Building a Roadmap From Patient-Reported Outcome Measures to Patient-Reported Outcome Performance Measures

TECHNICAL GUIDANCE – UPDATED FINAL DRAFT

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Executive Summary

The Building a Roadmap From Patient-Reported Outcome Measures to Patient-Reported Outcome Performance Measures initiative (henceforth referred to as Building a Roadmap) provides guidance to measure developers on the development of digital patient-reported outcome performance measures (PRO-PMs). This work is funded by the Centers for Medicare & Medicaid Services (CMS) and led by the National Quality Forum (NQF), both of whom have a long-standing shared commitment to advance the use of patient-reported outcomes (PROs) and digital measurement.

CMS identifies the importance of digital quality measurement across its priority initiatives (e.g., the Digital Quality Measures [dQM] Strategic Roadmap and the Meaningful Measures Framework). Digital PRO-PMs are an important component of the future of healthcare, elevating the patient’s voice in the assessment of healthcare quality while reducing measurement burden. However, PRO-PMs compose only 7 percent of all NQF-endorsed quality measures. To increasingly incorporate the patient perspective, measure developers (i.e., the person or team that is developing the digital PRO-PM) need meaningful resources to help them navigate the development of digital PRO-PMs.

This Technical Guidance Report (henceforth referred to as the Roadmap) is a resource for measure developers who are developing digital PRO-PMs. It compiles existing resources on measure development, endorsement, and interoperability and frames them through the lens of patient-reported performance measurement. NQF convened a Technical Expert Panel (TEP) (Appendix A) to inform the Roadmap, which consists of four stages composed of 17 tasks:

- **Stage 1: Definition of Measurement Goals.** Measure developers set goals for the measure concept and the development of a PRO-PM.

- **Stage 2: Exploration and Assessment of Patient-Reported Outcome Measures.** Measure developers identify, assess, and select the patient-reported outcome measure(s) (PROM[s]) that will collect data for the PRO-PM.

- **Stage 3: Development and Testing of the PRO-PM.** Measure developers specify the details of the measure, select test sites, and create and execute testing plans.

- **Stage 4: Finalization and Implementation of the PRO-PM.** Measure developers prepare for implementation and the Consensus Development Process (CDP).

The Roadmap is designed so that measure developers can complete recommended tasks within or across stages, ensuring consistent but flexible guidance to individuals and organizations that approach measure development differently.
Figure 1. PRO-PM Roadmap: Each column contains one stage, and the bidirectional arrows indicate that tasks can move freely and be iterated across stages.

This Roadmap is an updated version of the original Roadmap published by NQF in November 2021. NQF updated the Roadmap based on a series of key informant interviews (KIIs) with measurement and interoperability experts and guidance from the TEP. Improvements include the following:

- A new stage 1 task titled “Review Relevant Resources,” which guides measure developers to a collection of source-of-truth resources. The addition of this task ensures the Roadmap will stay relevant and meaningful by linking to permanent webpages (when possible) for general guidance on measure development, NQF endorsement, and interoperability. NQF removed detailed information that mirrored source-of-truth resources.
- Tailored information on elements of development, testing, and endorsement that are unique to digital PRO-PMs.
- Additional examples of how patients can contribute to digital PRO-PM development.
- New information on digital measurement, including references to the Electronic Clinical Quality Improvement (eCQI) Resource Center page on dQMs and an example of how machine-readable and narrative specifications differ.

The Roadmap is a guide to help measure developers navigate the steps required to develop a high quality PRO-PM. It is also a catalyst to build a more robust database of digital PRO-PMs, which will ultimately elevate patient voices and prioritize outcomes that matter to patients.
Introduction

Digital PRO-PMs offer an important opportunity to ensure that patient voices are captured in assessments of quality while supporting learning health systems. Policymakers, payers, and healthcare providers are increasingly using PRO-PMs to inform clinical decision making, improve quality of care, modify provider payment, and evaluate the value of medical technologies. This growth aligns with the CMS Meaningful Measures Framework, which includes Meaningful Measures 2.0 (e.g., prioritizing outcome measures and PROs), and the CMS National Quality Strategy goals (e.g., foster engagement between individuals and their care teams, embrace the digital age, and increase alignment of performance metrics across stakeholders).

Despite this growth, PRO-PMs make up only 7 percent of all NQF-endorsed quality measures, and only one PRO-PM was included on CMS’ list of Merit-Based Incentive Payment System (MIPS) quality measures for 2022. Challenges hamper the broad adoption of these measures. PROMs, the tools on which PRO-PMs are based, have not yet become standard practice in clinical use, and some healthcare professionals do not yet understand their benefits. Patients lack awareness about the benefits of PROMs and PRO-PMs, such as the ability to compare outcomes across different clinicians, hospitals, and health plans. PRO-PMs are complex measures, and measure developers lack thorough yet accessible technical guidance to develop high quality performance measures based on patient-reported data.

Goals and Objectives

The Roadmap aims to provide measure developers with thorough yet accessible guidance on developing digital PRO-PMs, from identifying a measure concept to preparing to submit the PRO-PM for NQF endorsement review. The Roadmap is intended for novice and advanced measure developers alike. Although its guidance is generally applicable to all PRO-PMs, it specifically focuses on digital PRO-PMs that are intended for use in CMS’ value-based purchasing (VBP) programs and alternative payment models (APMs) and can be calculated and transmitted across interoperable health information technology (IT) systems. dQMs are standardized measures that utilize electronic data from multiple sources generated during the normal course of clinical care, and thus allow for more efficient capture, retrieval, and sharing of patient data. PRO-PMs are well suited to be dQMs because PROMs often have structured data fields, and PRO-PMs may require information from multiple data sources (e.g., PROM data, electronic health record [EHR] data, and claims data).

The Roadmap describes how to develop a digital PRO-PM that is:

- meaningful to patients and clinicians;
- aligned with best and promising practices for performance measurement;
- appropriate for use in CMS’ VBP programs and APMs;
- usable by public and private payers; and
- consistent with the standards of scientific acceptability.

The Roadmap is a high-level guide that helps measure developers understand a complex process, ask educated questions, and contribute to a more robust database of digital PRO-PMs. It is intended to be used as a companion alongside existing measure development resources (Appendix B).
Background

Although PRO-PMs make up a small percentage of the measures used in CMS’ VBP programs and APMs, CMS sees the broader development and use of these performance measures as an important part of its evolving initiatives to incentivize high quality care. Following the passage of the Affordable Care Act (ACA) in 2010, CMS adopted new VBP programs that shifted towards improving and rewarding value rather than volume. CMS designed the VBP programs to ensure that healthcare was more person centered by creating care that focuses on patients’ preferences and desired outcomes. Several VBP programs apply to different provider settings, such as hospitals and outpatient centers. CMS utilizes VBP programs and APMs to incentivize eligible participants to provide high quality and cost-efficient care. These programs are likely to interact during the shift toward improving value, given that incentives linked to APMs may be similar to VBP programs for some providers.

CMS has a long-standing commitment to PRO-PMs and has partnered with NQF for more than 10 years to advance the knowledge of patient-centered performance measurement through a series of foundational reports. This body of work includes, but is not limited to, the following:

- 2012: Methodological Issues in the Selection, Administration, and Use of Patient-Reported Outcomes in Performance Measurement in Health Care Settings (henceforth referred to as Methodological Issues; this report was updated in 2015 by its authors.)
- 2012: PRO-Based Performance Measures for Healthcare Accountable Entities
- 2013: Patient-Reported Outcomes in Performance Measurement
- 2020: Patient-Reported Outcomes: Best Practices on Selection and Data Collection (henceforth referred to as PRO Best Practices)

The Roadmap is the final report in the Building a Roadmap initiative, which is the most recent PRO-PM collaboration between CMS and NQF, and includes four distinct reports:

1. The Environmental Scan Report identifies and summarizes existing information relevant to the use of high quality PROMs as the basis for PRO-PMs in accountability programs. It is the first report in the Building a Roadmap initiative and provides background for the subsequent publications. The report was originally published in spring 2021 and updated in June 2022.
2. The Interim Report identifies and describes 12 attributes of PROMs that are suitable data collection instruments for high quality, digital PRO-PMs. It provides guidance to measure developers on selecting PROMs for use in a PRO-PM and is best read alongside the Roadmap.
3. The Developer Feedback Report summarizes improvement opportunities for the Roadmap from measure developers and health IT experts.
4. The Technical Guidance Report describes a series of four stages and 17 tasks that measure developers can follow when developing and testing digital PRO-PMs that are suitable for CMS’ VBP programs and APMs. The Roadmap was originally published in November 2021 and updated in November 2022.

Methodology

NQF began developing the Roadmap by convening a multistakeholder TEP that met 14 times over the course of two years. The TEP comprised diverse viewpoints that represented patients, patient advocates, measure developers, health IT professionals, clinicians, health systems, payers, purchasers,
and researchers. Due to the technical nature of the Roadmap, NQF intentionally included multiple PROM and PRO-PM developers on the TEP. These individuals acknowledged their work during the disclosure of interest portion of the TEP’s first web meeting, which was open to the public. In addition to the TEP, liaisons from federal agencies provided guidance on PRO-PMs from the federal perspective. Twelve experts participated in KII s that identified improvement opportunities for the Roadmap. (A detailed description of the KII methodology is available in the Developer Feedback Report.) Unless a fact or recommendation is explicitly attributed to a specific source, information in the Roadmap comes from the TEP and the KII s and is synthesized by NQF. The Building a Roadmap initiative does not recommend any PROM, nor does it identify any PROM as being “high quality.” NQF does not currently endorse, recommend, rank, or prioritize PROMs. This report includes specific PROMs only as examples.

Terminology

The definitions for PRO, PROM, and PRO-PM used in the Roadmap (Table 1) align with the definitions used in the CMS Measures Management System Blueprint (henceforth referred to as the CMS Blueprint). Because the terminology regarding PRO-PMs is highly technical, the report includes a glossary of key terms (Appendix C).

Table 1: Distinctions Among PROs, PROMs, and PRO-PMs

<table>
<thead>
<tr>
<th>Concept</th>
<th>Definition</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient-Reported Outcome (PRO)</td>
<td>What gets measured. The status of a patient’s health condition or health behavior that comes directly from the patient (i.e., outcome data)</td>
<td>Symptoms of depression</td>
</tr>
<tr>
<td>Patient-Reported Outcome Measure (PROM)</td>
<td>How PROs are measured. The tools/instruments used to collect data</td>
<td>Patient Health Questionnaire 9 (PHQ-9©, a standardized tool to assess depression</td>
</tr>
<tr>
<td>Patient-Reported Outcome Performance Measure (PRO-PM)</td>
<td>How PROs are calculated. A way to aggregate the information from patients into a reliable, valid measure of performance</td>
<td>Percentage of patients with a diagnosis of major depression or dysthymia and an initial PHQ-9 score greater than 9 with a follow-up PHQ-9 score less than 5 at 12 months (NQF #0710e)</td>
</tr>
</tbody>
</table>

The Roadmap focuses on digital PRO-PMs, which are a type of dQM. The Roadmap also discusses electronic clinical quality measures (eCQMs).

- **Digital quality measures:** Although the definition of dQMs is evolving, the draft definition is as follows: “Quality measures, organized as self-contained measure specifications and code packages, that use one or more sources of health information that is captured and can be transmitted electronically via interoperable systems. Data sources for dQMs may include administrative systems, electronically submitted clinical assessment data, case management systems, EHRs, laboratory systems, prescription drug monitoring programs, instruments (for example, medical devices and wearable devices), patient portals or applications (for example,
for collection of patient-generated data such as a home blood pressure monitor, or patient-reported health data), health information exchanges, or registries, and other sources."

- **Electronic clinical quality measures:** eCQMs are clinical quality measures expressed and formatted to use data primarily from EHRs to measure healthcare quality, ideally data captured in structured form during the process of patient care. For the measured entity to report an eCQM from an EHR, eCQM developers format the Health Quality Measure Format (HQMF) using the Quality Data Model (QDM) to define the data elements and Clinical Quality Language (CQL) to express the logic needed to evaluate a provider or organization’s performance.

The Roadmap discusses two different concepts that have similar terminologies, so it is important that readers understand these differences:

- **Attributes and the Attribute Grid:** Throughout the Building a Roadmap initiative, the word *attribute* is used to describe certain characteristics or traits of a high quality PROM that is suitable for use in a digital PRO-PM. These attributes are collected in a table (i.e., the Attribute Grid, Appendix D) that is designed to facilitate a side-by-side comparison of different PROMs.

- **Attribution:** A process used in quality measurement that aims to assign accountability for a patient’s outcomes to a clinician, groups of clinicians, or a facility.

Throughout the Roadmap, the term *measure developer* refers to the person/team that is developing the PRO-PM. The term *PROM developer* refers to the person/team that developed the PROM. The Roadmap notes the importance of active communication between these two distinct roles. In the uncommon scenario where a single organization acts as both the PROM developer and the measure developer, this communication remains important.

**Environmental Scan Findings**

The Environmental Scan summarized information relevant to the use of high quality PROMs as the basis for digital PRO-PMs in accountability programs and provided background for the other documents in the Building a Roadmap initiative. Key findings from the Environmental Scan Report included the following:

- **NQF-endorsed PRO-PMs are currently limited in number:** At the time of the Roadmap’s publication, only 7 percent of all NQF-endorsed measures were PRO-PMs. The gap between the total number of NQF-endorsed measures, the number of NQF-endorsed PRO-PMs, and the number of PRO-PMs used in CMS accountability programs constrains CMS’ ability to elevate the patient voice. Challenges in development, including resource limitations (e.g., finances, time, and staff) and a lack of clear guidance, contribute to this gap.

- **PRO-PMs can use data from one PROM or many different PROMs:** While most NQF-endorsed PRO-PMs collect data from a single PROM, there can be benefits to using two or more different PROMs to measure the quality concept of interest. Advantages and disadvantages to both approaches are discussed in the stage 2 task titled “Select PROMs for Inclusion in the PRO-PM.”

- **Digital PRO-PMs align with CMS’ priorities of reduced burden and elevated patient voices:** CMS’ Meaningful Measures 2.0 acknowledges the importance of collecting and using patient-reported data across digital systems, the transformation of measures to fully digital (i.e., allowing for data entry, storage, integration, calculation, and reporting to be conducted by EHRs and other health IT systems), and the reduction of measurement burden on clinicians and
patients. These priorities can be addressed, in part, through the use of digital PRO-PMs, which are intended to collect data generated during the normal course of clinical care.

- **Data collection methods are an important consideration in the development of PRO-PMs:** Stakeholders agree that modes of PROM administration and methods of data collection are important to consider when developing PRO-PMs. NQF’s PRO Best Practices Report focuses on the importance of using multiple modes to maximize patient participation and response. Patient-level factors may also impact PROM-collected data.

**Environmental Scan Findings on Interoperability**

The Environmental Scan Report also identified the importance of interoperability to the ongoing work to advance data standardization and data sharing, as well as opportunities for measure developers to contribute to national priority setting on advancing interoperability for specific data classes and data elements. The widespread use of PROMs and PRO-PMs will require improved integration of data collected from patients with data from EHRs and other health IT and data exchange systems that support quality measure calculation and public reporting. This can be achieved through the widespread use of interoperable data standards (e.g., Fast Healthcare Interoperability Resources [FHIR]) and standardized terminology systems (e.g., Logical Observation Identifiers Names and Codes [LOINC]).

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The U.S. is advancing data interoperability in multiple ways that will lower the burden associated with collecting and sharing data used for quality measurement. CMS recently released its dQM Strategic Roadmap ([executive summary](#) and [full report](#)), which sets forth how interoperability can be leveraged to advance and reduce the burden of quality measurement. The development and widespread support for FHIR have allowed it to emerge as the leading standard for interoperable healthcare data. FHIR is a Health Level Seven International (HL7) standard that defines how healthcare information can be standardized for exchange between different IT systems regardless of how it is stored in those systems. The Office of the National Coordinator for Health Information Technology (ONC) has also advanced the use of FHIR through policy. The 21st Century Cures Act Final Rule defined a minimum interoperable set of data classes and specific data elements: the United States Core Data for Interoperability (USCDI). The rule requires as part of the certification criteria that certain healthcare providers maintain access to the full scope of data defined in the USCDI Version 1 through standards-based application programming interfaces (APIs) for certain uses, such as providing patients access to their own data. These requirements will build upon the investments that providers and their vendors make to map their data to FHIR resources and make those data accessible in the FHIR standard.
ONC will expand the scope of available data over time through a USCDI Standards Version Advancement Process (SVAP). While the availability of standardized data relevant to PROMs and PRO-PMs will likely be minimal initially, this will evolve as the USCDI is updated and ONC adopts new versions for certification requirements. For example, Version 3 standards include a new class of data called Health Status/Aessment, which contain functional status data elements. These additions were supported by the quality measurement community seeking to advance digital PRO-PMs. In addition, ONC announced in 2021 that it is developing a set of standards to support a number of federal partners, including CMS and other agencies, implementing quality measurement and related use cases. This initiative will support the development of harmonized data sets for FHIR-based quality reporting. ONC is committed to engaging private as well as public stakeholders in this process.

ONC’s current structured process for updating USCDI and ONC, as well as other federal agencies’ work on USCDI+, creates opportunities for measure developers to advance interoperability for the measures they are developing by identifying and advocating for inclusion in both national core data sets and data elements as well as FHIR specifications. Information on how to submit data classes and elements for inclusion in future USCDI updates can be found in the USCDI ONC New Data Element and Class (ONDEC) Submission System.

In summary, advances in interoperability will ease data sharing, reduce clinical and administrative burden, create the infrastructure and specificity of standardized data and shared data formats needed to aggregate data across measured entities, and allow for the use of shared data in centralized measure score calculations. Measure developers must stay abreast of these advances as they design and implement digital PRO-PMs and can contribute to setting priorities for advancing the interoperable data needed for measurement.

Interim Report Recommendations
The Interim Report describes the attributes of high quality PROMs. The TEP and NQF recommend that measure developers use the Interim Report alongside the Roadmap. The goal of the Interim Report is to guide measure developers in selecting high quality PROMs that will be most conducive to the development and use of a digital PRO-PM. Key components of this report include the attributes of a high quality PROM, an Attribute Grid to facilitate analysis of PROMs, and prioritized PRO-PM domains.

Attributes of a High Quality PROM and the Attribute Grid
The Interim Report defines the attributes of high quality PROMs that are appropriate for use in a digital PRO-PM. These attributes are described and presented in the form of an Attribute Grid that measure developers can use to analyze and compare different PROMs that measure similar outcomes (Appendix D). The attributes identified in the Interim Report are as follows:

- Covers desired PROs from patient and/or caregiver perspective
- Outcome measured in PROM is the result of care for which relevant clinical quality is being measured
- Interpretable scores, defined and actionable cut points or targets, and anchors and/or defined meaningful change
- Clear conceptual and measurement models
- Psychometric soundness: Reliability
- Psychometric soundness: Validity
• Psychometric soundness: Responsiveness
• Usability/Feasibility of use: Low burden (e.g., length, time/effort to complete) and feasibility
• Usability/Feasibility of use: Fits with standard of care and related workflows (e.g., actionable, incorporated, and discussed at point of care)
• Usability/Feasibility of use: Cultural appropriateness, language, and translations with culturally appropriate items
• Usability/Feasibility of use: Availability of standardized clinical terminology and codes
• Usability/Feasibility of use: Guidance on standardized data collection (including modes and methods)

The Interim Report provides detail on the 12 attributes of a high quality PROM that is suitable for use in a digital PRO-PM. Some important themes within the attributes include identifying outcomes that matter to patients and/or caregivers and can be influenced by clinical care. High quality PROMs include well-documented scoring systems that indicate meaningful change (e.g., minimal clinically important difference [MCID]). A high quality PROM is also psychometrically sound, usable, and feasible in the clinical settings and populations that the instrument aims to address.

Although the TEP recognizes the attributes in the Interim Report as being important indicators of high quality PROMs, the Attribute Grid is not intended to be prescriptive. The Attribute Grid does not generate a score or a pass/fail determination, and the TEP opted against defining any “must-have” attributes. The Attribute Grid also allows measure developers to identify and assess additional attributes that are pertinent to the PRO-PM being developed.

Prioritized PRO-PM Domains

PRO-PMs can be grouped within five domains, or areas of patient-reported health status: HRQoL, functional status, symptoms and symptom burden, health behaviors, and patient experience. While all five domains represent important dimensions of performance, the Interim Report describes three primary drivers for the Roadmap’s focus on HRQoL, functional status, and symptoms and symptom burden:

1. **Representation of each domain in currently endorsed NQF measures**: More than one-third of NQF-endorsed PRO-PMs fall within the patient experience domain, while HRQoL and symptom-based PRO-PMs are underrepresented.

2. **Assessment of healthcare entity performance**: The Building a Roadmap initiative focuses on areas in which the clinical performance of healthcare entities most directly influences outcomes. These areas are most effectively captured by the domains of HRQoL, functional status, and symptoms and symptom burden. The patient behavior domain centers more on the actions and behaviors of patients than the clinical performance of the entities being assessed. The patient experience domain typically measures how well a healthcare entity serves patients (e.g., communication or ease of scheduling) by assessing the impact of processes of care on the patient’s experience.

3. **Data collection methodology and suitability for digital measurement**: PROMs within the patient experience domain, such as the Consumer Assessment of Healthcare Providers and Systems (CAHPS), utilize methodologies that significantly differ from other domains. To help prevent the possibility of retaliation against patients who report poor experiences with a provider or health system, patient experience measures typically depend on external partners to
confidentially collect and analyze patient-reported data using mixed modes and methods of data collection (e.g., self-administered mail and/or web questionnaires, with telephonic questionnaires for patients who do not respond). As a result of these methodological differences, the patient experience domain may not be as applicable to digital PRO-PMs as other domains.

This prioritization is not intended to diminish the importance of the patient behavior or patient experience domains, and many of the tasks identified in the Roadmap are applicable to these domains.

**Roadmap for the Development of a Digital PRO-PM**

**Overview and Visualization of the Roadmap**

The Roadmap provides guidance on the development of digital PRO-PMs, with a focus on those that are intended for NQF endorsement and suitable for use in CMS' VBP programs and APMs. The Roadmap is organized into four stages of development:

- Stage 1: Definition of Measurement Goals
- Stage 2: Exploration and Assessment of PROMs
- Stage 3: Development and Testing of the PRO-PM
- Stage 4: Finalization and Implementation of the PRO-PM

The visualization of the Roadmap (Figure 1) illustrates that each stage is broken into discrete tasks, with a total of 17 tasks that represent the entire digital PRO-PM development process.

*Figure 1. PRO-PM Roadmap: Each column contains one stage, and the bidirectional arrows indicate that tasks can move freely and be iterated across stages.*
In the same way that an actual roadmap offers many routes between two locations, the digital PRO-PM Roadmap recognizes that there are many routes to get from identifying the need for a measure to submitting a fully tested digital PRO-PM for NQF endorsement review.

The digital PRO-PM Roadmap recognizes that there are many routes to get from identifying the need for a measure to submitting a fully tested digital PRO-PM for NQF endorsement review. While the four stages generally occur in the sequence listed above, the arrows in the visualization indicate that the 17 tasks are not entirely linear and are not bound to any specific stage. Different measure developers and organizations may follow different processes when developing digital PRO-PMs; therefore, the Roadmap is designed to allow tasks to move across stages based on individual and organizational style, preference, and need. For example, attribution is addressed in the stage 3 task titled “Specify Key Measure Details”; however, measure developers on the TEP expressed different preferences on when this task should occur, with some designing the attribution model as part of the preliminary work that occurs during stage 1.

Because these variations occur across different people and organizations, the Roadmap is not prescriptive as to when each task must be performed, and it acknowledges that significant flexibility exists within these tasks. However, every task is an important part of the digital PRO-PM development process that should be planned proactively and addressed by the measure developer in order to avoid unnecessary recursive activities. The measure developer should also ensure considerations of health equity have been included in every task. If a task is not relevant to the PRO-PM being developed, the measure developer should thoroughly and transparently explain the rationale within the measure’s documentation.

In addition to the Roadmap’s flexibility, the TEP recognizes that many tasks related to digital PRO-PM development are iterative and may repeat across multiple stages. For example, testing a digital PRO-PM is an expensive and complex process that occurs in stage 3, but developers should begin planning for this task during the earliest stages and expand upon it regularly throughout the life of the project. The stages and tasks are described in detail in the Roadmap, and this high-level summary may assist readers in understanding what to expect during each stage.

- **Stage 1: Definition of Measurement Goals.** The measure developer becomes familiar with key reference documents, then assembles a stakeholder advisory group that is composed of multiple expert perspectives, including patients, patient advocacy groups, caregivers, and/or consumer groups. The stakeholder advisory group helps to identify the measure’s outcomes and audience, ensure the measure is novel, evaluate its intent and intended use, and ascertain that a PRO-PM is the proper measurement approach. The tasks within stage 1 establish a clear plan for the entire measure development life cycle. Execution of the tasks within stage 1 can influence the success of the entire digital PRO-PM development life cycle.

- **Stage 2: Exploration and Assessment of PROMs.** This stage of the Roadmap includes the identification, assessment, and selection of the PROM(s) that will collect data for the PRO-PM.
The Interim Report describes, in detail, the process and criteria for assessing the attributes of a PROM and determining whether it is appropriate for use in a PRO-PM. The Interim Report, along with the Attribute Grid it contains, can help a measure developer select high quality PROM(s) for use with the performance measure. The Interim Report is designed to be used during stage 2 of the Roadmap.

- **Stage 3: Development and Testing of the PRO-PM.** The third stage of the Roadmap addresses the specification of details about the measure, including risk adjustment and attribution; the selection of test sites; and the creation and execution of testing plans, including alpha and beta testing. Depending on the measure developer’s experience, preference, or organizational policies, the tasks within this stage may vary. However, every task in stage 3 must be considered within the development process. If the measure developer and the stakeholder advisory group determine that a detail is not appropriate for the PRO-PM specification (e.g., risk adjustment), the measure developer should thoroughly document the rationale for these decisions for future reference.

- **Stage 4: Finalization and Implementation of the PRO-PM.** During this stage, the measure developer creates the implementation documentation and prepares for NQF’s CDP depending on whether the PRO-PM will be submitted for NQF endorsement review. The preparation for tasks in stage 4 will likely begin early in the development process, even though many of these tasks are not completed until late in the process. Careful planning and preparation for these tasks will improve the likelihood of successfully testing the PRO-PM and submitting it for NQF endorsement review.

It is critical for readers of this report to understand that the Roadmap is a guide for development, not a guarantee of NQF measure endorsement. Many factors influence whether a measure is endorsed by NQF. The authors of this report encourage measure developers to carefully review and follow the official NQF endorsement processes.

**Health Equity**

Health equity is a critical element of measurement and is prioritized among both public and private stakeholders. Assessments of interventions to reduce health disparities are an important step to achieving health equity, and PRO-PMs pose a unique patient-centered measurement opportunity. As measure developers engage in the development process, they should continuously and iteratively consider how equity is integrated in the measure.

At each stage and task in the Roadmap, the measure developer should refer to health equity resources for best practices to ensure the highest level of inclusion (Appendix B, e.g., NQF’s Best Practices for Developing and Testing Risk Adjustment Models and the CMS Framework for Health Equity). Measure developers should actively aim to prevent internal biases and exclusion of populations from data by planning and managing equity considerations through stage 1 tasks. The development of PRO-PMs stems from the data and data collection practices of PROMs; therefore, developers should plan to review these practices for equity during stage 2 tasks. For stage 3 tasks, developers should engage in an iterative process to ensure plans for equity made during stage 1 are accomplished. Stage 3 tasks also present developers with the opportunity to ensure the digital PRO-PM aligns with current best practices for risk adjustment or stratification as well as equitable testing. To help support stage 4 (i.e., preparing
for implementation of the PRO-PM and submission for NQF endorsement), the measure developer should document how the measure addresses health equity.

Stage 1: Definition of Measurement Goals

As with any major project, early planning and preparation can improve the entire measure development life cycle. A measure developer should complete these tasks before beginning the development of a digital PRO-PM. This early work helps to ensure the measure will assist in improving the desired outcomes. While the measure developer may decide to move other tasks into stage 1, particularly those that will benefit from early planning or iterations, the six tasks listed below should be completed before moving beyond stage 1.

Review Relevant Resources

The TEP designed the Roadmap as a high-level guide that measure developers can use alongside source-of-truth resources on quality measures, PRO-PMs, equity, and interoperability. Developers who are working on digital PRO-PMs should familiarize themselves with these resources, which are developed and maintained by organizations with expertise on specific aspects of measure development. Developers should reference these resources throughout the four stages of digital PRO-PM development. The Roadmap contains a comprehensive list of recommended resources to aid in the development of digital PRO-PMs (Appendix B).

Establish Stakeholder Advisory Group and Feedback Processes

The creation of the stakeholder advisory group is a critical early step in developing PRO-PMs because it ensures key perspectives are represented throughout the measure development life cycle. The stakeholder advisory group offers expert feedback and guidance to the measure developer. A responsibility of the stakeholder advisory group is to ensure that the new measure will provide value to a broad range of stakeholders.

Although the Roadmap generally refers to a single measure advisory group, measure developers can choose to have multiple stakeholder advisory groups, depending on the feedback needed. The measure developer can also identify specialized subgroups within a stakeholder advisory group. A single stakeholder advisory group allows the measure developer to gather diverse viewpoints and seek consensus on complex issues, while multiple groups or subgroups allow the measure developer to gain targeted feedback. In situations where multiple stakeholder advisory groups are appropriate, the measure developer should consider scheduling a mixture of separate and combined advisory group meetings to maximize opportunities for transparency, diverse feedback, and consensus.

Measure developers might consider creating a patient-focused subgroup to address issues that are pertinent to patients, families, and caregivers. A patient-focused subgroup likely spends less time on technical facets of measure development and more time helping the measure developer ensure the measure is meaningful and valuable to patients. Tasks for which measure developers might engage a patient-focused subgroup include the identification of outcomes and audiences (stage 1, task 3), the exploration and assessment of PROMs (stage 2), and the selection of test sites (stage 3, task 3). If the measure developer creates a patient subgroup, it should regularly engage with the full stakeholder advisory group.
The stakeholder advisory group must include representation from patients, patient advocacy groups, caregivers, and/or consumer groups, as appropriate to the measurement goals: These are the people whose perspectives drive the performance measure.

The composition of the stakeholder advisory group will depend on the measurement goal. The measure developer should first identify which perspectives are mandatory and which are optional, then select individuals to represent these perspectives. The stakeholder advisory group must include representation from patients, patient advocacy groups, caregivers, and/or consumer groups, as appropriate to the measurement goals: These are the people whose perspectives drive the performance measure. Depending upon the PRO-PM’s measurement goals, a balanced stakeholder advisory group might also include clinicians, administrative staff (e.g., staff who will administer or assist with PROM completion), payers, health IT staff, EHR and/or other pertinent software vendors, data scientists, policy experts who represent the entities that will be affected by the measure, public program administrators, and other stakeholders who will be directly affected by the PROM completion and collection process and/or the PRO-PM. Measure developers should consider including PROM developer(s) in the stakeholder advisory group, although this might not be possible until PROM(s) are selected during stage 2.

Health equity is the attainment of the highest level of health for all people, and it warrants consideration throughout the measure development process. This consideration begins with the stakeholder advisory group. During this task, measure developers can determine other stakeholder perspectives that are necessary to add to the group to ensure the PRO-PM is equitable. Depending on the measure’s outcomes and audience, members of the stakeholder advisory group could include patients and/or clinicians from rural settings, people in the measure’s target population who speak English as an additional language, people who have physical or cognitive disabilities, patients and/or clinicians with conditions or diagnoses targeted by the measure (e.g., people with depression or who had hip replacement surgery), and other representatives from vulnerable or underserved populations. This task is an opportunity for the measure developer to ensure the inclusion of typically underrepresented voices and integrate a practice of equity throughout the development process.

The measure developer should clearly define processes to ensure stakeholders can submit feedback in a timely manner throughout the development life cycle and that other interested parties can provide input at key milestones (e.g., public commenting periods). The type and complexity of the proposed measure will influence the involvement of the stakeholder advisory group, with complex measures requiring more input.

Identify Outcomes and Audience

The measure developer, with guidance from the stakeholder advisory group, should identify a consensus definition for the measure concept (i.e., a high-level recommendation that includes a clear description of what should be measured) and the measure intent (i.e., what the measure aims to achieve or
improve). The measure developer should identify the desired outcomes that will be monitored, such as functional status after a procedure, evaluation of symptom remission, or assessment of pain control throughout the care continuum. During this task, the measure developer should also identify the primary audience of users who will be assessed by the measure (i.e., the measured entities). The measure developer should conduct an environmental scan and use empirical data, literature, and/or guidelines to determine and support the measure concept and target population.

Measure developers need to provide supporting evidence that the PRO within the PRO-PM is meaningful to the target population of patients and/or caregivers. The stakeholder advisory group and any patient-focused subgroups might participate in these activities, and the measure developer may also choose to engage members of the target population through surveys, focus groups, or other information-gathering activities. These activities help the measure developer to demonstrate that the measure concept is meaningful to patients.

Confim Concept Fills a Measurement Gap

A new PRO-PM must address a measurement gap. It might measure an important issue with significant variation that is not currently measured, or it might be an improvement to an existing measure. A measure developer can determine whether a measurement gap exists by scanning existing measure databases (e.g., NQF’s Quality Positioning System [QPS]; the CMS Measure Inventory Tool [CMIT]; and CMS’ Pre-Rulemaking webpage, which includes the Measures Under Consideration [MUC] list) to identify and assess any similar measures and ensure the definition, outcome, and audience are either new or will significantly improve upon existing measures. By conducting an environmental scan of measures, the measure developer can avoid duplicating existing work and adding unnecessary burden on measured entities. If a similar measure(s) already exists, the measure developer can consult with the stakeholder advisory group as well as the steward of the existing measure to determine whether the concept justifies another measure.

Determine the Intended Use of the Measure

Different measures serve different purposes. Some performance measures are used solely for quality improvement, while others might be used for accreditation, comparison of provider performance, or payment in APMs or VBP programs. Due to the time, effort, and cost required for developing a digital PRO-PM, the measure developer and the stakeholder advisory group should determine how the measure will be used before moving beyond stage 1. Development and testing processes will vary depending on this determination, and requirements are more stringent for accountability than for internal quality improvement use cases.

Assess the Best Type of Quality Measure

The measure developer will need to determine whether the measure concept can be assessed with information gathered directly from patients. For example, if the measure assesses how frequently a process was completed or focuses on certain types of structured clinical data (e.g., a lab value, blood pressure result, or weight), a PRO-PM is most likely not appropriate. However, if the measure concept is based on PROs, assessed through data collected via a PROM, and aggregated for an accountable healthcare entity, then a PRO-PM may be the ideal type of performance measure.
Developing PRO-PMs, and specifically digital PRO-PMs, can be complex, time consuming, and expensive. Once the measure developer has determined that the measure concept should proceed as a digital PRO-PM, reviewing the full Roadmap and planning for several tasks in stages 2, 3, and 4 can help mitigate challenges and delays later in the development process. Some important questions to begin considering now include the following:

- What health equity issues affect the target population (e.g., language requirements and access to broadband)? (stages 2 and 3)
- Will data collection be limited to a single PROM, or can different PROMs collect data for the PRO-PM? (stage 2, task 3)
- What is the measure’s attribution model? (stage 3, task 1)
- How will the measure be risk-adjusted or stratified? (stage 3, task 1)
- What sampling methodology might ensure a representative sample, and what strategies might lessen the chances of a low response rate or measurement bias? (stage 3, task 1)
- What is the potential testing plan? (stage 3, task 2)
- Which test sites might be appropriate for alpha and beta testing? (stage 3, tasks 3 and 4)
- When will the measure developer have adequate information to complete the Intent to Submit process? (stage 4, task 2)

Additionally, as the measure developer prepares to assess the test results of PROMs and plan for the testing of the PRO-PM, the developer needs to understand the differences between PROM testing and PRO-PM testing. Table 2 offers a high-level description of these differences.

**Table 2: Distinctions in Testing PROMs and PRO-PMs**

<table>
<thead>
<tr>
<th>Distinction</th>
<th>PROM</th>
<th>PRO-PM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Who is responsible for testing?</td>
<td>The PROM developer is responsible for testing a PROM</td>
<td>The measure developer is responsible for testing a PRO-PM</td>
</tr>
<tr>
<td>What are some key testing components?</td>
<td>Psychometric soundness, meaningfulness, usability, and feasibility</td>
<td>Importance to measure and report, scientific acceptability</td>
</tr>
<tr>
<td>Where is this discussed in the report?</td>
<td>PROM testing is discussed further in stage 2</td>
<td>PRO-PM testing is discussed further in stage 3</td>
</tr>
<tr>
<td>What additional resources exist?</td>
<td>• The 2013 NQF Committee Final Report titled Patient-Reported Outcomes in Performance Measurement</td>
<td>• NQF Measure Evaluation Criteria webpage</td>
</tr>
<tr>
<td></td>
<td>• The 2015 report written by David Cella et al, also titled Patient-Reported Outcomes in Performance Measurement</td>
<td>• NQF Measure Evaluation Criteria and Guidance for Evaluating Measures for Endorsement (PDF, September 2021)</td>
</tr>
<tr>
<td></td>
<td>• The 2021 Building a Roadmap Interim Report on high quality PROMs for use in PRO-PMs</td>
<td>• The CMS Blueprint Measure Lifecycle Overview webpage</td>
</tr>
<tr>
<td></td>
<td>• The CMS Blueprint QuickStart Guide (PDF, May 2022)</td>
<td>• The CMS Blueprint QuickStart Guide (PDF, May 2022)</td>
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</table>
Stage 2: Exploration and Assessment of PROMs

Once the initial strategic work is completed, the measure developer can begin an objective assessment and selection process to determine which PROM(s) are the most appropriate data collection tools for the PRO-PM. Hundreds of PROMs already exist, including instruments that are specific to a disease or condition as well as those that are disease-agnostic and can be used across a broad population of patients. The Roadmap focuses its guidance on measure developers who are selecting from existing PROMs rather than those who are working with PROM developers to create a de novo PROM for use with a PRO-PM, although the attributes of a high quality PROM are applicable to either scenario.

Both the Interim Report and Roadmap are designed to be used together, and the Interim Report provides more extensive detail on the tasks that occur during stage 2 of the Roadmap.

**Identify Candidate PROMs**

The measure developer must first identify at least one PROM that is a candidate for use in the digital PRO-PM. When identifying candidate PROMs, measure developers should keep stage 1 findings top of mind. For example, if the stage 1 assessments determined that the PROM must have a validated computerized adaptive testing (CAT) version available in both English and Spanish in order to reduce burden and inequities for the target population, the measure developer can immediately disregard PROMs that do not meet those initial requirements.

The measure developer can consult several resources to identify candidate PROMs that might be suitable data collection tools for the desired PRO-PM:

- The International Consortium for Health Outcomes (ICHOM) sets of Patient-Centered Outcome Measures provide lists of recommended PROMs that have been vetted by diverse expert panels.
- The Consensus-Based Standards for the Selection of Health Measurement Instruments (COSMIN) website contains extensive resources on identifying candidate PROMs.
- The Mapi Research Trust PROQOLID database and catalog of clinical outcome assessment (COA) instruments
- Many professional societies have convened working groups of experts to review and recommend PROMs that are relevant within that specialty or discipline.
- Articles from peer-reviewed journals can provide insights into which PROMs are most widely used within a discipline and which novel PROMs are offering noteworthy contributions.
- The descriptions or specifications of PRO-PMs typically list which PROM(s) are used to collect data for the PRO-PM; websites such as QPS and CMIT include these details if they are available.
- Some payment models in the CMS Quality Payment Program (QPP), such as APMs, identify required PROMs.

**Assess PROMs Using the Attribute Grid and Ensure Sufficient Testing Has Occurred**

Once the measure developer has identified candidate PROMs, the Attribute Grid (Appendix D; described in detail in the Interim Report) can help to compare PROMs and determine which are most suitable for use in the digital PRO-PM. The Attribute Grid provides a structure for performing a side-by-side comparison of the benefits and limitations of each PROM by prompting the measure developer to
review the literature and gather information that indicates whether a PROM is high quality for use in a digital PRO-PM (i.e., its ability to collect data—including patient responses, scores, or threshold data—that are appropriate for use in a digital PRO-PM that is suitable for inclusion in payer assessments, such as one of CMS’ VBP programs or APMs).

Patients, caregivers, advocates, and/or consumers from the stakeholder advisory group will be important participants in assessing the first attribute (i.e., whether the PROM addresses the desired outcomes from the patient and/or caregiver perspective), while clinician advisors should be closely engaged when assessing the second attribute. However, each of the 12 attributes are important to consider, and the stakeholder advisory group should be engaged during this process.

The attributes of a high quality PROM that are well suited for use in a digital PRO-PM are as follows:

- Covers desired PROs from patient and/or caregiver perspective
- Outcome measured in PROM is the result of care for which relevant clinical quality is being measured
- Interpretable scores, defined and actionable cut points or targets, and anchors and/or defined meaningful change
- Clear conceptual and measurement models
- Psychometric soundness: Reliability
- Psychometric soundness: Validity
- Psychometric soundness: Responsiveness
- Usability/Feasibility of use: Low burden (e.g., length, time/effort to complete) and feasibility
- Usability/Feasibility of use: Fits with standard of care and related workflows (e.g., actionable; incorporated and discussed at point of care)
- Usability/Feasibility of use: Cultural appropriateness, language, and translations with culturally appropriate items
- Usability/Feasibility of use: Availability of standardized clinical terminology and codes
- Usability/Feasibility of use: Guidance on standardized data collection (including modes and methods)

The TEP strongly encourages the measure developer to consider each of the 12 attributes in the Attribute Grid when selecting a PROM. However, the TEP opted against utilizing a scoring mechanism or identifying any must-have attributes, as these could vary depending on the specific need and intent of the PROM. As such, the measure developer has the flexibility to prioritize the 12 attributes based on the needs of the PRO-PM that is being developed. The measure developer may also identify additional attributes that are relevant to the performance measure and add new rows to accommodate these attributes. Health equity considerations are addressed across multiple attributes (e.g., covers desired PROs from the patient perspective and cultural appropriateness), and measure developers should utilize the Attribute Grid to assess those considerations.

The TEP strongly encourages the measure developer to consider each of the 12 attributes in the Attribute Grid when selecting a PROM.
The Interim Report does not explicitly address the use of proxy responses (i.e., one person completing a PROM on behalf of another person, such as a caregiver completing a questionnaire on behalf of a person with cognitive disabilities or a parent completing a PROM for their child), though it does highlight the importance of guidance on standardized data collection practices. When assessing a PROM for use in a digital PRO-PM, the measure developer needs to examine the PROM developer’s guidance on capturing proxy responses and incorporating them alongside data captured directly from the patient. If the existing guidance does not align with the planned use of the PRO-PM, the measure developer should work with the PROM developer to ensure the approach to proxy responses aligns with the intent of the PRO-PM. When proxy responses are not sufficiently addressed, issues of equity and scientific acceptability can arise.

Some of the attributes will have objective, quantifiable results, and others will be subjective. For example, the attribute of “Psychometric Soundness: Reliability” should always include a numeric result. While a measure developer can quantifiably measure the number of languages in which a PROM is translated, the cultural appropriateness of those translations may contain more subjectivity. However, PROM developers have access to well-developed processes for establishing the linguistic and cultural appropriateness of each PROM translation, including pilot testing/cognitive debriefing with patients and clinicians, so that even potentially subjective results should be assessed against established scientific methods. The subjective attributes allow the measure developer to think holistically about a performance measure and consider the perspectives of multiple stakeholders, particularly patients and caregivers, when assessing PROMs. The quantifiable attributes are more structured, and existing literature can provide guidance in interpreting the results. For example, the 2012 NQF Methodological Issues Report recommends that PROMs should have a reliability estimate of greater than or equal to 0.70 for group-level purposes and greater than or equal to 0.90 for individual-level purposes. However, the Attribute Grid itself does not specify which results should be considered acceptable. The measure developer should utilize professional judgement and expertise in selecting PROMs while analyzing both the quantifiable and subjective attributes, knowing that the pool of instruments and literature is constantly expanding. The measure developer should refer to existing tools and recommendations when selecting PROMs, as well as previous NQF reports.

Many elements of testing—including psychometric soundness and mode of administration—are reviewed using the Attribute Grid. However, the NQF endorsement process for a digital PRO-PM could require a PROM to undergo additional testing. For example, endorsement of a PRO-PM that is also an eCQM will require validity testing for each question of a PROM, and the endorsement process will scrutinize any substantial differences in validity across questions. As a result, the measure developer should examine additional evidence supporting the PROM’s suitability for use in the PRO-PM, such as the determination of whether the PROM was tested with the population in the setting (e.g., inpatient surgery, ambulatory surgery center) that the PRO-PM will be implemented. Because PRO-PM developers typically do not have the resources to independently test the reliability and validity of each PROM, developers should consider seeking PROMs in which the tests of psychometric soundness were performed in settings and with populations that align with the entities being measured by the PRO-PM. If the original testing of the PROM does not match the current intended use, the PRO-PM developer should contact the PROM developer to determine whether additional information is available. The
The interdependent relationship between PROMs and PRO-PMs is why the measure developer should attempt to engage the PROM developer as a member of the stakeholder advisory group.

Because PRO-PM developers typically do not have the resources to independently test the reliability and validity of each PROM, developers should consider seeking PROMs in which the tests of psychometric soundness were performed in settings and with populations that align with the entities being measured by the PRO-PM.

Select PROM(s) for Inclusion in PRO-PM

Based on the analysis of the Attribute Grid, the measure developer should determine which PROM(s) best meet the needs of the PRO-PM. During this task, the measure developer and stakeholder advisory group will determine whether the PRO-PM will utilize data from a single PROM or from multiple different PROMs. There are advantages and disadvantages to either approach:

- **A PRO-PM that utilizes data from a single PROM:**
  - The measure developer can select one PROM that is singularly well suited to collect data for the PRO-PM (e.g., a PROM with multiple validated translations).
  - The measure developer can tailor the PRO-PM specification to the data structure, scoring, and cut points (i.e., markers that indicate clinically meaningful change) of a single PROM, potentially resulting in a more straightforward development process.
  - The measure steward’s maintenance of a single-PROM performance measure is typically less burdensome.

- **A PRO-PM that can utilize data from multiple, different PROMs:**
  - Clinical settings have increased autonomy to implement the PROM that best meets their population and business needs (e.g., PROMs that are culturally sensitive and translated into languages that are most relevant to the patient population or PROMs that are free versus those with a licensing cost).
  - The measure developer has the flexibility to create a PRO-PM that collects data from different instruments that measure the same underlying domain (e.g., HRQoL, symptoms and symptom burden, or health functioning); however, a reliable and valid crosswalk must exist or be developed to ensure the different PROMs consistently assess outcomes (e.g., harmonized cut points and scoring).

Stage 3: Development and Testing of the PRO-PM

The measure developer can begin stage 3, “Development and Testing of the PRO-PM,” after assessing and selecting the PROM(s) and confirming that defined instrument parameters and psychometric properties are sufficient for the proposed measurement concept. Measure development involves designing a quality measure that is suitable for assessing the performance of the accountable entity; it
includes defining the care targeted for measurement, the level and type of provider being evaluated, the approach to attributing patients to measured entities, and any model development required to adjust for difference in case mix across measured entities. Measure testing involves formally evaluating the measure’s reliability and validity for use as a quality measure.

Some developers and organizations will prefer to address certain tasks (e.g., attribution or risk adjustment) at different points in the development life cycle. Regardless of at which point the measure developer initiates these tasks, each must be addressed and iteratively updated as new information is gained throughout the development process. If the measure developer and the stakeholder advisory group determine that certain tasks are unnecessary for a specific PRO-PM, the measure developer should thoroughly document the rationale for these decisions in the PRO-PM’s supporting documentation.

**Specify Key Measure Details**

The technical specification for a PRO-PM identifies the information and data that are needed to generate the measure result. This includes the population (i.e., the patients whose PROM data will be aggregated for the measure), the inclusion and exclusion criteria, and the data sources (e.g., PROMs and claims data). Measure specifications provide technical instructions on how the measure should be collected and used consistently, reliably, and effectively. Developing measure specifications is an iterative process that requires a variety of stakeholder perspectives. The measure developer should engage the stakeholder advisory group in creating and refining the measure specifications and may consider conducting public commenting once the initial draft of the measure specifications is complete in order to acquire additional feedback before investing in the testing process.

Elements that are typically part of the technical specifications include the following:

- Measure name/title
- Description
- Population
- Numerator and denominator statements
- Inclusion and exclusion criteria and exceptions, including rationale as appropriate
- Data sources (e.g., PROMs, registries, and claims data)
- Key terms
- Data elements, codes, and code systems
- Unit of measurement of analysis
- Sampling methodology
- Attribution model
- Risk adjustment
- Time intervals
- The calculation algorithm

**Specify Information for Related PROMs**

Technical specifications for PRO-PMs require information about the PROM(s) that are used as data collection instruments. The measure developer will access some of this information during the stage 2 tasks, but other details may be needed, such as acceptable modes of PROM administration, instructions for recording PROM results that are blank or contain multiple responses to a single question, and the
Although the following examples are not PRO-PMs, they present a side-by-side comparison of a digital (i.e., machine-readable) specification and a narrative (i.e., human-readable) specification for a quality measure. The examples compare the “Initial Patient Population” sections of CMS #146 Appropriate Testing for Children With Pharyngitis. Links are provided to specifications on the HL7 CQL website and the CMS Quality Payment Program website.

**Machine-readable format:**

```
* Initial Patient Population =
  * AND: "Patient Characteristic: Birthdate: birth date" ≥ 2 year(s) starts before start of "Measurement Period"
  * AND: "Patient Characteristic: Birthdate: birth date" < 18 year(s) starts before start of "Measurement Period"
  * AND:
    * AND: "Occurrence A of Encounter, Performed: Ambulatory/ED Visit" during "Measurement Period"
    * AND: "Medication, Order: Antibiotic Medications" starts after start of "Occurrence A of Encounter, Performed: Ambulatory/ED Visit"
    * AND:
      * OR: "Occurrence A of Encounter, Performed: Ambulatory/ED Visit" during
        * OR: "Occurrence A of Diagnosis, Active: Acute Pharyngitis"
        * OR: "Occurrence A of Diagnosis, Active: Acute Tonsillitis"
      * OR:
        * OR: "Occurrence A of Diagnosis, Active: Acute Tonsillitis"
        * starts during "Occurrence A of Encounter, Performed: Ambulatory/ED Visit"
```

**Human-readable format:**

Initial Population: children 3-18 years of age who had an outpatient or emergency department (ED) visit with a diagnosis of pharyngitis during the measurement period and an antibiotic ordered on or three days after the visit.

As described in the section on interoperability, FHIR will facilitate the use of dQMs by advancing data and specification standards. To date, however, eCQMs are primarily developed and reported using the QDM rather than FHIR. CMS and others are working to streamline the specification process and offer enhanced tools for measure specification development and testing. NQF is considering its role in developing standards for evaluating FHIR-based measures. Measure developers should align their efforts with current NQF, CMS, and/or ONC guidelines (e.g., the dQM Strategic Roadmap).
Determine Appropriate Attribution Model

The measure developer must answer an important question during the digital PRO-PM development process: Which provider types and organizations are responsible for the patient outcomes? Attribution is used in quality measurement to assign accountability for a patient’s outcomes to the accountable entity being assessed by the measure; in other words, the model assigns specific patients to the specific parties being measured (e.g. a clinician, group of clinicians, facility, or accountable care organizations [ACOs] and health plans). An attribution model is a set of rules that defines the accountable unit for a patient’s healthcare outcomes. It may utilize, for example, a patient’s attestation of who their provider is or analyses of claims data. Attribution models are needed for PRO-PMs because patients may acquire healthcare related to the measured outcome from multiple providers serving various payers; an attribution model specifies how patients will be assigned to providers when the measure is implemented to ensure that providers’ measure scores will reliably reflect patients whom the measured providers cared for and outcomes the providers had the ability to influence. NQF’s Attribution: Principles and Approaches Report includes the Attribution Model Selection Guide, a useful resource for determining the appropriate model selection. The NQF report on Attribution for Critical Illness and Injury offers additional guidance on geographic/population-based attribution models.

During this task, measure developers should engage the stakeholder advisory group to discuss effects on accountable entities. This will help clarify the connection between the PRO-PM data and the accountable entity’s performance.

Develop and Test Risk Adjustment Model

For outcome measures, including PRO-PMs, NQF endorsement requires either an evidence-based explanation of why risk adjustment is not required or an evidence-based risk adjustment strategy to account for differences in case mix across measured entities that affect the risk of the outcome independent of quality. Patients may have demographic, clinical, and social risk factors that vary widely across the measured entities and need to be considered for risk adjustment in model development. For the risk model, developers must provide a demonstrated conceptual and statistical rationale that considers patient factors outside of the provider’s control, which influence the measured outcome and are present at the start of care. Differences in individuals’ responses related to instruments or methods, modes, and languages of administration need to be analyzed and potentially included in the risk adjustment model. If risk adjustment is not appropriate for the PRO-PM or unrealistic within the development timeline, the measure developer must document and provide rationale and evidence to support the lack of risk adjustment and/or provide the intended plan for future risk adjustment as more PRO-PM data become available after implementation and use in the field.

Guidance on risk adjustment modeling is evolving. Risk-adjusting performance measures, namely outcome and cost/resources use measures, has traditionally accounted for differences in patient health status and clinical factors (e.g., comorbidities or severity of illness) that are present at the start of care. This approach to risk adjustment has been widely accepted and implemented within measure development. However, patients can also bring certain social characteristics (e.g., income, education, housing instability, food insecurity, and urbanicity/rurality) into their engagement with the healthcare system, which can influence healthcare outcomes and are outside the provider’s control. The idea of incorporating these social factors into risk adjustment models for quality measurement has been debated, in brief, due to competing concerns: On one hand, not adjusting may be unfair to providers caring for more patients with these risk factors, while on the other hand, adjusting could lead to
potential unintended consequences, such as obscuring differences in quality among providers and institutionalizing differing health outcome expectations for different patient groups.

Stratification of measure scores can be used to enhance transparency by highlighting known areas where disparities exist, or a need is present to identify differences in performance measure results. Stratification refers to the division of a population into distinct, mutually exclusive strata or groups defined by particular patient characteristics represented in data, thus enabling analysis of the specific subgroups. Stratification can be a tool for reporting on within-entity differences in measure results between subgroups of patients. It can also be used to compare measure scores calculated for subgroups of patients across accountable entities, which is referred to as between-entity differences. Additionally, stratification can refer to segmenting accountable entities rather than patients, which is known as “peer grouping.”

NQF is publishing updated recommendations and analyses of best practices for risk adjustment models, which includes recommendations for stratification, in late 2022 on the Risk Adjustment Guidance project page.

**Sampling Methodology**

The sampling methodology describes the approach to identifying a representative sample of accountable entities that will be measured by the PRO-PM and quantifying the implications of nonresponse rates observed in testing. Although nonresponse is a consideration for PRO-PMs that does not apply to most other types of performance measures (e.g., measures based on laboratory values are not dependent on a patient voluntarily completing a questionnaire), measure developers must assess nonresponse rates for PRO-PMs because they can bias the measure score. These concerns can be mitigated with thoughtful planning of a sampling strategy for the PRO-PM with the goal of establishing a minimal sample size that supports a reliable measure score. The sampling strategy should identify adequate numbers, approaches for randomization, and be representative of the intended patient population. NQF does not provide prescriptive rules for identifying a sampling methodology; therefore, the measure developer needs to carefully consider and prepare to illustrate how risks of bias are addressed.

**Create Testing Plan(s)**

Comprehensive and accurate testing plans that closely follow NQF’s measure evaluation criteria are an important aspect of developing measures. To ensure the practical and logistical success of the digital PRO-PM, the test plan should identify and address the goals of the testing process. The testing plan should identify data requirements, including preliminary strategies for addressing issues of data availability, data accuracy, insufficient sample sizes, or other common challenges.

Testing plans can vary depending on measure type and the complexity of the measure. Separate test plans are created for alpha testing and beta testing. Alpha testing, or formative testing, determines the feasibility of testing and implementing the draft specifications; subsequently, this testing plan is usually prepared early in the measure development life cycle. Beta testing, or field testing, expands feasibility testing and also assesses scientific acceptability and usability. A testing plan for beta testing is usually created later in the project since beta testing should not occur until the measure specifications are almost final. Testing plans for beta testing generally include the following:
Select Test Sites and Prepare for Testing

Beta testing is a resource-intensive and expensive process that is critical to the success of the digital PRO-PM. Although beta testing is listed as a stage 3 task, the measure developer should plan for and coordinate beta testing of the digital PRO-PM early in the development process.

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Because PRO-PMs rely on data that are voluntarily provided by patients, selecting PRO-PM test sites carries unique challenges. Test sites are hesitant to accept the burden of testing PRO-PMs that might not be endorsed or adopted. Measure developers involved with the Building a Roadmap initiative identified this as one of the most significant challenges that faces PRO-PM developers. Unlike testing a new performance measure that depends on information that is routinely collected during clinical care, collecting PROs may require new clinical workflows, staff training, patient education, and/or IT infrastructure. Strategic approaches to this challenge do exist nonetheless:

Identify candidate test sites that are already using the PROM(s): Once the measure developer completes the stage 2 task of identifying PROM(s) for data collection, begin surveying potential test sites to identify who is already using the PROM as part of their established clinical workflows. By focusing on test sites that are already using the PROM, the burdens of implementing a new PROM are significantly reduced, if not eliminated. Measure developers and PROM developers can collaborate in this effort, which reinforces the benefits of including PROM developers on the stakeholder advisory group. Journal articles about the PROM can also help measure developers identify individuals and clinics who have already adopted a PROM. Notably, digital PRO-PMs must be tested at a minimum of two sites with at
least two different EHRs. If the measure developer is only able to identify one suitable test site, other strategies in this section might apply.

**Collaborate with partners to incentivize test sites:** Some stakeholders in the measure development process, such as federal agencies, EHR vendors, private companies, or foundations, can be well positioned to develop unique methods of incentivization for test sites. As an example, an EHR vendor who is seeking a competitive advantage in the market might be willing to incur costs related to building a digital PROM in the EHR. Federal agencies, such as CMS, can be uniquely positioned to incentivize test sites that participate in VBP programs, APMs, or other innovative payment programs. Private companies or foundations may be able to provide funding to offset the costs of training staff and implement clinical workflows that support the PROM. While stakeholders need to be aware of and avoid potential conflicts of interest, incentives can help alleviate the burden that test sites face.

**Explore creative solutions to reduce the burden of implementation:** Measure developers are well positioned to find solutions that will lower barriers resulting from training staff, educating patients, and developing IT infrastructure for a new PROM. As an example, a measure developer could offset the burden of building a PROM into the test site’s EHR by designing a spreadsheet-based method to analyze and aggregate PROM data and providing staff to capture and manage the data. A different approach might be to utilize PROM data from existing registries under a data use agreement (DUA) or business associates agreement (BAA). A third approach could be to engage a vendor who administers PROMs and stores patient responses without affecting the test site’s workflows or data systems. While creative solutions such as these might not meet the testing requirements of endorsing a digital PRO-PM, they might enable the measure developer to apply for trial use under NQF’s trial use criteria for eCQMs, as described below.

**Consider NQF Trial Use endorsement of a digital PRO-PM:** The NQF Trial Use program recognizes the testing burden of eCQMs that are ready for implementation but cannot be adequately tested to meet NQF endorsement criteria. A digital PRO-PM with Trial Use Approval receives a three-year period for additional testing. While not every eCQM will proceed to endorsement (some fail due to the results of testing during the Trial Use period), it is an important program that can offset the burden of testing digital PRO-PMs.

Patients are valuable partners in this task. Their insights and experiences are critical on topics related to workflows regarding PROM administration that affect testing. For example, patients who are part of the target population are more likely than clinicians to understand issues related to social determinants of health and health equity (e.g., barriers that prevent patients from completing PROMs digitally via patient portals or smart phones) or convenience (e.g., challenges with arriving early for appointments to fill out PROMs in the waiting room).

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**Patients are valuable partners in this task. Their insights and experiences are critical on topics related to workflows regarding PROM administration that affect testing.**
**Conduct Alpha Testing**

Alpha testing is an initial assessment that allows the measure developer to determine the feasibility of assembling the valid and reliable data required for the measure and calculating the measure score using the measure calculation software. According to NQF’s criteria for evaluation, feasibility is the “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.” For PRO-PMs, measure developers must demonstrate that the burden to respondents (i.e., people providing the data) is minimized (e.g., availability and accessibility enhanced by multiple languages, methods, and modes), and the infrastructure to collect instrument-level data is integrated into workflows and EHRs. For digital measures, preliminary feasibility assessments (consisting of both the data model and how various EHR systems map and store the data elements) need to confirm the desired information is available within an EHR or can be added, ideally in a structured format. Measure developers can use NQF’s Feasibility Scorecard to rate the feasibility of a measure’s data elements using four domains: data availability, data accuracy, data standard, and workflow. The measure developer should perform testing of the measure calculation software to demonstrate that the measure logic will work (e.g., BONNIE testing for eCQMs). Although the current state of PROM data collection can create feasibility challenges, the measure developer can mitigate these risks by using the Attribute Grid in the Interim Report (Appendix D) to identify high quality PROMs with LOINC codes and an evidence base that supports successful data collection in the relevant setting from the specified patient population.

**Conduct Beta Testing**

Beta testing is more extensive than alpha testing and is used to gather additional information on feasibility and to assess the scientific acceptability and usability of a measure. It includes testing of the measure logic with data in a live environment and/or care setting and often includes testing of measure results against a “gold standard” reference.

**Assess Feasibility and Usability**

Feasibility should be assessed again once the measure specifications are updated and ready for beta testing to confirm that the finalized data elements can be pulled from the selected data source. Usability should also be evaluated during beta testing. Usability refers to the extent to which potential audiences can interpret and understand performance results. The measure developer should consider unintended consequences and a plan to assess whether they outweigh the evidence of improving healthcare quality.

**Evaluate PRO-PM for Scientific Acceptability**

The scientific acceptability of measure properties is one of the criteria for evaluation that NQF Standing Committees assess when considering a PRO-PM for endorsement. These criteria determine whether the PRO-PM, as specified, produces consistent (i.e., reliable) and credible (i.e., valid) results. The measure developer must not conflate the reliability and validity of the PRO-PM with the reliability and validity of the chosen PROM(s) because these are completely separate concepts.

Reliability comprises two subcriteria, and validity comprises six subcriteria, which are described in detail in NQF’s Measure Evaluation Criteria on the Submitting Standards webpage.
The measure developer must not conflate the reliability and validity of the PRO-PM with the reliability and validity of the chosen PROM(s) because these are completely separate concepts.

Evaluate PRO-PM for Usability and Use
Criteria on usability and use assess the extent to which potential audiences (e.g., patients, clinicians, health systems, health plans, and policymakers) are using or could use the results of the performance measure to achieve the goal of delivering or choosing high quality, efficient healthcare. Details on testing for usability and use are also addressed in NQF’s Measure Evaluation Criteria on the Submitting Standards webpage.

As with many of the tasks in stage 3, the evaluation for scientific acceptability and usability and use is a complex process. The measure developer should begin planning for beta testing as early as possible in the development life cycle and should recognize the iterative nature of this task.

Refine Digital PRO-PM As Needed
Measure developers should use the testing results to determine whether the measure is ready for implementation. If the results show the measure is both reliable and valid, and the benefits outweigh any potential negative consequences, the measure developer should move forward with the measure and begin the implementation process. If the results show low reliability and/or validity, or the stakeholder advisory group expresses concerns about the measure, the measure developer should evaluate what improvements are needed to strengthen the measure in order to prepare it for implementation. The measure developer should also create a mechanism for gathering feedback from those being measured and to consider this feedback when changes are made to the measure.

Stage 4: Finalization and Implementation of the PRO-PM
The final stage of the Roadmap addresses preparing for implementation of the measure and preparation for NQF endorsement review. As with the other stages, the measure developer may address these tasks earlier in the development process based on preference and organizational policies. In fact, the success of the tasks in stage 4 heavily depends on the planning that is recommended during stage 1, as it can be difficult to remedy a shortcoming at the end of the development life cycle if it was overlooked earlier in the process.

Prepare for Implementation
A systematic approach to implementation can minimize unique challenges from varied clinical settings and contribute to the success of implementation across diverse clinical environments. The measure developer is responsible for developing implementation guidance and should consider using an iterative process that incorporates information from every stage of the PRO-PM development life cycle, including context on data collection via the PROM(s) and lessons learned during testing. It should contain specific examples that are applicable in the relevant settings and with the targeted populations.
guides must provide additional information outside of the dQM technical specification that elucidates the measure developer’s intent for each cohort definition and how to ensure the reliability of the information sought in local data systems.

The implementation guidance is a resource document that prepares implementers to put the new measure into practice. By educating implementers on suggested resources and other services that might support measure implementation, the measure developer should collaborate with representatives from testing sites as well as the advisory stakeholders to prepare a guide that will help the entities adapt to the new measure and, ultimately, facilitate improved patient outcomes.

*Prepare for Endorsement Submission*

Measures endorsed by NQF have undergone careful evaluation through a multistakeholder consensus-building endorsement process, a process designed intentionally to garner highly diverse stakeholder perspectives. These stakeholders consist of clinicians, hospitals and other healthcare providers, employers, health plans, public agencies, community coalitions, patients, and caregivers. This process ensures all NQF measures meet thorough standards for performance measures.

Preparing for NQF endorsement review is specific to the measure steward or developer. As referenced in stage 1 of this report, the measure developer should determine whether they intend to submit the PRO-PM for NQF endorsement review prior to moving forward with the early stages of development. An important consideration throughout the development process is whether testing of the digital PRO-PM is robust enough to support endorsement review.

If the measure developer and the stakeholder advisory group determine in stage 1 that NQF endorsement is appropriate for the digital PRO-PM, it is essential that the measure developer understand the NQF Intent to Submit process. Developers who have guided PRO-PMs through the endorsement process note that Intent to Submit is an intensive process that should be started as early as possible. The Submitting Standards webpage provides an overview of the Intent to Submit process and links to an Intent to Submit checklist.

*The Roadmap is not a comprehensive resource for seeking NQF endorsement.* Extensive information on both the endorsement process and measure evaluation criteria is available on the NQF website, including the following pages that may be particularly useful to measure developers:

- The Submitting Standards page
- A description of the CDP
- The Measure Evaluation Criteria
- The Measure Developer Guidebook

As mentioned previously in the Roadmap, the CMS Measures Management System Blueprint is also a valuable source of guidance, and the measure developer should closely review this website.

In addition to the tasks outlined in the Roadmap, the measure developer will benefit from documenting information throughout the PRO-PM development process, including rationale for key decisions, lessons learned throughout the development life cycle, and a proposed plan to maintain and update the PRO-PM in the future.
Next Steps

The Roadmap is the first iteration of a guidance document on digital PRO-PMs. Although this document is intended to be evergreen by linking to external resources, the TEP recognized how the evolving nature of measurement will affect the Roadmap over time. The TEP highlighted two particularly dynamic elements in the Roadmap, digital measurement and measurement of initiatives to reduce health disparities. Additionally, new information about key areas of measure development (e.g., attribution, risk adjustment, and feasibility and usability) will emerge as more PRO-PMs are developed, piloted, and incorporated into payment models. For these reasons, the TEP recommended that NQF and CMS periodically revisit and update the Roadmap to ensure it remains meaningful to measure developers, accurately highlights the challenges and solutions that are unique to PRO-PMs, and continues to act as a catalyst for new PRO-PMs.

The TEP also recognized areas of potential confusion related to patient-reported measurement. The similarity of terminology for PROs, PROMs, and PRO-PMs can hinder understanding of patient-reported measurement at a time when CMS is prioritizing the elevation of patients’ voices. Similarly, the TEP and public commenters throughout the Building a Roadmap initiative note that patient-reported experience and PROs should be recognized as serving different purposes in the body of measurement. The TEP recommends focused expert guidance to drive clarity and improvement in both of these areas.

Conclusion

The Roadmap is both a guide for measure developers and a catalyst to elevate patients’ voices. It is a guide because it offers measure developers straightforward information on the unique aspects of developing PRO-PMs that are suitable for NQF endorsement and use in CMS’ VBP programs and APMs. It is a catalyst because it will facilitate the creation of additional PRO-PMs that measure what matters to patients, using data that patients provide.

When combined with the CMS Blueprint, NQF’s Measure Developer Guidebook, NQF’s Measure Evaluation Criteria, and the Attribute Grid in the Interim Report, this Roadmap is a resource that can help measure developers navigate the development of high quality digital PRO-PMs. By familiarizing themselves with the four stages of digital PRO-PM development (i.e., Definition of Measurement Goals, Exploration and Assessment of PROMs, Development and Testing of the PRO-PM, and Finalization and Implementation of the PRO-PM) and the 17 tasks that exist within those stages, measure developers can gain an overview of the PRO-PM development process in a relatively short period of time. The Roadmap is not a textbook that will answer every question related to the development of digital PRO-PMs, but it is a primer that will help measure developers understand a complicated process, ask informed questions, and ultimately aid in building a more robust database of digital PRO-PMs.
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Appendices

Appendix A: Technical Expert Panel (TEP) Members, Federal Liaisons, Key Informant Interviewees, NQF Staff, and CMS Staff

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Appendix B: Relevant Resources

NQF Resources

Resources and publications about PROs and PRO-PMs

- The 2012 white paper, PRO-Based Performance Measures for Healthcare Accountable Entities
- The 2012 white paper, Methodological Issues in the Selection, Administration, and Use of Patient-Reported Outcomes in Performance Measurement in Health Care Settings
- The 2015 update to the 2012 Methodological Issues white paper written by David Cella et al, titled Patient-Reported Outcomes in Performance Measurement
- The 2013 Committee Final Report, Patient-Reported Outcomes in Performance Measurement
- The 2020 Committee Final Report, Patient-Reported Outcomes: Best Practices on Selection and Data Collection
- The 2021 Building a Roadmap Interim Report on high quality PROMs for use in PRO-PMs

Resources and publications about the Consensus Development Process

- CDP homepage
- Submitting Standards
- Measure Evaluation Criteria
- Measure Developer Guidebook for Submitting Measures to NQF (PDF, Version 6.5, July 2022)
- Intent to Submit Checklist and Guidance (PDF, May 2021)

NQF reports about pertinent areas of measure development

- Risk Adjustment for Sociodemographic Factors (2014)
- Attribution - Principles and Approaches (2016)
- Attribution for Critical Illness and Injury (2021)

U.S. Department of Health & Human Services Resources

CMS reports about pertinent areas of measure development

- CMS Measures Management System
- CMS Blueprint Measure Lifecycle Overview
- CMS Blueprint QuickStart Guide (PDF, May 2022)
- Supplement on Patient-Reported Outcome Measures (PDF, May 2022)
- Supplement on Risk Adjustment in Quality Measures (PDF, May 2022)

CMS and eCQI resources on digital quality measures

- eCQI Resource Center
- Overview of electronic clinical quality measures (eCQMs)
- dQM Strategic Roadmap
- Digital Quality Measurement Strategic Roadmap (PDF, March 2022)
- Digital Quality Measurement Strategic Roadmap – Executive Summary (PDF, March 2022)
- Explanations of Qualified Clinical Data Registries (QCDR)
- Overview of Clinical Quality Language (CQL)
- Overview of United States Core Data for Interoperability (USCDI)
- The ONC USCDI+ website, which describes the initiative and provides links to additional resources

**Additional Resources on Interoperability and Technical Issues**
- Fast Healthcare Interoperability Resources (FHIR)
- FHIR Implementation Guide Registry
- FHIR Confluence site
- The Regenstrief Institute homepage on Logical Observation Identifiers, Names and Codes (LOINC)

**Other Resources**
- The Council of Medical Specialty Societies (CMSS) resources on clinical registries
- The Assistant Secretary for Planning and Evaluation (ASPE) Report to Congress: Social Risk Factors and Performance Under Medicare’s Value-Based Purchasing Programs (2016)
- The ASPE Second Report to Congress: Social Risk Factors and Performance Under Medicare’s Value-Based Purchasing Programs (2020)
- CMS Framework for Health Equity (2022)
Appendix C: Glossary of Terms

Accountable Unit
The entity whose performance is being measured, which could be a hospital, health plan, clinician, etc. Performance measurement can be applied to any setting and level of analysis.  

Alternative Payment Models (APMs)
A payment approach that gives added incentive payments to provide high quality and cost-efficient care. APMs can apply to a specific clinical condition, a care episode, or a population.

Anchors
Anchor-based methods are one of three types of methods used to determine minimal clinically important difference (MCID). A numerical scale for an outcome is “anchored” to a subjective and independent assessment of improvement. For example, a response of “a little better” to a question about how the patient feels post-treatment can be anchored to a numeric outcome.

Attribute
A characteristic or trait of a PROM. Past National Quality Forum (NQF) reports have used attribute and characteristic synonymously. Throughout the Roadmap, attributes primarily refer to the characteristics that make a PROM suitable for use in a PRO-PM.

Attribute Grid
A table in the Interim Report designed to facilitate a side-by-side comparison of different PROMs against 12 attributes of high quality PROMs.

Attribution
A process used in quality measurement that aims to assign accountability for a patient’s outcomes to a clinician, groups of clinicians, or a facility.

Burden
Burden refers to the time, effort, or other demands placed on respondents or those administering the PROM. This can include the number and complexity of items and the literacy level needed to understand and complete the measure.

Crosswalk
A concordance table to convert scores from one scale to the other and vice versa. Crosswalks can allow harmonization of PROMs that measure similar outcomes (e.g., HRQoL after a knee replacement surgery), which may facilitate multicenter collaboration or allow sites to switch PROMs without loss of historic comparison data.

Cut Points
Clinically meaningful thresholds of a score change within a PROM that is often associated with either improvement in patient outcome or indication of need for treatment.
Digital Quality Measures (dQMs)

Quality measures, organized as self-contained measure specifications and code packages, that use one or more sources of health information that is captured and can be transmitted electronically via interoperable systems. Data sources for dQMs may include administrative systems, electronically submitted clinical assessment data, case management systems, electronic health records, laboratory systems, prescription drug monitoring programs, instruments (for example, medical devices and wearable devices), patient portals or applications (for example, for collection of patient-generated data such as a home blood pressure monitor, or patient-reported health data), health information exchanges, or registries, and other sources. 13

Electronic Clinical Quality Measures (eCQMs)

An electronic clinical quality measure (eCQM) is a clinical quality measure expressed and formatted to use data from electronic health record (EHRs) and/or health information technology systems to measure healthcare quality, ideally data captured in a structured form during the process of patient care. For the measured entity to report an eCQM from an EHR, eCQM developers format the Health Quality Measure Format using the Quality Data Model to define the data elements and Clinical Quality Language to express the logic needed to evaluate a provider or organization’s performance.

Interpretability

The degree to which the meaning of the scores can be easily understood by any group requiring use of the scores. A PRO measure should have documentation to support interpretation of scores, including the meaning of low and high scores, representative mean(s) and standard deviation(s) in the reference population, and guidance on the minimally important difference in scores between groups and/or over time. 46

Logical Observation Identifiers, Names, and Codes (LOINC)

LOINC is a database and universal standard for identifying medical laboratory observations. It was developed in 1994 and is maintained by the Regenstrief Institute, a U.S. nonprofit medical research organization. LOINC was created in response to the demand for an electronic database for clinical care and management and is publicly available at no cost. 41,42

Measured Entities

Measured entities are the front-line clinicians and their organizations, including health information technology, collecting quality measurement data. Measured entities are the implementers of quality measures. The effect of quality measure data collection on clinician workflow can be negative. There may be effects on their payments, positive and negative, with respect to reporting and actual performance on quality measures. Because of these potential effects, measured entities should be involved in all aspects of the Measure Lifecycle. 43

Minimal Clinically Important Difference (MCID)

This is the smallest improvement needed after treatment that would be considered worthwhile from the patient’s perspective. 37 MCID can be calculated using three different methods: consensus or delphi method, which depends on consensus of an expert panel; anchors (described above); and a distribution-based method, which relies on the statistical analysis of the distribution of outcome scores. 37
**Patient-Reported Outcome (PRO)**
The status of a patient’s (or person’s) health or behavioral condition that comes directly from the patient without interpretation of the patient’s response by a clinician. 9

**Patient-Reported Outcome Measure (PROM)**
The tools and instruments that are used to collect PRO data. 9 Depending on the measurement concept, PROMs can be used to collect data for PRO-PMs. 9

**Patient-Reported Outcome Performance Measure (PRO-PM)**
How PROs are calculated. A way to aggregate the information from patients into a reliable, valid (tested) measure of performance at the healthcare entity level (e.g., a hospital, health plan, or clinician). 5

**Performance Measures (PMs)**
These are standards that can be used to measure and quantify healthcare processes, outcomes, patient perceptions, organizational structure, and/or systems that are associated with the ability to provide high quality care. 44

**Psychometric Soundness**
How consistently and accurately an assessment measures what it purports to measure. 34 Validity and reliability are key aspects to attaining psychometric soundness. Psychometrics is a scientific discipline concerned with how psychological constructs (e.g., intelligence, neuroticism, or depression) can be optimally related to observables (e.g., outcomes of psychological tests, genetic profiles, and neuroscientific information). 45

**Value-Based Purchasing (VBP) Program**
Value-based programs reward healthcare providers with incentive payments or penalties for the quality of care they give to people with Medicare. These programs are part of CMS’ larger quality strategy to reform how healthcare is delivered and paid for. 46
Appendix D: Attributes of High Quality PROMs for Use in PRO-PMs

Originally, this was Appendix C from the Building a Roadmap Interim Report. For samples of a completed Attribute Grid, please see Appendices D and E in the Interim Report. This appendix shows the Attribute Grid with columns for four PROMs that could be compared side by side. (Any number of PROMs can be compared in the grid by adding or removing columns.) The asterisks in this appendix are placeholders for where the measure developer would enter relevant information about each PROM.

<table>
<thead>
<tr>
<th>ATTRIBUTE</th>
<th>PROM 1</th>
<th>PROM 2</th>
<th>PROM 3</th>
<th>PROM 4</th>
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<tbody>
<tr>
<td>Covers desired PROs from patient and/or caregiver perspective</td>
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<td>Outcome measured in PROM is the result of care for which relevant</td>
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<td>clinical quality is being measured</td>
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<td>Interpretable scores, defined and actionable cut points or targets, and</td>
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<td>anchors and/or defined meaningful change</td>
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<td>Clear conceptual and measurement models</td>
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Appendix E: Public Comments and Responses

The draft Technical Guidance Report for the Building a Roadmap From Patient-Reported Outcome Measures to Patient-Reported Outcome Performance Measures initiative was posted on the National Quality Forum (NQF) project webpage for public and NQF member comment from September 21, 2022, through October 13, 2022. Six prompts were offered to guide public commenters on key areas of interest. Seventeen comments from three organizations are grouped below by organization, and the responses from NQF and the TEP are included beneath each comment. Unless otherwise noted, public comments are presented as they were received by NQF and have not been edited, except for minor updates to spacing, spelling, and punctuation. Page numbers in the public comments refer to a draft version of the Roadmap and may not align with the final version.

American Association on Health and Disability

Comment:
The American Association on Health and Disability and the Lakeshore Foundation appreciate the opportunity to provide comments on draft #2 PRO-PM.

The American Association on Health and Disability (AAHD) (www.aahd.us) is a national non-profit organization of public health professionals, both practitioners and academics, with a primary concern for persons with disabilities. The AAHD mission is to advance health promotion and wellness initiatives for persons with disabilities. AAHD is specifically dedicated to integrating public health and disability into the overall public health agenda.

The Lakeshore Foundation (www.lakeshore.org) mission is to enable people with physical disability and chronic health conditions to lead healthy, active, and independent lifestyles through physical activity, sport, recreation, and research. Lakeshore is a U.S. Olympic and Paralympic Training Site; the UAB/Lakeshore Research Collaborative is a world-class research program in physical activity, health promotion and disability linking Lakeshore’s programs with the University of Alabama, Birmingham’s research expertise. The American Association on Health and Disability and the Lakeshore Foundation comments have several observations and purposes:

1. Reinforce the report’s recognition of the importance of patient/beneficiary/program participant/consumer engagement (and their caregivers, families, and advocates).
2. Advocate for greater focus on health equity in the report’s work and recommendations.
3. Advocate for more substantial and ongoing involvement by public program administrators who have actual experience implementing PROs and PRO-PMs in the public domain.
4. Raise the question of the balance between promoting the whole-person health and wellness of persons with disabilities vs use of PROs/PRO-PMs in targeted (siloeed) public programs, particularly Medicaid and Medicare.
5. Observe, as we have done previously, that the major PROs/PRO-PMs currently serving persons with disabilities are missing as reference points and lessons learned.
6. Question who gets left out with the report, CMS, and NQF emphasis on digital quality measures.

Response:
Thank you for your ongoing support for and input on the Building a Roadmap initiative. Responses to each of the six themes of your comments are below.
Comment:
Reinforce the report’s recognition of the importance of patient / beneficiary / program participant / consumer engagement (and their caregivers, families, and advocates).

We appreciate the Roadmap intent to be both a guide for measure developers and “a catalyst to elevate patients’ voices” (page 30). We agree with the observation (page 5) that patients, in general, lack awareness about the benefits of PRO-PMs.

We support the recommended establishment of stakeholder advisory groups (pages 14 and 15). Yes, the composition of the stakeholder advisory group will depend on the measure. It is important that the report states that the stakeholder advisory group must include representation from the patients, patient advocacy groups, caregivers, and (or) consumer groups. We suggest the report delete “or” and exclusively use “and.”

We further endorse the page 15 recommendation of the importance of ensuring stakeholders can submit feedback in a timely manner throughout the development life cycle. We support the suggestion that stakeholders identify a consensus definition for the measure concept-desired outcomes, “that will be monitored.” We strongly concur with the pages 15-16 statement that PRO-PMs must be meaningful to patients and/or caregivers and must be meaningful to the program’s target population. As above, we recommend both patients and caregivers, not “or.” Please delete “or.”

Response:
The language in the report is updated to “patients, patient advocacy groups, caregivers, and/or consumer groups, as appropriate.” This edit acknowledges your comment while still recognizing measure developers’ need to make meaningful decisions about the composition of the stakeholder advisory group.

Comment:
Advocate for greater focus on health equity in the report’s work and recommendations.

We appreciate and agree with the page 15 observation: Health equity is the attainment of the highest level of health for all people, and it warrants consideration throughout the measure development process. However, this observation ignores the current priority focus of much of the health care field; advocates, patients, families; most of the quality measurement entities; and federal agencies such as ACL, AHRQ, CDC, CMS, HRSA, and SAMHSA. Addressing health equity requires much greater attention.

We note the page 24 observation that guidance on risk adjustment is evolving. But the urgency and importance of health equity is missing.

Response:
The TEP discussed this comment in detail during its sixth web meeting on October 21, 2022. A new subsection of the report, titled “Health Equity” is now included in the Roadmap. This section includes brief information on the widespread prioritization of health equity across healthcare stakeholders, as well as guidance for measure developers to incorporate health equity considerations across all stages and tasks in the Roadmap. NQF also updated Appendix B to include additional sources on health equity.
Comment:
Advocate for more substantial and ongoing involvement by public program administrators who have actual experience implementing PROs and PRO-PMs in the public domain.

Page 19 lists candidate PRO-PMs from lists of academics and quality measure specialists. MISSING – public program administrators. While most public program administrators bring a conservatism based on available resources, budgeting, and other factors, they have the experience and lessons of using PRO-PMs in actual use (and limitations and barriers). In the area of disability, public program administrators of Medicaid, especially Medicaid Home and Community-Based Services (HCBS), special Medicare programs, and state government service delivery systems including agencies serving persons with ID/DD, mental illness, and related behavioral health conditions.

We agree with the observations (page 24) regarding attribution – for which providers is the patient outcome a signal of the quality of care? Attribution is used in quality measurement to assign accountability for a patient’s outcomes to the accountable entity being assessed by the measure. MISSING: the public program administrators.

Response:
Public program administrators are now included in the list of considerations for the stakeholder advisory group, described in stage 1, task 2 (“Establish Stakeholder Advisory Group and Feedback Processes”). The list of resources in stage 2, task 1 (“Identify Candidate PROMs”) is not comprehensive, and the inclusion of professional societies is intended to include those societies and organizations that represent public program administrators. Because attribution is not unique to PRO-PMs, the report does not address detailed information and instead links to source-of-truth resources on attribution.

Comment:
Raise the question of the balance between promoting the whole-person health and wellness of persons with disabilities vs use of PROs/PRO-PMs in targeted (siloed) public programs, particularly Medicaid and Medicare.

In 2020, 7.5 million people (persons with disabilities, mental illnesses, and aging with challenges) received Medicaid home-and-community-based services (HCBS) through both Medicaid waiver programs and state plan benefits. [CMS; Medicaid Beneficiaries Who Use LTSS; July 22, 2022.] In 2020, HCBS expenditures accounted for $125 billion, or 62%, of the $199 billion spent nationally on Medicaid LTSS (CMS: SMD 22-003, HCBS Quality Measures Set, July 22, 2022]. In addition to these 7.5 million persons served, 39 states have HCBS waiting lists of 665,015 persons, largely persons with disabilities. [Kaiser Family Foundation: State Policy Choices About Medicaid HCBS Amid the Pandemic; March 4, 2022.]

So, the report does not address the major programs that serve persons with disabilities, mental illnesses, aging with challenges, and related life situations.

Persons with disabilities and mental illnesses are served by the general health care system. But there are few appropriate individualized services and supports for such persons. Additionally, many of these persons face lack of accessibility and accommodation, discrimination, lack of privacy and confidentiality, and lack of provider knowledge. General health services and specialized behavioral health and disability
services are rarely coordinated, much less integrated. These factors are not mentioned in the report. PRO-PMs need expertise and experience in addressing the whole-person health and wellness of these special populations. And, largely missing are bridges between specialized Medicaid and Medicare programs (silos) and the general health care arena.

Response:
Although the scope of the Roadmap is on CMS value-based purchasing programs and alternative payment models, the guidance on developing digital PRO-PMs is applicable to all programs that utilize PRO-PMs. Therefore, the Roadmap does not specifically list programs in which PRO-PMs can be used, beyond general examples. The intersection of these programs and PRO-PMs is an important opportunity for future work, and NQF hopes to be involved in those efforts.

Comment:
Observe, as we have done previously, that the major PROs/PRO-PMs currently serving persons with disabilities are missing as reference points and lessons learned.

Medicaid home-and-community-based services (HCBS) programs have over 20 years PRO-PM experience in multiple states through the National Core Indicators (NCI) and Personal Outcome Measures (POMs). During the past several years, almost 20 states have implemented a new HCBS-CAHPS (Consumer Assessment of Healthcare Providers and Systems) PRO-PMs. These programs have been discussed, documented, and even endorsed by the NQF. This experience is missing from the NQF PRO-PM guide. The Administration for Community Living (ACL) sponsors several projects examining these HCBS-based PRO-PMs.

Response:
Although the four-stage, 17-task development process described in the Roadmap can be applied to any PRO-PM regardless of programmatic intent, the focus of the Roadmap is on CMS value-based purchasing programs and alternative payment models. The TEP agrees that opportunities exist to learn from HCBS measurement, but that is beyond the scope of this initiative. NQF and the TEP encourage future work that investigates and disseminates these learning opportunities.

Comment:
Question who gets left out with the report, CMS, and NQF emphasis on digital quality measures.

The report emphasizes the Roadmap for the Development of a Digital PRO-PM (page 12), and need for machine-readable specifications (page 23), and measures captured and transmitted electronically (page 42). Digital measures is the hot/buzz topic and development in health care delivery, led by CMS and NQF. But rarely do the advocates, including the NQF PRO-PM report ask: who gets left out? The entire Medicaid HCBS enterprise, the public mental illness and substance use systems largely do not have such digitally based systems – left out by silo thinking and lack of attention. The few operational HCBS systems generally lack any inter-operability. The report should acknowledge this challenge and situation.

Response:
The TEP discussed this comment in detail during its sixth web meeting on October 21, 2022. A new subsection of the report, titled “Health Equity” is now included in the Roadmap, and a brief discussion of proxy responses is added to stage 2, task 2 (“Assess PROMs Using the Attribute Grid and Ensure
Sufficient Testing Has Occurred”). Although the important question of who is left out of digital measurement is beyond the scope of the Building a Roadmap initiative, these additions are concrete reminders to measure developers of the importance of considering the full target population, including people who cannot access digital resources.

*IPO4Health (Improving Patient Outcomes For Health)*

Comment:

General Comment: Review of the current list of Patient Reported Outcome Measures available in the NQF Quality Positioning System yield two categories of quality measures: Patient Reported Experience Measures (PREMs) and true Patient Reported Outcome Measures (PROMs). Only 9 of these measures that are currently endorsed fit the definition of a true PROM, with the remaining endorsed measures being pure PREMs (see attachment). According to Kingsley and Patel,

“PROMs are tools used to measure patient-reported outcomes (PROs). PROMs are standardized, validated questionnaires that are completed by patients’ during the perioperative period to ascertain perceptions of their health status, perceived level of impairment, disability, and health-related quality of life. They allow the efficacy of a clinical intervention to be measured from the patients’ perspective. Questionnaires are given to patients both pre and post operatively to allow comparison of outcomes pre and post procedure. In addition to outcomes relating to interventions, PROMs measure patients’ perceptions of their general health or their health in relation to a specific disease. PROMs are a means of measuring clinical effectiveness and safety.”

And:

“PREMs gather information on patients’ views of their experience whilst receiving care. They are an indicator of the quality of patient care, although do not measure it directly. PREMs are most commonly in the form of questionnaires. In contrast to PROMs, PREMs do not look at the outcomes of care but the impact of the process of the care on the patient’s experience, e.g., communication and timeliness of assistance. They differ from satisfaction surveys by reporting objective patient experiences, removing the ability to report subjective views.”


Also see the COSMIN Database, which does not include PREMs: https://www.cosmin.nl/tools/database-systematic-reviews/

Response:

The “Prioritized PRO-PM Domains” section of the Roadmap discusses the TEP’s decision to focus the technical guidance on PRO-PMs for health related quality of life, functional status, and symptoms and symptom burden. While the TEP and NQF value the importance of patient experience and health behavior measures, the aforementioned domains are priorities, in part due to their underrepresentation in the current body of NQF-endorsed PRO-PMs. The reference you provided is now cited in the Roadmap.
Comment:
Page 28: Evidence evaluation (both foundational rationale for the importance of the PRO-PM and measurable impact from its direct usage): Refer to “Submitting Standards” document pages 23-26 “Importance to Measure and Report: Evidence (Outcomes) (1a.01-1a03). Standardized Logic Models and formal Systematic Reviews are not often adequately presented in submissions of many measure developers in the Consensus Development Process performed by Standing Committees, Technical Expert Panels or the NQF Scientific Methods Panel. Evidence is often simply provided by measure developers via a non-structured, annotated “literature review”, followed by narrative, subjective and non-scientific inferential interpretations of association and/or causation.

Response:
Standardized logic models and formal systematic reviews are applicable to all quality measures. The Roadmap intentionally directs readers to external source-of-truth documentation for guidance on best practices for measure development and endorsement preparation that widely apply to all outcome measures. NQF has shared your comment with its CDP leadership for consideration in future iterations of CDP documentation.

Comment:
Pages 18 and 27: The report only briefly discusses the importance of the usage of PROMs for active and effective shared decision making

From page 18 of Patient-Reported Outcomes: Best Practices on Selection and Data Collection:

“Generally, patients are amenable to completing PROMs that are utilized and discussed by the clinician during the clinical encounter. Reducing barriers to the use of PROMs during patient care (e.g., making scores easy to find by placing them on the EHR’s vital signs screen) is key. For example, ‘Ms. Smith, today your KCCQ score is 54, because of your persistent shortness of breath and swelling that are limiting your physical activities. I am going to prescribe a new medicine and we will see how your KCCQ score changes when I see you again in two months; that will help us know if the new medicine is working.’ Having physicians integrate PROMs into their evaluations and conversations with patients can be transformational.”

This example is not emblematic of true shared decision-making (SDM) because the patient is only informed/told by the heart failure specialist, in essence, what will be done, as opposed to having a structured two-way conversation about the evidence-based rationale for the proposed new treatment option. According to Chan, et al, PROMs and PRO-PMs are often not presented during daily practice to patients by busy clinicians due to time constraints and lack of experience and expertise with SDM. Patients rarely get direct feedback on the importance and meaning of PROM/PRO-PMs when completing and submitting questionnaire instruments (now often done via digital apps).

Response:
While the TEP agrees on the opportunities for PROMs to play an active role in shared decision making, this is outside the scope of providing guidance to measure developers on developing digital PRO-PMs. Structured guidance on the intersection of PROMs and shared decision making is an important opportunity for future work.

Comment:
Page 25: Institutional Review Board evaluation of the risks to Human subject protection can most likely be achieved and encouraged through an expedited review in the evaluation and implementation phases of initial measure development.

Response:
The bullet on IRB evaluation in stage 3 task 2 (“Create Testing Plans”) is updated with new language.

Comment:
Pages 10, 12 & 15: Identification of evidence-based causal linkages between meaningful/reliable/validated change scores of PROMs and associated PRO-PMs and important relevant clinical outcomes (e.g., mortality, measurable improvements in biometrics, hospitalization, etc.) is necessary for successful implementation and use in daily practice.

Response:
A brief description of meaningfully important clinical difference has been added to the “Interim Report Recommendations” section of the report.

Comment:
Page 20: Please be more specific in specifying a robust framework for “psychometric soundness”, including inclusion of evidence based systematic reviews and an explicitly stated standardized logic model for the implementation and use of a PRO-PM.

Response:
While the TEP acknowledges your suggestions, a framework for assessing the psychometric soundness of PROMs is out of scope for the Roadmap. The Roadmap addresses the fact that there are multiple approaches to developing a quality measure, and the same holds true with demonstrating psychometric soundness of a PROM. Notably, NQF does not evaluate the psychometric soundness of PROMs unless the PROM is specified as a data collection tool for a PRO-PM that is under evaluation for endorsement.

Comment:
Page 17 and 26: Standardized clinical registries, especially CMS approved Qualified Clinical Data Registries (QCDRs) are ideal sources for PROM and PRO-PM data element collection, automated calculations, and tracking. Registry functions better sources.

Response:
New language is added to the bulleted list under stage 3, task 1 (“Specify Key Measure Details”) to include registries and other potential data sources.
Comment:
Page 22: Rationale for inclusions, exclusions from Denominators and Numerators. Also consider the relative risk stratification (separate from risk adjustment) e.g., stratification of ACC/AHA Stage C & D Heart Failure for the Kansas City Cardiomyopathy Questionnaire (KCCQ).

Response:
Regarding numerator and denominator definitions, new language is added under stage 3, task 1 (“Specify Key Measure Details”) about including rationale for inclusion and exclusion criteria. Regarding stratification, a new paragraph is added under the “Develop and Test Risk Adjustment Model” in stage 3, task 1.

Comment:
Page 26 Digital PROMs/EHR rationale: Take into account the likely usage of Clinical Decision Support software that may be used in electronic health records and other digitally deployed Application Program Interfaces (APIs) in the process of implementation of PROMs and PRO-PMs. Such digitally deployed resources may require clarification by the Food and Drug Administration (FDA) regarding the types of clinical decision support (CDS) software functions that are excluded from the definition of device by the criteria in section 520(o)(1)(E) of the FD&C Act (“Non-Device CDS criteria”). FDA has recently published final guidance further clarifying that FDA’s existing digital health policies continue to apply to software functions that meet the definition of a device, including those that are intended for use by patients or caregivers. See https://www.fda.gov/regulatory-information/search-fda-guidance-documents/clinical-decision-support-software.

Response:
Data sources, including but not limited to clinical decision support software, are regularly emerging and evolving. The Roadmap does not attempt to identify each data source. Measure developers should stay up-to-date on new data sources and the implications of each, including unique considerations by the FDA and/or other federal agencies.

Kidney Care Partners

Comment:
Kidney Care Partners (KCP) appreciates the opportunity to comment on NQF’s second draft technical guidance report, Building a Roadmap From Patient-Reported Outcome Measures (PROMs) to Patient-Reported Outcome Performance Measures (PRO-PMs). KCP is a coalition of 32 organizations comprised of patient advocates, dialysis professionals, healthcare providers, researchers, and manufacturers organized to advance policies that support the provision of high-quality care for individuals with chronic kidney disease (CKD) and end stage renal disease (ESRD).

KCP applauds NQF for undertaking this important and timely work. We support measurement and welcome guidance and innovation in this increasingly important area, and appreciate NQF’s work to establish guiding principles for these inherently complex measures. We believe this new NQF report provides a solid technical framework that will help guide measure developers in both the selection of appropriate and psychometrically sound existing surveys/questionnaires (i.e., PROMs) around which to build a PRO-PM, as well as in the design and development of the PRO-PM itself in a manner that will satisfy NQF’s rigorous endorsement criteria.
KCP believes it is critically important to assess patients’ reported experiences related to their dialysis treatments and their interaction with nephrologists. We thus support use of PRO-PMs in the federal ESRD programs. However, we note that the considerable flaws we and other stakeholders have identified with the only PRO-PM included in the ESRD Quality Incentive Program (QIP) to date, the In-Center Hemodialysis Consumer Assessment of Healthcare Providers and Systems (ICH-CAHPS) Survey, fundamentally compromise both its validity as a publicly reported measure and its utility as an instrument of quality improvement. These shortcomings highlight the intrinsic complexity of these metrics and the thoughtfulness with which they must be applied to yield valid, meaningful results—particularly in a penalty-based quality program such as the QIP. Of particular concern are patient response rates, which had been steadily declining since the measure’s implementation and were at only 35 percent prior to COVID; rates have continued to decline during the pandemic. Such low response rates, which necessarily threaten measure validity, are a result of the burdensome manner in which the lengthy survey is fielded to patients—i.e., patients are asked to complete the full 62-item questionnaire twice annually, generating considerable survey fatigue. While ensuring anonymity, the requirement that dialysis facilities field the survey using an independent third-party vendor likely contributes to low response rates as well. Additionally, in a 2018 cross-sectional analysis of ICH-CAHPS survey administration to 11,055 eligible in-center hemodialysis patients across the U.S., Dad et al.[1] reported that non-responders (6,541 [59 percent]) significantly differed from responders, broadly spanning individuals with fewer socioeconomic advantages and greater illness burden—further highlighting limitations in interpreting facility survey results and raising concerns about the risk of the measure paradoxically promulgating health inequities. (Dad T et al. Evaluation of non-response to the In-Center Hemodialysis Consumer Assessment of Healthcare Providers and Systems [ICH CAHPS] survey. BMC Health Services Research. 2018;18:790.) Another significant deficiency with the ICH-CAHPS is that facilities and patients are not granted the opportunity to use the survey results to improve care; facilities never see individual patients’ results and thus cannot communicate with patients or effectively address their identified concerns, leaving patients feeling as if they wasted their time completing the survey and providers at a loss as to how to improve performance. Finally, while we have long advocated for the inclusion of home dialysis patients in the existing instrument (or the adoption of a corollary home measure into the QIP), the measure continues to exclude this increasingly important and rapidly growing population.

KCP thus appreciates NQF’s thoughtful recommendations outlined in this draft report. In particular, we welcome NQF’s acknowledgement of the fielding burden issue with PRO-PMs and we support future exploration of suggested remedies such as increased digital interoperability, shared data formats, short forms, and computerized adaptive testing [CAT]; however, we note that simpler strategies such as reducing survey administration frequency should not be overlooked. Additionally, we urge NQF to consider adding specific guidance on how disparities in response rates stemming from intrinsic social drivers of health can best be effectively addressed, and on how data collected through PRO-PMs can and should be used—specifically, to offer providers direct insight from their patients on how to improve the quality of their care.

KCP looks forward to working with NQF in this exciting area in the future. In the interim, we would like to call attention to our own prior work addressing this important topic, KCP’s Patient-Reported Outcomes for End-Stage Renal Disease: A Framework for Priorities and Measurement. This multi-stakeholder consensus document was developed by KCP in 2017 to identify priorities and outline guiding
principles for PRO-PM development, and to establish a framework for what should be measured to effectively assess patients’ perspectives of their dialysis care.

KCP again thanks you for the opportunity to comment on this important work.

**Response:**
Thank you for your support for and input on the Building a Roadmap initiative. The TEP understands the concerns you raise about the dearth of PRO-PMs specific to chronic kidney disease and end-stage renal disease. These concerns align with the Roadmap’s goal of fostering new PRO-PM development by providing measure developers with specific and actionable guidance on developing PRO-PMs. The TEP appreciates your work to develop a framework for PRO-PMs in CKD and ESRD.