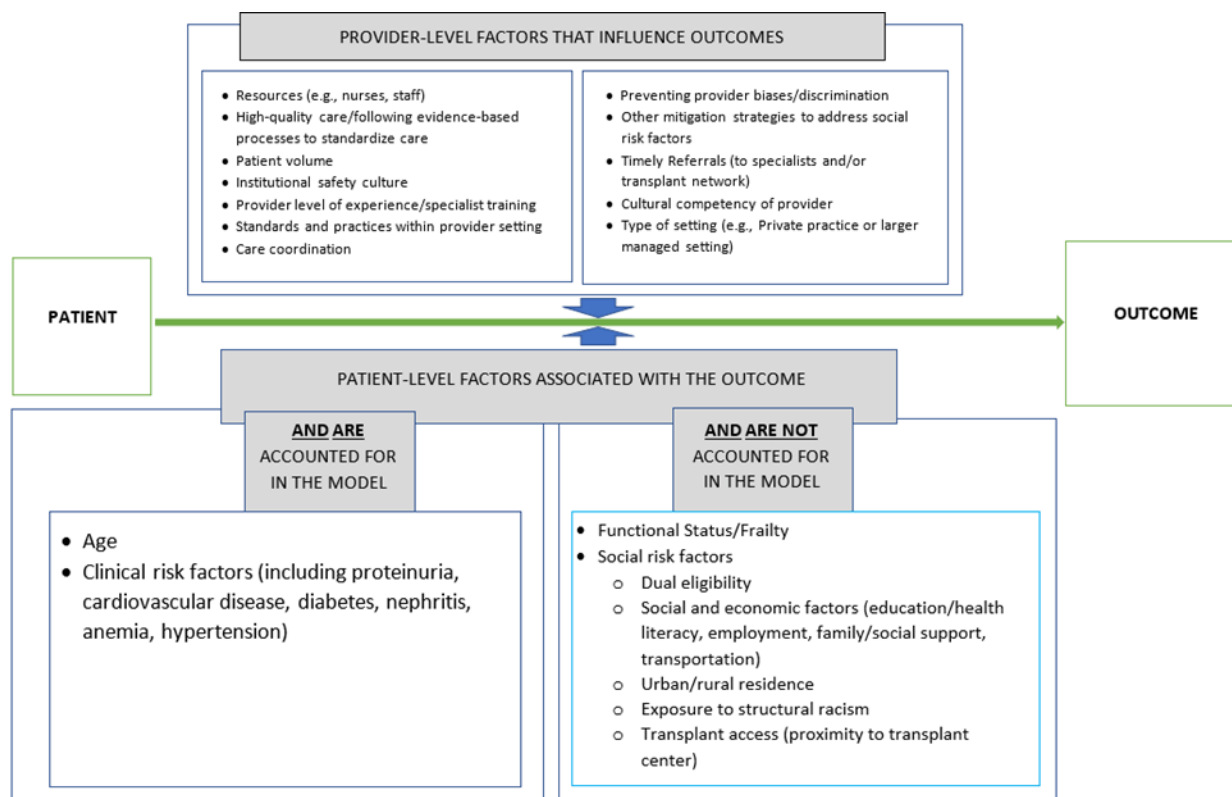


Figure 2b.23. Conceptual Model

There are widely recognized ethnic and racial disparities in both the incidence of kidney disease and in many outcomes (including time to diagnosis, time to progression, and transplant-related outcomes), however, a review of the literature shows mixed evidence regarding the relationship between race/ethnicity and the outcome of mortality in patients with stage 4 or 5 CKD. For example, a scoping review of the literature published in 2020 found that most studies (18 of 27) showed no difference in mortality rates in patients from different race/ethnicities who had pre-dialysis kidney disease; five of the 27 studies showed higher rates in Black patients, four showed lower rates in Black patients [2]. The most recent (2022) United States Renal Data systems report showed that adjusted mortality rates in white patients with stage 4 or 5 CKD were higher compared with Black patients [3].

There is some evidence, however, supporting a relationship between socioeconomic status and the outcome of mortality in patients with CKD. For example, one study found that patients experiencing homelessness had a higher adjusted mortality rates compared with patients with stable housing [4]; another study found that patients in higher income neighborhoods had lower mortality rates [5]. In another study, socioeconomic status was found to be independently associated with a higher risk of mortality, in both Black and white patients [6].

We have also included functional status/frailty, Medicaid dual-eligibility, urban/rural residence, and proximity/access to transplant centers in our conceptual model, based in part on the literature (where available) as well based on expert consideration. Dual-eligible status is an indicator of low income and a proxy indicator of various socioeconomic factors that may affect patients' options for CKD treatment. Similarly, urban vs. rural residence may affect the proximity to effective CKD care that may be associated with worse outcomes. Patient's functional status and frailty may also play a role in patients' ability to access care for CKD. Finally, patients' ability to receive kidney transplants (a censoring event for this measure) may be affected by their proximity to transplant centers.

To define a list of SRF indicators that would be feasible to test, we first compiled an initial list of SRFs to consider, using the National Academies of Sciences, Engineering, and Medicine (NASEM) report framework, which categorized social risk factors into four domains:

- Socioeconomic position;

- Race, ethnicity (not as biological factors but as proxy for the social risk factor of exposure to systemic racism), and cultural factors;
- Social relationships; and
- Residential and community context

Second, we identified candidate SRFs for analyses, based on:

- Internal hypotheses regarding the relationships between the SRF and mortality for patients with CKD or ESRD;
- Potential / perceived ability of a kidney care provider to mitigate the SRF; and
- Data availability and feasibility, including level of analysis (availability of patient-level or area-level data).

Among candidate SRFs, we identified the corresponding variable from different data sources and linked them to the test dataset based on the related patient information. The candidate social risk variables considered are listed in Section 2a.08, and below in Table 2b.23. Candidate social risk factors from Medicare FFS claims including Medicaid dual-eligibility and Black race, while Agency for Healthcare Research and Quality (AHRQ) socioeconomic status (SES) index, and urban residence could be linked to patients at the ZIP code and county level respectively. We did not identify a suitable and feasible SRF indicator for functional status/frailty or proximity to transplant centers.

Variable	Description	Data level
Dual-eligible	Dual-eligible for Medicare and Medicaid vs. Medicare-only (reference variable)	Patient
AHRQ SES index	Lowest AHRQ quartile for socioeconomic status indicator (higher score = less social risk) vs. other quartiles (reference variable)	Zip code
Race	Black race variable vs. non-Black race variables (reference variable). Note: Medicare administrative claims data are not a reliable source for accurate race information except for Black race, as noted in the literature. Included here as above to explore general impact using available data.	Patient
Urban resident	Residence in metro area county vs. non-metro county (suburban and rural are considered non-urban) (reference variable)	County

Table 2b.23. Candidate Social Risk Factors

Methods for testing each social risk factor included examining the prevalence and distribution of SRFs, bivariate (unadjusted) associations of SRFs with mortality, risk adjusted associations, and risk model performance when incorporating SRFs, including impact on provider performance scores.

Some patient level factors potentially associated with the outcome but not accounted for the in the model were ultimately not tested for the following reasons. Social and economic factors such as health literacy and transportation do not have enough available data to accurately capture and adjust for their influence. Transplant is considered a censoring event in our measure and does not count as an adverse event rather it is encouraged as a positive event. There are a large number of external and individual factors that can impact transplant access other than geographic location, including patient willingness to receive care and the availability of kidneys for transplant once on the waitlist. Frailty is potentially associated with mortality but is not accounted for in this measure due to a lack of suitable data; however, aspects of frailty will be captured in the clinical factors that are adjusted for in the model.

References:

1. NQF 2022: National Quality Forum (NQF). Developing and Testing Risk Adjustment Models for Social and Functional Status-Related Risk within Healthcare Performance Measurement.; 2022. <https://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=96087>. Accessed December 24, 2022.
2. Hounkpatin HO, Fraser SDS, Honney R, Dreyer G, Brettle A, Roderick PJ. Ethnic minority disparities in progression and mortality of pre-dialysis chronic kidney disease: a systematic scoping review. *BMC Nephrol*. 2020 Jun 9;21(1):217.
3. United States Renal Data System: USRDS annual data report: Epidemiology of kidney disease in the United States (2022). Available at <https://usrds-adr.niddk.nih.gov/2022/supplements-covid-19-disparities/14-racial-and-ethnic-disparities>; accessed January 5, 2023.
4. Hall YN, Choi AI, Himmelfarb J, Chertow GM, Bindman AB. Homelessness and CKD: a cohort study. *Clinical journal of the American Society of Nephrology : CJASN*. 2012 Jul;7(7):1094–1102.
5. Garg PP, Diener-West M, Powe NR. Income-based disparities in outcomes for patients with chronic kidney disease. *Seminars in nephrology*. 2001 Jul;21(4):377–385.
6. Fedewa SA, McClellan WM, Judd S, Gutiérrez OM, Crews DC. The association between race and income on risk of mortality in patients with moderate chronic kidney disease. *BMC Nephrol*. 2014 Aug 23;15:136.

[Response Ends]

2b.24. Detail the statistical results of the analyses used to test and select risk factors for inclusion in or exclusion from the risk model/stratification.

[Response Begins]

Description (CC#)	Percentage	Estimate (Standard Error)	Hazard Ratio (95% confidence interval)
Age (mean (standard deviation))	70.14 (13.76)	0.02 (0.00)	1.018 (1.017-1.019)
Proteinuria: DX Code of R80.9	16.05	-0.09 (0.01)	0.911 (0.887-0.936)
ESRD-Dialysis Enrollment	39.28	-0.15 (0.02)	0.856 (0.823-0.892)
Metastatic Cancer and Acute Leukemia (CC 8)	1.62	0.48 (0.03)	1.612 (1.523-1.706)
Lung and Other Severe Cancers (CC 9)	3.61	0.21 (0.02)	1.231 (1.184-1.28)
Lymphoma and Other Cancers (CC 10)	2.72	0.15 (0.02)	1.163 (1.110-1.218)
Colorectal, Bladder, and Other Cancers (CC 11)	6.09	0.01 (0.02)	1.012 (0.977-1.048)
Other Digestive and Urinary Neoplasms (CC 14)	10.04	-0.10 (0.01)	0.902 (0.878-0.927)
Diabetes with Acute Complications (CC 17)	2.62	0.12 (0.02)	1.129 (1.084-1.177)
Diabetes with Chronic Complications (CC 18)	59.76	0.11 (0.01)	1.118 (1.089-1.148)
Diabetes without Complication (CC 19)	58.15	0.04 (0.01)	1.039 (1.013-1.066)
Protein-Calorie Malnutrition (CC 21)	9.39	0.11 (0.01)	1.114 (1.088-1.14)
Morbid Obesity (CC 22)	14.12	-0.10 (0.01)	0.902 (0.881-0.924)
Disorders of Fluid/Electrolyte/Acid-Base Balance (CC 24)	57.29	0.10 (0.01)	1.104 (1.081-1.127)

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Description (CC#)	Percentage	Estimate (Standard Error)	Hazard Ratio (95% confidence interval)
Disorders of Lipoid Metabolism (CC 25)	80.52	-0.17 (0.01)	0.844 (0.824-0.863)
End-Stage Liver Disease (CC 27)	1.94	0.25 (0.03)	1.286 (1.224-1.351)
Cirrhosis of Liver (CC 28)	3.27	0.31 (0.02)	1.365 (1.312-1.421)
Disorders of the Vertebrae and Spinal Discs (CC 41)	23.67	-0.09 (0.01)	0.918 (0.900-0.936)
Osteoporosis and Other Bone/Cartilage Disorders (CC 43)	25.81	-0.09 (0.01)	0.910 (0.893-0.927)
Severe Hematological Disorders (CC 46)	2.71	0.17 (0.02)	1.181 (1.134-1.23)
Coagulation Defects and Other Specified Hematological Disorders (CC 48)	17.63	0.09 (0.01)	1.09 (1.068-1.111)
Iron Deficiency and Other/Unspecified Anemias and Blood Disease (CC 49)	81.54	0.12 (0.02)	1.127 (1.094-1.162)
Delirium and Encephalopathy (CC 50)	11.39	0.10 (0.01)	1.104 (1.079-1.13)
Dementia With Complications (CC 51)	2.28	0.13 (0.02)	1.140 (1.090-1.193)
Dementia Without Complication (CC 52)	10.96	0.11 (0.01)	1.121 (1.094-1.149)
Depression (CC 61)	18.38	0.00 (0.01)	0.999 (0.978-1.019)
Parkinson's and Huntington's Diseases (CC 78)	1.88	0.02 (0.03)	1.016 (0.963-1.072)
Seizure Disorders and Convulsions (CC 79)	5.71	0.05 (0.02)	1.056 (1.025-1.088)
Cardio-Respiratory Failure and Shock (CC 84)	18.28	0.18 (0.01)	1.193 (1.167-1.22)
Congestive Heart Failure (CC 85)	49.53	0.36 (0.01)	1.427 (1.396-1.458)
Acute Myocardial Infarction (CC 86)	8.88	0.18 (0.01)	1.202 (1.174-1.231)
Unstable Angina and Other Acute Ischemic Heart Disease (CC 87)	7.69	0.03 (0.01)	1.026 (1-1.053)
Coronary Atherosclerosis/Other Chronic Ischemic Heart Disease (CC 89)	48.43	0.15 (0.01)	1.160 (1.137-1.183)
Valvular and Rheumatic Heart Disease (CC 91)	32.03	0.12 (0.01)	1.131 (1.111-1.152)
Hypertension (CC 95)	91.62	-0.22 (0.02)	0.805 (0.779-0.832)
Specified Heart Arrhythmias (CC 96)	33.16	0.22 (0.01)	1.252 (1.229-1.275)
Atherosclerosis of the Extremities with Ulceration or Gangrene (CC 106)	4.12	0.07 (0.02)	1.073 (1.036-1.111)
Vascular Disease (CC 108)	43.41	0.08 (0.01)	1.086 (1.066-1.106)
Other Circulatory Disease (CC 109)	37.61	0.08 (0.01)	1.081 (1.062-1.101)
Chronic Obstructive Pulmonary Disease (CC 111)	25.81	0.16 (0.01)	1.173 (1.151-1.196)
Fibrosis of Lung and Other Chronic Lung Disorders (CC 112)	5.40	0.08 (0.02)	1.087 (1.055-1.121)
Asthma (CC 113)	11.18	-0.10 (0.01)	0.902 (0.880-0.924)

#3754 Risk Standardized Mortality Ratio for Late-Stage Chronic Kidney Disease (CKD) and End Stage Renal Disease (ESRD) , Submission Last Updated: Jan 09, 2023

Description (CC#)	Percentage	Estimate (Standard Error)	Hazard Ratio (95% confidence interval)
Viral and Unspecified Pneumonia, Pleurisy (CC 116)	23.04	0.09 (0.01)	1.097 (1.074-1.12)
Pleural Effusion/Pneumothorax (CC 117)	18.25	0.24 (0.01)	1.270 (1.244-1.297)
Other Respiratory Disorders (CC 118)	43.32	-0.04 (0.01)	0.963 (0.945-0.981)
Other Ear, Nose, Throat, and Mouth Disorders (CC 131)	37.88	-0.09 (0.01)	0.911 (0.895-0.926)
Kidney Transplant Status: ICD-10-CM codes beginning with 'Z' (CC 132Z) (Z4822 Encounter for aftercare following kidney transplant; and Z940 Kidney transplant status)	6.98	-0.26 (0.02)	0.769 (0.739-0.801)
Dialysis Status (CC 134)	39.74	0.19 (0.02)	1.214 (1.165-1.265)
Acute Renal Failure (CC 135)	35.29	0.13 (0.01)	1.137 (1.114-1.16)
Chronic Kidney Disease, Stage 5 (CC 136)	49.74	0.06 (0.02)	1.066 (1.034-1.099)
Chronic Kidney Disease, Severe (Stage 4) (CC 137)	61.70	-0.46 (0.01)	0.630 (0.615-0.644)
Chronic Kidney Disease, Moderate (Stage 3) (CC 138)	48.60	-0.13 (0.01)	0.877 (0.857-0.897)
Chronic Kidney Disease, Mild or Unspecified (Stages 1-2 or Unspecified) (CC 139)	72.26	-0.08 (0.01)	0.928 (0.909-0.947)
Unspecified Renal Failure (CC 140)	16.19	0.01 (0.01)	1.008 (0.987-1.028)
Nephritis (CC 141)	6.51	-0.10 (0.02)	0.904 (0.873-0.936)
Other Urinary Tract Disorders (CC 145)	42.04	-0.07 (0.01)	0.930 (0.914-0.947)
Pressure Ulcer of Skin with Full Thickness Skin Loss (CC 158)	2.83	0.11 (0.02)	1.114 (1.072-1.157)
Pressure Ulcer of Skin with Partial Thickness Skin Loss (CC 159)	3.02	0.12 (0.02)	1.130 (1.089-1.171)
Pressure Pre-Ulcer Skin Changes or Unspecified Stage (CC 160)	3.82	0.14 (0.02)	1.151 (1.112-1.190)
Chronic Ulcer of Skin, Except Pressure (CC 161)	11.62	0.21 (0.01)	1.235 (1.203-1.267)
Cellulitis, Local Skin Infection (CC 164)	19.28	0.10 (0.01)	1.101 (1.078-1.124)
Vertebral Fractures without Spinal Cord Injury (CC 169)	2.27	0.15 (0.02)	1.167 (1.116-1.221)
Other Injuries (CC 174)	35.56	0.09 (0.01)	1.094 (1.074-1.114)
Major Symptoms, Abnormalities (CC 178)	82.78	0.15 (0.02)	1.158 (1.123-1.195)
Amputation Status, Lower Limb/Amputation Complications (CC 189)	5.45	0.18 (0.02)	1.200 (1.163-1.237)
Chemotherapy (CC 193)	5.17	0.02 (0.02)	1.016 (0.977-1.057)
Screening/Observation/Special Exams (CC 195)	90.69	-0.25 (0.01)	0.782 (0.759-0.805)

Description (CC#)	Percentage	Estimate (Standard Error)	Hazard Ratio (95% confidence interval)
Supplemental Oxygen (CC 197)	6.54	0.24 (0.01)	1.266 (1.233-1.299)
Wheelchairs, Commodes (CC 200)	1.57	0.16 (0.02)	1.179 (1.126-1.235)
Alcohol/Cannabis Use or Use Disorder, Mild or Uncomplicated; Non-Psychoactive Substance Abuse; Nicotine Dependence (CC 203)	12.70	0.11 (0.01)	1.114 (1.089-1.140)

Table 2b.24. Parameter Estimates for Final Risk Variables Using Cox Proportional Hazard Model with Frailty Regression Model, Mortality Development Dataset (N= 758,162 Patients)

CC = condition category (groups of ICD-10 codes)

[Response Ends]

2b.25. Describe the analyses and interpretation resulting in the decision to select or not select social risk factors.

Examples may include prevalence of the factor across measured entities, availability of the data source, empirical association with the outcome, contribution of unique variation in the outcome, or assessment of between-unit effects and within-unit effects. Also describe the impact of adjusting for risk (or making no adjustment) on providers at high or low extremes of risk.

[Response Begins]

Social risk factor testing included:

1. Examining the distribution of SRFs among providers (Table 2b.25:1);
2. Examined bivariate (unadjusted) and multivariate (adjusted) relationships of SRFs with mortality (Table 2b.25:2);
3. Examined risk model performance with and without each SRF (Table 2b.25:2); and
4. Examined provider performance stratified across quintiles of each SRF (Table 2b.25:3, Figures 2b.25:1-4).

The prevalence of SRFs in the Progression Measure cohort varies across 2,854 measured entities as shown in Table 2b.25:1. At the median provider, 23.1% of patients are dual-eligible; 22.2% have bottom-quartile AHRQ SES; 14.3% are of Black race; and 97.3% live in urban areas.

SRF	Median provider-level SRF prevalence (IQR)
Dual Eligibility	23.1% (14.5%-37.2%)
Low AHRQ SES	22.2% (6.7%-42.9%)
Race (Black)	14.3% (3.7%-36.0%)
Urban	97.3% (76.1%-100%)

Table 2b.25:1 Provider-level distribution of social risk factors in the Mortality Measure cohort

Next, Table 2b.25:2 examines the bivariate (unadjusted) relationships of SRFs with mortality and the risk-adjusted relationships and risk model performance when incorporating SRFs. The c-statistic of the clinical model with 71 risk variables using a Cox Proportional Hazard Regression Model for comparison is 0.735, which indicates strong model discrimination.

Social Risk Factors	Unadjusted (Bivariate) estimate (SE)	Unadjusted Hazard Ratio (95% CI)	Adjusted (Multivariate) estimate (SE)	Adjusted Hazard Ratio (95% CI)	C-statistic
None (clinical risk model, 71 factors)	*	*	*	*	0.735
Dual Eligibility	0.238 (0.009)	1.268 (1.246-1.29)	0.047 (0.01)	1.048 (1.028-1.068)	0.735
Low AHRQ SES	0.057 (0.009)	1.059 (1.04-1.078)	0.021 (0.009)	1.021 (1.003-1.039)	0.735
Race (Black)	-0.041 (0.01)	0.959 (0.942-0.978)	-0.078 (0.01)	0.925 (0.906-0.944)	0.736
Urban	-0.074 (0.01)	0.929 (0.911-0.947)	-0.117 (0.01)	0.890 (0.872-0.908)	0.736

Table 2b.25:2. Bivariate Associations and Multivariate Associations Using Cox Proportional Hazard Regression Models Between SRF and Outcome (Mortality) (N= 758,162)

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- Dual eligibility: The unadjusted hazard ratio (1.268) suggests dual-eligible patients have a higher risk of death than those who are not dual-eligible. Adjustment for comorbidities greatly attenuates that risk (adjusted hazard ratio 1.048), though it remains statistically significant. This suggests that the increased risk of death associated with dual eligibility is largely explained by greater comorbidity among dual-eligible patients captured by the clinical risk model.
- AHRQ SES Index: Lower neighborhood economic status is slightly (though significantly) associated with mortality in the unadjusted model (hazard ratio 1.059); this is attenuated (though still significant) in the adjusted model (hazard ratio 1.021). This suggests that some association between SES and mortality may exist that is not explained by comorbidities, but the practical significance of this is minimal.
- Black Race: The unadjusted hazard ratio (0.959) suggests Black patients have a slightly but significantly lower risk of death than non-Black patients. Adjustment for comorbidities further reduces the relative risk for Black patients (hazard ratio 0.925, still significant). This suggests that non-Black patients have a higher risk of death that is not fully explained by comorbidities.
- Urban: Patients living in urban areas have a significantly lower risk of mortality than patients in non-urban areas (unadjusted hazard ratio 0.929, adjusted ratio 0.890). This suggests that patients in non-urban areas may have higher risk of CKD or ESRD mortality that is not fully explained by comorbidities.

Finally, we examined the correlations between provider's RSMR and the proportion of their patients in the "disadvantaged" group by each SRF, both overall and within each quintile of the SRF. As shown in Table 2b.25:3 and Figures 2b.25:1-4 below, there is no significant association between a provider's risk adjusted score and their proportion of dual-eligible or low-SES, or urban patients, a small significant association with percent Black race, and a modest significant association with percent urban residence. The lack of systematic variation based on the distribution of each SRF among providers indicates that variation in providers' performance is dominated by other factors and there is no need for further adjustment. These findings complement our previously stated conceptual rationales for not adjusting for these factors and provide reassurance that, given the constraints of feasibly available data, there are no key patient-level factors which the measure fails to account for.

Quintile	Black race (p-value)	Dual-eligible (p-value)	Low SES (p-value)	Urban residence (p-value)
1st	-0.135 (0.003)	-0.062 (0.176)	-0.040 (0.380)	-0.044 (0.331)
2nd	0.002 (0.967)	0.001 (0.989)	-0.050 (0.271)	-0.036 (0.425)
3rd	-0.051 (0.264)	0.005 (0.910)	-0.030 (0.513)	-0.010 (0.822)

Quintile	Black race (p-value)	Dual-eligible (p-value)	Low SES (p-value)	Urban residence (p-value)
4th	0.015 (0.746)	0.009 (0.844)	0.023 (0.618)	-0.021 (0.687)
5th	-0.029 (0.522)	-0.020 (0.664)	0.072 (0.116)	n/a*
Total	-0.060 (0.003)	-0.015 (0.463)	0.021 (0.306)	-0.176 (<.0001)

Table 2b.25:3. Correlation coefficients between RSMR (provider-level score) & proportion of patients in disadvantaged group by SRF.

*More than 20% of providers have 100% urban patients so there is no variation within the top quintile.

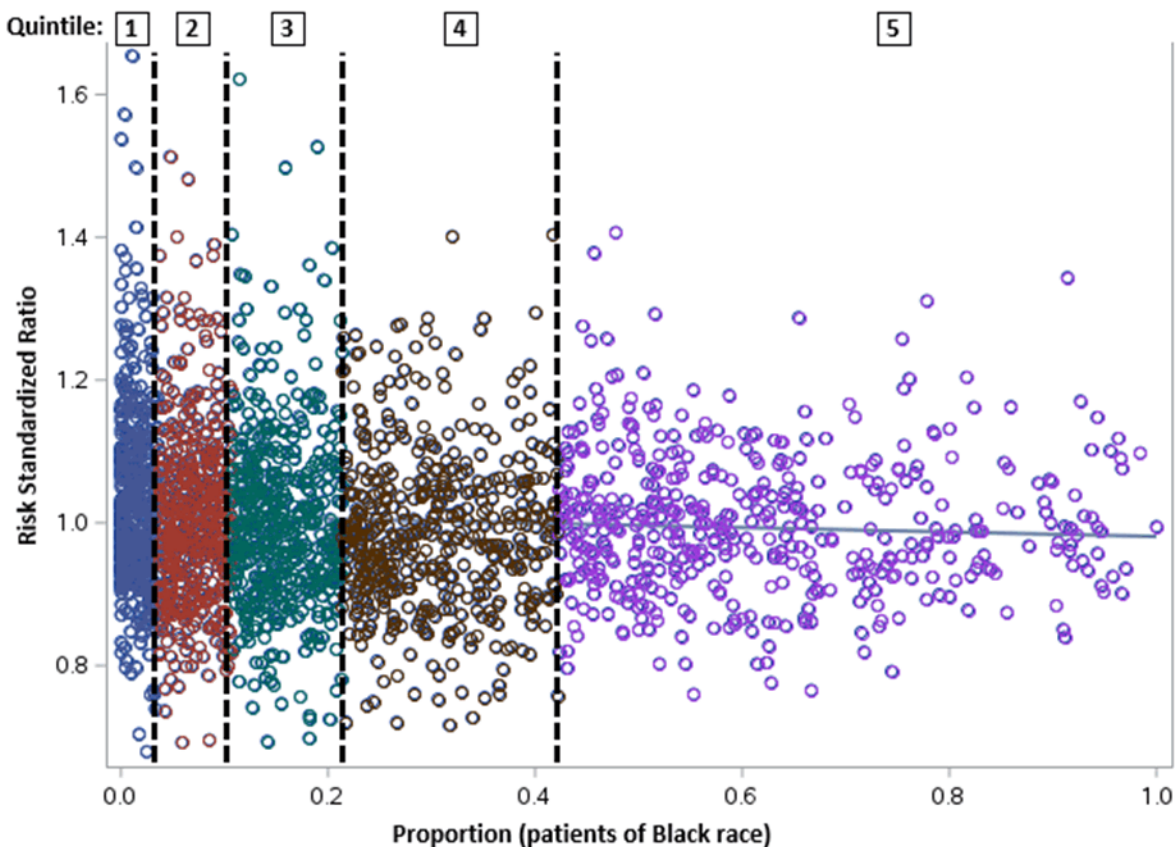


Figure 2b.25:1. RSMR by Percent Patients of Black Race (Mortality Development Dataset)

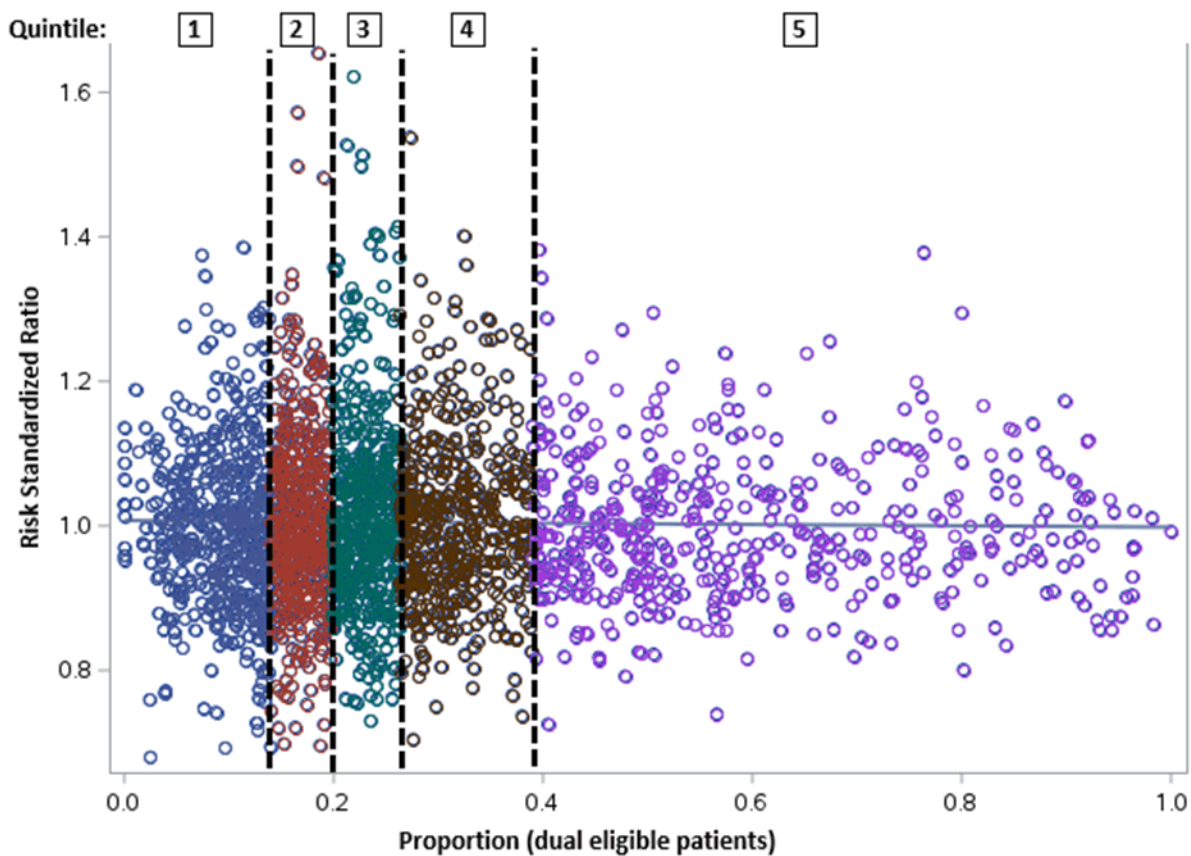


Figure 2b.25:2. RSMR by Percent Dual-eligible Patients (Mortality Development Dataset)

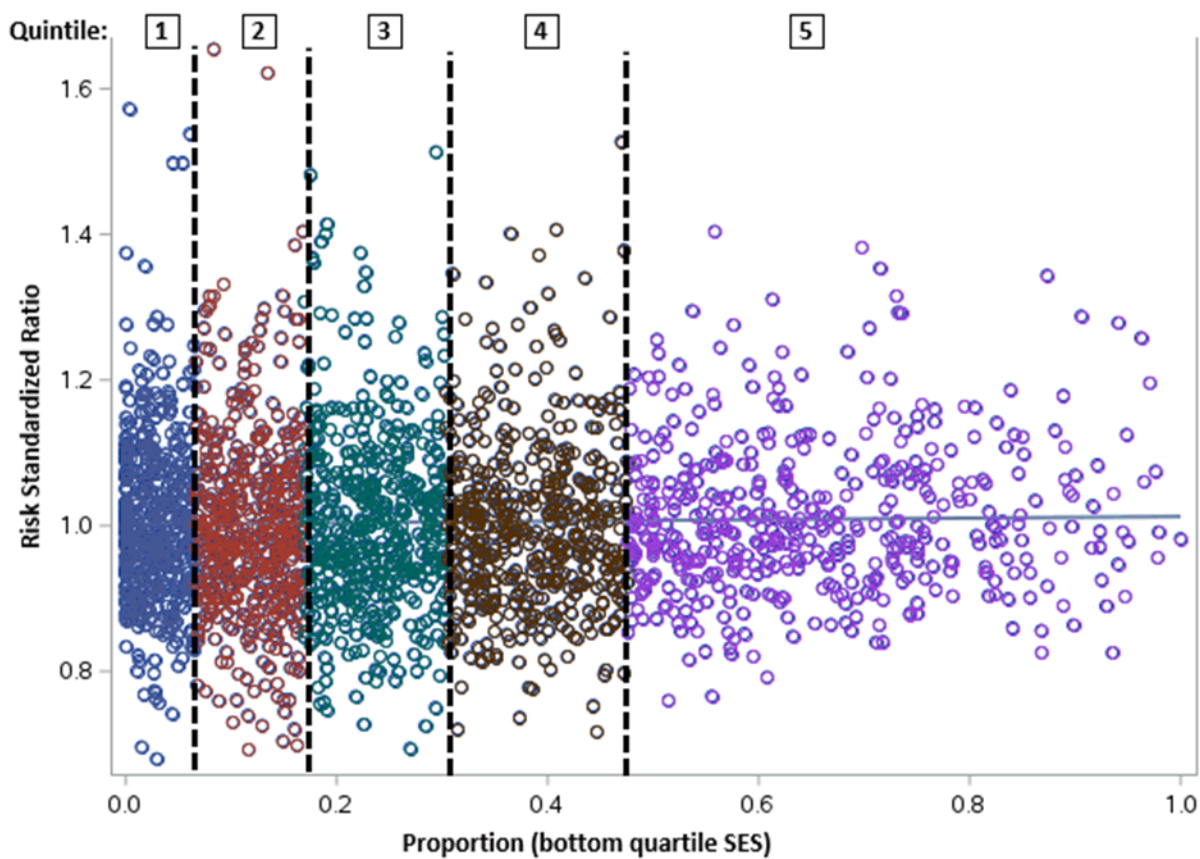


Figure 2b.25:3. RSMR by Percent Patients of Low SES (Mortality Development Dataset)

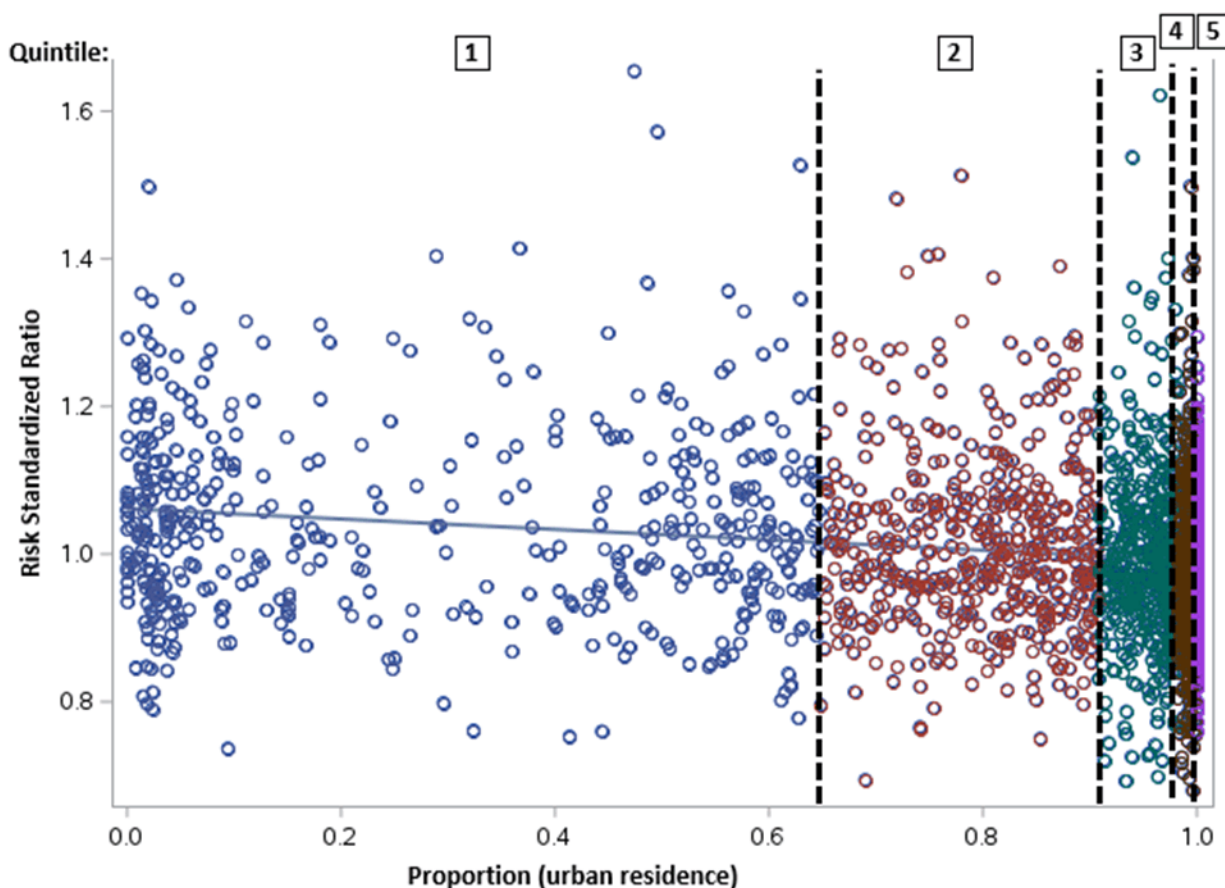


Figure 2b.25:4. RSMR by Percent Urban Patients (Mortality Development Dataset)

Summary and Conclusion

Based on our conceptual model, we examined the impact of including four social risk factors (dual eligibility, low AHRQ SES, race, and urbanicity) on the CKD and ESRD Mortality Measure. Importantly, we did not see a substantial relationship between any of the social risk factors we tested and measure scores, even among nephrology practices with the highest proportion of disadvantaged patients. We found that while odds of the outcome in a bivariate model are higher among patients with dual eligibility and low AHRQ SES, the relationship between the variable and the outcome is greatly attenuated in a multivariable model, suggesting that the clinical risk variables account for most of the risk. After risk adjustment, non-urban residents are somewhat more likely to die compared to those patients living in urban areas, while odds of death were less for Black than for non-Black patients.

Importantly, there is no statistically significant relationship between any of the social risk factors we tested and measure scores among nephrology practices with the highest proportion of patients with social risk factors. Therefore, because there is minimal impact on provider scores, and due to the tradeoff between unintended consequences of adjusting for social risk factors and potentially masking differential care for patients with social risk factors, in particular for Black patients, we did not include social risk factors in the final model. We will revisit this decision during periodic re-evaluation of the measure.

[Response Ends]

2b.26. Describe the method of testing/analysis used to develop and validate the adequacy of the statistical model or stratification approach (describe the steps—do not just name a method; what statistical analysis was used). Provide the statistical results from testing the approach to control for differences in patient

characteristics (i.e., case mix) below. If stratified ONLY, enter “N/A” for questions about the statistical risk model discrimination and calibration statistics.

Validation testing should be conducted in a data set that is separate from the one used to develop the model.

[Response Begins]

To test model performance, we divided the mortality development dataset randomly in half into a “derivation sample” and a “validation sample.” We fit the model to the data in the derivation sample to specify the model coefficients in a hierarchical logistic regression model, then used those coefficients in the validation sample to confirm the model is generalizable and well-calibrated.

We computed three summary statistics for assessing model performance¹.

Discrimination Statistics

(1) Area under the receiver operating characteristic (ROC) curve (the c-statistic) is the probability that predicting the outcome is better than chance, which is a measure of how accurately a statistical model is able to distinguish between a patient with and without an outcome)

(2) Predictive ability (discrimination in predictive ability measures the ability to distinguish high-risk subjects from low-risk subjects; therefore, we would hope to see a wide range between the lowest decile and highest decile.)

Calibration Statistics

(3) Over-fitting indices (over-fitting refers to the phenomenon in which a model accurately describes the relationship between predictive variables and outcome in the development dataset but fails to provide valid predictions in new patients)

In addition, we plotted calibration curves (Figures 2b.29:1-2) comparing the observed to predicted mortality at the patient level, within each decile of predicted mortality, in both the derivation and validation samples. A well-specified and calibrated model will demonstrate a) clear increasing trend; b) close correspondence between the predictions and the observations; and c) similar results in the validation as the derivation sample.

Reference:

1. Harrell FE and Shih YC. Using full probability models to compute probabilities of actual interest to decision makers, Int. J. Technol. Assess. Health Care 17 (2001), pp. 17–26.

[Response Ends]

2b.27. Provide risk model discrimination statistics.

For example, provide c-statistics or R-squared values.

[Response Begins]

The Harrell’s C-statistic for the full Mortality Development Dataset sample (n=758,162), evaluating the risk model using Cox proportional hazard model, is 0.735. Table 2b.27 shows our model testing results for the derivation and validation samples.

Model Performance Statistic	Derivation Sample	Validation Sample
Number of Patients	379,081	379,081
Mortality Risk	7.92%	7.98%
Calibration (y0, y1)	(0, 1)	(-0.001, 0.996)
Discrimination- Predictive ability (lowest decile %- highest decile %)	(1.2%, 22.9%)	(1.2%, 23.1%)

Model Performance Statistic	Derivation Sample	Validation Sample
C-statistic	0.734	0.734

Table 2b.27. Risk Model Performance, Mortality Development Dataset (N=758,162)

[Response Ends]

2b.28. Provide the statistical risk model calibration statistics (e.g., Hosmer-Lemeshow statistic).

[Response Begins]

Please see table 2b.27 above for calibration (overfitting) results; please see section 2b.29 for risk-decile plots.

[Response Ends]

2b.29. Provide the risk decile plots or calibration curves used in calibrating the statistical risk model.

The preferred file format is .png, but most image formats are acceptable.

[Response Begins]

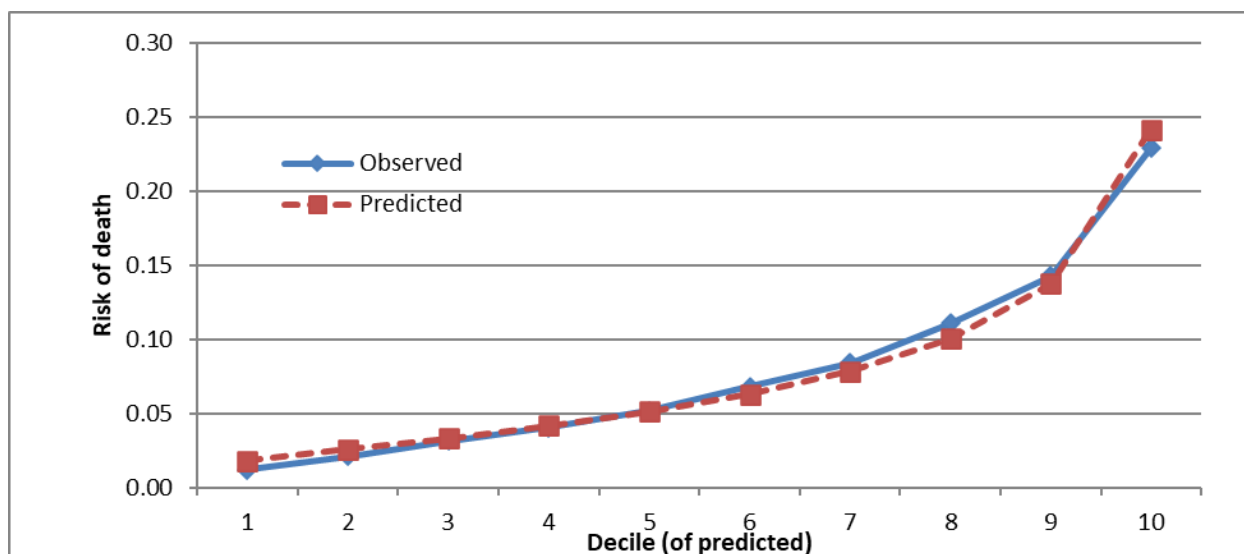


Figure 2b.29:1. Observed vs. predicted mortality risk by decile of predicted (derivation sample)

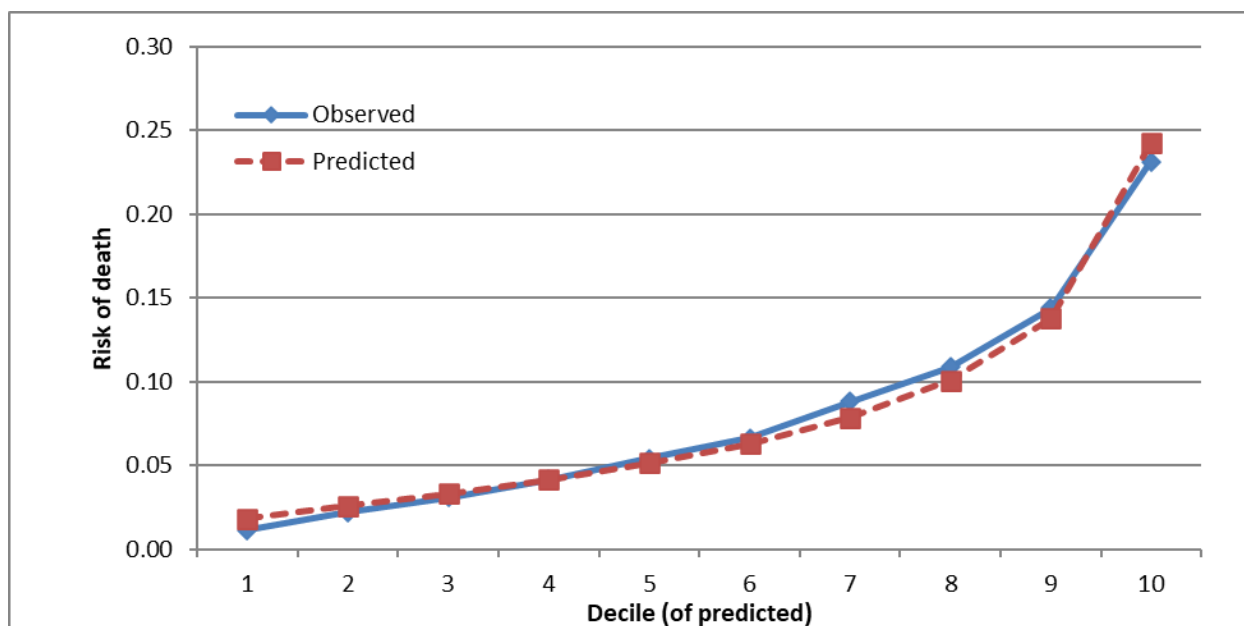


Figure 2b.29:2. Observed vs. predicted mortality risk by decile of predicted (validation sample)

[Response Ends]

2b.30. Provide the results of the risk stratification analysis.

[Response Begins]

Not applicable; measure is not stratified.

[Response Ends]

2b.31. Provide your interpretation of the results, in terms of demonstrating adequacy of controlling for differences in patient characteristics (i.e., case mix).

In other words, what do the results mean and what are the norms for the test conducted?

[Response Begins]

Discrimination Statistics

The C-statistics were 0.735 in the full dataset and 0.734 in both the derivation and validation sample, indicating good model discrimination. The model's predictive ability shows a wide range between the lowest decile and highest decile in both samples, indicating the ability to distinguish high-risk subjects from low-risk subjects.

Calibration (over-fitting) Statistics (γ_0 , γ_1)

If the γ_0 in the validation samples are substantially far from zero and the γ_1 is substantially far from 1, there is potential evidence of over-fitting. The validation sample calibration values of $\gamma_0 = -0.001$ and $\gamma_1 = 0.996$ indicate good calibration of the model.

Risk Decile Plots

Higher deciles of the predicted outcomes are closely associated with higher observed outcomes, which show a good calibration of the model. These plots indicates excellent discrimination of the model and good predictive ability. The similar results in both the derivation and validation samples indicates the model has high generalizability.

Overall Interpretation

Interpreted together, our diagnostic results demonstrate the risk-adjustment model adequately controls for differences in patient characteristics.

[Response Ends]

2b.32. Describe any additional testing conducted to justify the risk adjustment approach used in specifying the measure.

Not required but would provide additional support of adequacy of the risk model, e.g., testing of risk model in another data set; sensitivity analysis for missing data; other methods that were assessed.

[Response Begins]

No additional testing.

[Response Ends]

3. Feasibility

Extent to which the specifications including measure logic, require data that are readily available or could be captured without undue burden and can be implemented for performance measurement.

3.01. Check all methods below that are used to generate the data elements needed to compute the measure score.

[Response Begins]

[Response Ends]

3.02. Detail to what extent the specified data elements are available electronically in defined fields.

In other words, indicate whether data elements that are needed to compute the performance measure score are in defined, computer-readable fields.

[Response Begins]

[Response Ends]

3.03. If ALL the data elements needed to compute the performance measure score are not from electronic sources, specify a credible, near-term path to electronic capture, OR provide a rationale for using data elements not from electronic sources.

[Response Begins]

[Response Ends]

3.04. Describe any efforts to develop an eCQM.

[Response Begins]

[Response Ends]

3.06. Describe difficulties (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.

[Response Begins]

[Response Ends]

Consider implications for both individuals providing data (patients, service recipients, respondents) and those whose performance is being measured.

3.07. Detail any fees, licensing, or other requirements to use any aspect of the measure as specified (e.g., value/code set, risk model, programming code, algorithm),

Attach the fee schedule here, if applicable.

[Response Begins]

[Response Ends]

4. Usability and Use

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

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.

NQF-endorsed measures are expected to be used in at least one accountability application within 3 years and publicly reported within 6 years of initial endorsement, in addition to demonstrating performance improvement.

4a.01. Check all current uses. For each current use checked, please provide:

- Name of program and sponsor
- URL
- Purpose
- Geographic area and number and percentage of accountable entities and patients included
- Level of measurement and setting

[Response Begins]

[Response Ends]

4a.02. Check all planned uses.

[Response Begins]

[Response Ends]

4a.03. If not currently publicly reported OR used in at least one other accountability application (e.g., payment program, certification, licensing), explain why the measure is not in use.

For example, do policies or actions of the developer/steward or accountable entities restrict access to performance results or block implementation?

[Response Begins]

[Response Ends]

4a.04. If not currently publicly reported OR used in at least one other accountability application, provide a credible plan for implementation within the expected timeframes: used in any accountability application within 3 years, and publicly reported within 6 years of initial endorsement.

A credible plan includes the specific program, purpose, intended audience, and timeline for implementing the measure within the specified timeframes. A plan for accountability applications addresses mechanisms for data aggregation and reporting.

[Response Begins]

[Response Ends]

4a.05. Describe how performance results, data, and assistance with interpretation have been provided to those being measured or other users during development or implementation.

Detail how many and which types of measured entities and/or others were included. If only a sample of measured entities were included, describe the full population and how the sample was selected.

[Response Begins]

[Response Ends]

4a.06. Describe the process for providing measure results, including when/how often results were provided, what data were provided, what educational/explanatory efforts were made, etc.

[Response Begins]

[Response Ends]

4a.07. Summarize the feedback on measure performance and implementation from the measured entities and others. Describe how feedback was obtained.

[Response Begins]

[Response Ends]

4a.08. Summarize the feedback obtained from those being measured.

[Response Begins]

[Response Ends]

4a.09. Summarize the feedback obtained from other users.

[Response Begins]

[Response Ends]

4a.10. Describe how the feedback described has been considered when developing or revising the measure specifications or implementation, including whether the measure was modified and why or why not.

[Response Begins]

[Response Ends]

4b.01. You may refer to data provided in Importance to Measure and Report: Gap in Care/Disparities, but do not repeat here. Discuss any progress on improvement (trends in performance results, number and percentage of people receiving high-quality healthcare; Geographic area and number and percentage of accountable entities and patients included). If no improvement was demonstrated, provide an explanation. If not in use for performance improvement at the time of initial endorsement, provide a credible rationale that describes how the performance results could be used to further the goal of high-quality, efficient healthcare for individuals or populations.

[Response Begins]

[Response Ends]

4b.02. Explain any unexpected findings (positive or negative) during implementation of this measure, including unintended impacts on patients.

[Response Begins]

[Response Ends]

4b.03. Explain any unexpected benefits realized from implementation of this measure.

[Response Begins]

[Response Ends]

5. Comparison to Related or 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.

If you are updating a maintenance measure submission for the first time in MIMS, please note that the previous related and competing data appearing in question 5.03 may need to be entered in to 5.01 and 5.02, if the measures are NQF endorsed. Please review and update questions 5.01, 5.02, and 5.03 accordingly.

5.01. Search and select all NQF-endorsed related measures (conceptually, either same measure focus or target population).

NOTE: If there are no related measures, please select N/A.

(Can search and select measures.)

[Response Begins]

[Response Ends]

5.02. Search and select all NQF-endorsed competing measures (conceptually, the measures have both the same measure focus and target population).

NOTE: If there are no competing measures, please select N/A.

(Can search and select measures.)

[Response Begins]

[Response Ends]

5.03. If there are related or competing measures to this measure, but they are not NQF-endorsed, please indicate the measure title and steward.

[Response Begins]

[Response Ends]

5.04. If this measure conceptually addresses EITHER the same measure focus OR the same target population as NQF-endorsed measure(s), indicate whether the measure specifications are harmonized to the extent possible.

[Response Begins]

[Response Ends]

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

[Response Begins]

[Response Ends]

5.06. Describe why this measure is superior to competing measures (e.g., a more valid or efficient way to measure quality). Alternatively, justify endorsing an additional measure.

Provide analyses when possible.

[Response Begins]

[Response Ends]

Appendix

Supplemental materials may be provided in an appendix.:

Available in attached file

Contact Information

Measure Steward (Intellectual Property Owner): Centers for Medicare & Medicaid Services

Measure Steward Point of Contact: Day, Tim, timothy.day@cms.hhs.gov

Measure Developer if different from Measure Steward: Yale New Haven Health Services Corporation – Center for Outcomes Research and Evaluation (CORE)

Measure Developer Point(s) of Contact: Peter, Doris, doris.peter@yale.edu

Bagshaw, Kyle, kyle.bagshaw@yale.edu

Additional Information

1. Provide any supplemental materials, if needed, as an appendix. All supplemental materials (such as data collection instrument or methodology reports) should be collated one file with a table of contents or bookmarks. If material pertains to a specific criterion, that should be indicated.

[Response Begins]

Available in attached file

[Response Ends]

2. List the workgroup/panel members' names and organizations.

Describe the members' role in measure development.

[Response Begins]

[Response Ends]

3. Indicate the year the measure was first released.

[Response Begins]

[Response Ends]

4. Indicate the month and year of the most recent revision.

[Response Begins]

[Response Ends]

5. Indicate the frequency of review, or an update schedule, for this measure.

[Response Begins]

[Response Ends]

6. Indicate the next scheduled update or review of this measure.

[Response Begins]

[Response Ends]

7. Provide a copyright statement, if applicable. Otherwise, indicate "N/A".

[Response Begins]

[Response Ends]

8. State any disclaimers, if applicable. Otherwise, indicate "N/A".

[Response Begins]

[Response Ends]

9. Provide any additional information or comments, if applicable. Otherwise, indicate "N/A".

[Response Begins]

[Response Ends]