

# 2017 Consensus Development Process Redesign

#### FINAL REPORT

## Background

The National Quality Forum's multi-step Consensus Development Process (CDP) is essential to providing a usable portfolio of measures that meets NQF's rigorous measure evaluation criteria and ensures that measures integrated into HHS' public reporting and pay-for-performance initiatives are up-to-date, reflective of the current evidence, reliable and valid, useful for accountability and quality improvement, and feasible. Since the first version of the CDP (approved in July 2000), NQF has continuously refined its process to address the needs of CMS, NQF members, and the healthcare industry more broadly. Many of these refinements have been incremental and others more substantive, requiring pilot testing and substantial operational changes. However, CMS and other stakeholders have raised concerns about the agility of the CDP – specifically, the time from measure submission to measure endorsement and the timeliness of measure evaluation/wait time for available projects (which in some cases is three or more years).

## Approach

NQF hosted a process improvement, or Kaizen event on May 18-19, 2017, to explore opportunities for a more agile and efficient CDP for measure endorsement. Over the two-day event, NQF, in collaboration with the Centers for Medicare & Medicaid Services (CMS), sought to address:

- Improving coordination among CMS, developers, and NQF to better facilitate timely evaluation of measures
- Increasing opportunities for submission and timely review of measures
- Reducing cycle time of the CDP
- Improving flow of information between the CDP and Measure Applications Partnership (MAP) processes

## Objectives

Specifically, through the Kaizen, NQF was committed to exploring opportunities for:

- Continuous availability of CDP for all measure types
- Improved management of the CDP measure pipeline
- Improved utilization of standing committee expertise
- Improved leveraging of NQF and external expertise
- Significant reduction in overall endorsement time to about 6 months

More than 40 invited healthcare stakeholders from the public and private sectors participated in the event—including experts from CMS and other federal agencies, NQF's standing committees, and organizations that develop measures that represented a significant proportion of participants.

## **Proposed Redesign**

Based on the outputs from the Kaizen event, NQF will undergo a significant CDP redesign that incorporates on-going measure submission opportunities. (Currently 63% of standing committees experience an average of three years of dormancy.) Offering more continuous and predictable submission pathways can increase the timeliness of endorsement decisions for measures that will drive value and fill prioritized gaps. Recommended changes include:

Proposed Changes	Implementation Timing
Increase stakeholder training and education	Summer 2017
Improve information exchange and access	Summer 2017-2019 (phased approach)
Implement Intent to Submit process	Fall 2017
• Form a newly-convened NQF Scientific Methods panel	Fall 2017
<ul> <li>Implement continuous commenting period and NQF member support of expression</li> </ul>	Beginning Fall 2017
Revise the technical report— content and structure	Fall 2017
• Designate Standing Committee as the final endorsement body1	TBD
• CSAC Role change and disbandment of the Appeals Board 2	TBD

Some of the changes intended to compress the endorsement process will help to reinforce process changes that have already proven to be effective (i.e., standing committees and staff preliminary analyses). While other changes will establish new processes that reflect increased efficiencies in stakeholder participation and engagement.

NQF will not implement all changes immediately, as this will require significant resources, input from several stakeholders like NQF's Governance Committee and Board of Directors (e.g. changes to the standing committee, CSAC and Appeals Board's roles), design and testing to ensure that the process works as intended for all stakeholders. However, NQF will initiate a phased implementation in order to monitor these recommendations to assess outcomes and ensure a more agile and effective process.

<sup>&</sup>lt;sup>1</sup> If changes to the roles of the Standing Committee and the CSAC are approved, execution of these changes will be phased and implemented at a later date

<sup>&</sup>lt;sup>2</sup> If the change in the role of the Appeals Board is approved, execution of this change will be phased and implemented at a later date

## Increased Opportunities for Measure Submission: Scheduling/Frequency

NQF will offer two measure submission opportunities for each topic area each year; instead of one opportunity for a select, few topic areas each year per the current CDP schedule (see Figure 1). However, because there would be more opportunities for submission, NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures).<sup>3</sup> This was determined given that approximately 80% of the measures submitted for endorsement consideration are maintenance measures. The combination of maintenance and new measures may vary depending on the number of measures submitted, opportunities for related and competing measure review, and measure prioritization efforts. Per NQF's maintenance of endorsement policy, measures are due for reassessment every three years. NQF will remind measure stewards and developers of scheduled measure maintenance review several months prior to the review and notify each of their assigned review cycle.



#### Figure 1. Schedule of Measure Review Cycle

Due to the anticipated increased opportunities for measure submission, NQF has consolidated the 22 measure review topical areas to 15 topical areas as shown in Figure 2 below4:

<sup>&</sup>lt;sup>3</sup> NQF may consider including one or two additional measures within a cycle as needed.

<sup>&</sup>lt;sup>4</sup> These are recommended topic areas. Per NQF's current process, these topics will be reassessed periodically to ensure the appropriate measure groupings.

#### Figure 2. Measure Review Topical Areas



<sup>A</sup> Patient Safety will include acute infectious disease and critical measures

<sup>B</sup> Prevention and Population Health is formerly Health and Well Being

<sup>C</sup> Cost & Efficiency will include efficiency-focused measures from other domains

<sup>D</sup> Geriatric & Palliative Care includes pain-focused measures from other domains

Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures, while distributing measures to committees with the needed expertise to conduct an evaluation. As a result, many of the smaller portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more cross-cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care. Individual standing committees that will no longer convene for the following topical areas include:

- Person and Family-Centered Care
- Ears, Eyes, Nose and Throat Conditions
- Endocrine
- Musculoskeletal
- Infectious Disease
- Care Coordination
- Gastrointestinal
- Genitourinary

Each topical area will have a seated standing committee to help shape the endorsement project's scope, offer expert advice, ensure that input is considered from relevant stakeholders, and make

recommendations to the NQF Consensus Standards Approval Committee (CSAC) on measures proposed for endorsement. For larger topic areas that include multiple conditions or cross-cutting areas, NQF will utilize technical expertise in specific areas as needed. All committee members will be subject to NQF's current Conflict of Interest Policy.

To allow more frequent measure submissions, committees will convene more often. NQF will host a combination of in-person meetings and virtual web meetings to evaluate submitted measures. Since there will be two review cycles each year, the committee will convene via in-person meeting for one cycle and convene via virtual web meeting for the other cycle to be cost efficient. NQF will make every effort to standardize how both in-person meetings and web meetings are conducted to ensure consistency in the Committee's measure review and evaluation process.

### Measure Cycle Review

This report includes descriptions of revised processes to the extent possible. However, details and timing of these processes may change as implementation continues. Figure 3 below shows how a measure will move through the CDP for endorsement consideration.





Prior to the start of an evaluation cycle, NQF will announce staggered measure submission deadlines twice per contract year --- for any measure, any topic. During this time, any measure steward/developer, assuming responsibility for making the necessary updates to the measure, can submit a new measure for endorsement consideration. In addition to newly submitted measures, NQF-endorsed measures undergo evaluation for maintenance of endorsement approximately every three years. All measures must be submitted by the cycle submission deadline and will be evaluated against NQF's Measure Evaluation

Criteria. To submit a measure for an initial endorsement evaluation or a maintenance of endorsement evaluation, a measure steward must complete or update the online measure submission form and submit an *Intent to Submit* form.

### Intent to Submit

An intent to submit will require that all measure stewards/developers notify NQF of their readiness to submit measures for endorsement consideration. The *Intent to Submit* form will require the following information:

- **Submission Type**: maintenance measure (currently NQF-endorsed) or new measure (has never received NQF endorsement). Maintenance measures must indicate if new testing data will be available.
- **Measure type** measure categorization (e.g., structure, process, etc.) and level of complexity (e.g., outcomes, cost or resource use, instrument-based, etc.)
- Measure title concise description to convey who and what is being measured
- Level of analysis levels for which the measure is assessed—specified and tested
- Data source source(s) from which data are obtained for measurement
- **Measure description** brief narrative of the measure that includes the type of score, measure focus, target population, or time frame
- Numerator statement brief description of the measure focus or what is being measured
- Denominator statement brief description of the target population being measured
- **Planned submission date** cycle and year when all testing is completed and final submission is anticipated

Measure stewards/developers will need to notify NQF at least three months prior to the measure submission deadline to prepare for the committee's review in the upcoming cycle. This will allow NQF to adequately plan for measures in the pipeline and maintenance measures ready for evaluation in the various topic areas. NQF also welcomes measure stewards/developers to request technical assistance during this time. NQF staff will assess whether a measure is sufficiently 'complex' to require a methodological review by the Scientific Methods Panel, based on a set of criteria (details below). Because the newly formed Scientific Methods Panel will evaluate the *Scientific Acceptability* of new (and some previously endorsed) complex measures, measure stewards/developers must submit measure specifications and testing information along with the *Intent to Submit* form at least three months prior to the measure submission deadline.

### Technical Review: NQF and Scientific Methods Panel Review

Kaizen participants noted the challenges many committee members face when reviewing measures and applying NQF's measure evaluation criteria to the technical aspects of reliability and validity analyses and results, and therefore recommended removing this responsibility from the committee. NQF will operationalize this recommendation through a "methods review". As noted above, the methods review will conducted by the newly formed external NQF Scientific Methods Panel for complex measures. NQF will continue to provide a preliminary analysis, including the methods review for non-complex measures. The opportunity for a methods review is seen as a value add for the standing committee and developers

because it will reduce committee burden , particularly where committee members do not always have the needed expertise to adequately review and rate the scientific merits of a measure. Further, removal of this more technical review should encourage greater participation by consumers, patients, and purchasers in standing committees.

This methods review will apply to the *Scientific Acceptability* (reliability and validity) section of the measure, both are must-pass criteria. (It is important to note that the Scientific Methods Panel will not render endorsement recommendations. While important, the Scientific Methods Panel review will help to inform the standing committee's endorsement recommendation.) The Scientific Methods Panel would also provide guidance to NQF for methods/testing-related issues. NQF will convene the Scientific Methods Panel via an in-person meeting annually to discuss the methods/testing-related issues.

#### Scientific Methods Panel Composition, Terms, Policies, and Processes

The new NQF Scientific Methods Panel will consist of 15 to 25 statisticians, epidemiologists, psychometricians, economists, performance measure methodologists, and individuals with expertise related to eMeasures and disparities. NQF will solicit and identify nominees through NQF's standard nominations process. Preference will be given to individuals with previous experience on an NQF standing committee. As per NQF's current standing committee process, Scientific Methods Panel members will be randomly appointed to an initial two- or three-year term, with an optional three-year term to follow. All nominees will complete an annual general disclosure of interest (DOI) form, as well as measure-specific disclosure form to identify recusals from specific measures. NQF will assign measure review based on identified conflicts of interest, relevant expertise, and availability of panel members. Much like guidance for standing committees, NQF will provide standard guidance on assessing the *Scientific Acceptability* criterion for a measure, using the current decision algorithm from NQF's Measure Evaluation Criteria. To ensure impartiality, panel members will independently evaluate each measure undergoing an external panel review. The majority recommendation will serve as the overall assessment of reliability and validity. NQF will share all evaluations with the measure steward/developer.

#### Complex vs. Non-Complex Measures

Based on input from the Kaizen, the revised measure submission process will consider the complexity of the measure (see Figure 4). A measure will be categorized as 'complex' or 'non-complex' based on information provided in the *Intent to Submit* form.

The following types of measures are considered complex and therefore may require an evaluation by the Scientific Methods Panel:

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

For complex measures, the Scientific Methods Panel will evaluate the measure's reliability and validity (or *Scientific Acceptability* criterion) and provide a preliminary recommendation to NQF staff and the standing committee. Because updated reliability and validity testing is not required for maintenance measures, NQF staff will review previous testing results for complex maintenance measures and determine the adequacy of prior testing. If prior testing is inadequate, updated testing is provided, or NQF staff deems an external review necessary, the measure will be submitted to the external Scientific Methods Panel to evaluate the reliability and validity of the measure. Following the current process, NQF staff will perform a preliminary analysis against all of the other evaluation criteria for both new and maintenance measures. For non-complex measures (e.g., structure and process measures), NQF staff will complete the preliminary analysis against all measure evaluation criteria, including the *Scientific Acceptability* criterion.

For both complex and non-complex measures, when the preliminary analysis is complete, NQF staff will send the preliminary analysis to developers for review. Measures rated by NQF staff or the Scientific Methods Panel as "*Low*" or "*Insufficient*" for reliability or validity will be removed from the current evaluation cycle, allowing time for any additional testing, clarification or NQF technical support, or review prior to consideration of the measure in a future cycle. For all other measures, developers will have up to two weeks to provide further clarifications, if needed. NQF staff will then finalize the preliminary analysis and send the final submission materials to the standing committee for evaluation. If developers disagree with the staff or scientific methods panel review or ratings, they can use the two-week review period to provide additional clarification, which can be considered by staff when finalizing the preliminary analysis. Developers will also have the opportunity to introduce their measures during the committee evaluation meeting and answer questions from the committee during the discussion.



Measure Workflow

#### Figure 4. Measure Workflow

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The rating from the methods review—whether generated by NQF staff or the Scientific Methods Panel will be used to rate the *Scientific Acceptability* of the measure. However, standing committees may raise concerns with the specifications of the measure or with potential threats to validity (e.g., selection of variables for risk adjustment model) and can therefore overturn the staff or Scientific Methods Panel rating. As part of its ongoing education efforts, NQF will provide clear guidance to standing committees regarding the circumstances wherein an overturn of the rating would be permissible.

The relevant standing committee will conduct a detailed review of all measures and its preliminary analysis. During this review process, the committee may meet several times, via web meetings, conference calls and/or in-person meetings, to discuss and evaluate the submitted measures in accordance with NQF criteria and guidance. After a standing committee completes its initial review of the submitted candidate standards, a draft of the committee's recommendations—or draft report—will be posted on the NQF website for review and comment by members of NQF and the public.

### Measure Evaluation Technical Report – Content and Structure

After the standing committee completes its initial measure review, a draft of the committee's recommendations – or "draft report" – will be posted on the NQF website for the public and NQF membership to review and comment. To minimize the length and density of the technical report, NQF will revise the content and structure of the report.

This report will include:

- an executive summary that indicates the endorsement decision
- brief summaries of each measure reviewed
- details of the committee's deliberations on each measure against NQF's measure evaluation criteria (in appendix)
- full measure specifications for each measure reviewed (in appendix)

Any remaining background information on the topic area, including its alignment with the National Quality Strategy, and the NQF portfolio of topic- specific measures will be posted on NQF's public website. In addition, at the end of each two-cycle year, NQF will develop an annual cross-cutting report across all of the topic areas that will summarize trends and performance, high-priority gap areas in measurement for future development, and measure concepts submitted during the solicitation process for measures.

### Continuous Public Commenting Period with Member Expression of Support

As part of NQF's commitment to transparency, both NQF members and interested members of the public can submit comments on the standing committee's recommendations through the NQF website. In place of two separate public commenting periods (14-day pre-meeting commenting and 30-day post-meeting commenting), NQF will have one continuous public commenting period. This commenting period will span 12 weeks to allow adequate time for the public and NQF member commenting. The commenting period would open approximately three weeks prior to the committee evaluation meeting and close 30 days after NQF posts the draft technical report on the NQF website. NQF will include all comments received at least one week prior to the committee evaluation meeting into the committee materials for discussion during

the meeting. NQF will ensure the measure steward/developer receives the submitted comments in a timely manner to prepare for the committee evaluation meeting. Measure steward/developers are not required to provide written responses to the comments received prior to the measure evaluation meeting. The committee will review any comments received after the committee evaluation meeting during the post-commenting period call. All submitted comments during this time will receive written responses from the standing committee, measure steward/developers, and/or NQF, as appropriate. The standing committee may revise its recommendations in response to a specific comment or series of comments submitted during this phase of the process.

As part of this process, Kaizen participants recommended that NQF membership voting should no longer be a separate 15-day voting period. NQF members would have the opportunity to express their support (*'Support'* or *'Do Not Support'*) for each measure to inform the committee's recommendations. If desired, members can change their support decision at any time during the public commenting period. This earlier and more continuous expression of support/non-support from NQF members will promote membership engagement in the endorsement process. In order to implement this proposed change, approval is required from NQF's Governance Committee and Board of Directors. Depending on the outcome of this initiative, NQF could potentially implement this recommendation at a later time.

### **Endorsement Decision**

#### Consensus Standards Approval Committee

As of early 2017, the Consensus Standards Approval Committee (CSAC) makes the final endorsement decisions on measures under review by NQF standing committees, following public and NQF Member comment and Member voting. The CSAC, a standing committee appointed by the NQF Board of Directors, serves in an advisory capacity to NQF leadership regarding enhancements to the CDP, the measure evaluation criteria, and emerging issues in performance measurement.

Kaizen participants recommended that standing committees make the final endorsement decisions, without ratification by the CSAC. Participants noted that the CSAC rarely overturns the measure recommendations of the committee. NQF appreciates the comments received on the recommendation of this endorsement body. However, given important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. Currently, the CSAC is comprised of a simple majority of consumers and purchasers. In order to ensure those two stakeholder perspectives are a key part of the endorsement process, NQF will need to make certain there is adequate representation of these groups on each standing committee. NQF is committed to implementing a plan to identify and solicit ongoing engagement and participation opportunities from these stakeholder groups. Depending on the outcome of this initiative, NQF could potentially implement this proposed change at a later time.

#### Adjudication of Appeals

Once the CSAC's endorsement decisions are made public via the NQF website, a 30-day appeals period begins. Any interested party may file an appeal on an endorsed measure with the Appeals Board during this period. The Appeals Board reviews all appeals submitted to NQF for consideration. All decisions made by the NQF Appeals Board are final.

Kaizen participants recommended that the CSAC should adjudicate all submitted appeals for endorsed and non-endorsed measures, instead of serving as the endorsement body. Implementing this recommendation would result in disbanding the Appeals Board, which was established in fall 2016. NQF appreciates the comments received on this recommendation. However, given important strategic considerations, NQF will not be able to implement a change of this magnitude at this time.

# **Enhancing Training and Education**

NQF currently provides various educational resources for stakeholders involved in the CDP. This includes virtual meetings and written materials for committee members, developers and staff. At the beginning of each CDP, NQF virtually convenes standing committees for an orientation to the CDP and an overview of the measure evaluation criteria. Prior to all committee calls and meetings, committee co-chairs meet with NQF staff to assist in anticipating questions and identifying additional information that may be useful to the committee. In addition, NQF convenes measure developers on monthly webinars to provide educational and informational updates on ongoing NQF activities. NQF also conducts bimonthly internal staff training and education sessions that focuses on the CDP.

Kaizen participants expressed a need for increased training and education for all stakeholders engaged in the CDP. NQF will expand and strengthen the current range of educational resources tailored to specific audiences and more opportunities for on-demand virtual references available for review at any time.

### Committee Co-Chairs and Members

Prior to the onset of a measure evaluation cycle, standing committee co-chairs and all other members of standing committee will receive on-boarding education and training on changes to the process and expectations on their roles and responsibilities by webinar and one-on-one conference calls, as needed. Routine meeting facilitation training conducted by an experienced NQF facilitator will be required for standing committee co-chairs to promote consistency across measure evaluation meetings. In addition, NQF will provide committee members access to electronic materials including an updated committee guidebook, recording archives and a Frequently Asked Questions (FAQs) web page containing all necessary materials essential to being an effective committee member.

### Measure Developers

NQF currently provides technical assistance to measure developers on the measure submission process through one-on-one calls and written guidance materials. NQF will continue to conduct ongoing webinars specifically targeted to developers to inform and educate them on changes to the process and information relevant at specific stages in the process. For example, prior to the initial Scientific Methods Panel review phase, NQF will conduct an in-depth tutorial of this process.

While NQF currently offers a monthly measure developer webinar, additional efforts will focus on engaging less experienced developers. NQF will host an education series by webinar (live and pre-recorded). Relevant topics will include:

- an introduction to the CDP;
- understanding the NQF measure evaluation criteria;
- best practices for measure submission;
- roles/responsibilities and expectations throughout the CDP; and
- other topics as requested.

NQF will also offer developer-focused orientation sessions that will allow developers to pose specific questions, meet NQF staff, and discuss technical assistance needs.

#### NQF Members and the Public

NQF will create a specific set of educational materials targeted to NQF membership and interested stakeholders to promote awareness and encourage more engagement throughout the process. These materials will be easily accessible and available on the NQF website. NQF staff will be available on an asneeded basis to answer questions or provide additional, one-to-one training to interested parties.

Stakeholders will have the opportunity to attend live webinars addressing the process changes and updates; this information will also be available on the NQF website that will include guidance documents applicable to promote stakeholder participation.

#### NQF Staff

Finally, NQF will work to improve consistency across projects by expanding internal educational resources offerings for staff on the process, measure evaluation criteria and meeting facilitation. NQF will implement specific staff-focused trainings on meeting facilitation conducted by internal and external trained facilitators. All staff will receive trainings on the updates and changes to the CDP. These resources will include video trainings providing an overview/refresher course on each step of the CDP; enhanced written guidance; and ongoing small group and/or one on one training, on the specific steps within the CDP. NQF will also hold biweekly education sessions on measure methodology, which will be conducted by senior staff.

### **Improvements in Information Exchange and Access**

NQF currently conducts two separate measure review processes: measurement endorsement through the CDP and input on measure use and selection through the Measure Applications Partnership (MAP). While each process has a different purpose and goal, there is significant overlap in the information submitted and produced. For MAP, brief measure specifications are provided by CMS in the form of the Measures under Consideration (MUC) list, and the MAP's final recommendations for each review year are stored in Excel files and reports on NQF's public website. For the CDP, developers provide measure specifications through NQF's online measure submission form (MSF), and the endorsement decisions and summaries of committee discussions are stored in reports on various project-specific webpages on NQF's public website. Summary information for endorsed or previously endorsed measures is included on NQF's public measure repository, the Quality Positioning System (QPS).

Kaizen participants recommended a centralized information system that would allow for a comprehensive and longitudinal view of a measure. This system would allow staff, developers, and the public to access information, including both MAP and CDP data, as the information is updated in real-time. Participants emphasized attributes such as version control, consistency between NQF projects, and the ability to easily pull and edit information as key to an ideal-state measure information repository. Kaizen participants also recommended creating a more consistent, transparent, and user-friendly tool for submitting, reviewing, and analyzing measures and comments. Lastly, participants recommended that NQF should purposefully incorporate methods to ensure the tool provides an intuitive user-friendly experience.

#### **Ongoing Improvements: Short and Long Term Solutions**

NQF will adopt a two-fold approach to addressing recommendations from Kaizen participants. Some aspects of the recommendations are resolvable through short-term solutions and adaptations of existing platforms. Other recommendations will be addressed through a long-term product development approach.

NQF will advance a short-term initiative to aggregate information by grouping MAP measure recommendations and rationale issued each year into one comprehensive and filterable document, accessible from the existing MAP homepage on the NQF website. Similarly, NQF will work to consolidate existing information from CDP reports to make it easier for users to access measure information.

NQF will also advance a short-term initiative to improve business rules around publishing timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in NQF's processes. Commenting opportunities will also be enhanced by increasing the character limit to 10,000 characters, real-time updates on comments forwarded to developers, and better regulated public comment periods during evaluation meetings.

NQF will begin to specify components and features of a centralized measure information system, for long-term implementation. This system could feature:

- comprehensive information about a measure, linking CDP and MAP evaluations in one central repository;
- alternative search tools, including measure identification or tags, to improve information accessibility;
- upgrades to facilitate user experience, particularly improving the speed of searches, and contextual information available to explain key terms.

Other considerations geared towards developer-oriented enhancements, include the usability and transmission of measure submission form content, and opportunities for "cross-talk" between major measure databases in use currently.

### **Public Comment**

NQF solicited comments on the proposed recommendations from the Kaizen via an online tool located on the NQF website. The public comment period opened on June 6, 2017 and closed on June 23, 2017. A total

of 33 organizations and individuals submitted comments (<u>Appendix A</u>), including but not limited to, consumers, purchasers, health professionals and providers. The majority of the comments focused on the Scientific Methods Panel. Specifically, commenters requested additional information about the role in relation to the standing committee, process, composition and other operational details. Overall, the comments were generally supportive of the proposed recommendations. Comments are included in Appendix A in the order in which they were received.

Finally, NQF presented the recommendations to the Consensus Standards Approval Committee (CSAC) during their June 21, 2017 monthly conference call. The CSAC was generally supportive of the majority of recommendations but requested an additional, detailed update at their July 11-12, 2017 in-person meeting. As NQF continues to plan for the implementation of this new process, NQF will solicit input from other stakeholder groups.

## **Appendix A: Comments Received**

Comment	Commenter	NQF Response
Having a scheduled submission/review process is seen as an advantage. It should be easier for organizations to plan accordingly. Thank you	Robert Dent, Midland Memorial Hospital	Thank you for your comment. We appreciate your feedback on the proposed recommendations for the CDP Redesign.
Thank you for the opportunity to comment on this important proposal. I agree with all of the proposed changes and would like to comment on one in particular. Regarding the improvements in information exchange and access, I recommend the NQF reconsider the prioritization of the proposed changes. From the perspective of end user of the quality measures, having a coordinated, centralized system for a comprehensive longitudinal view of the measure would be extremely helpful. The current process requires a significant amount of time and work effort on the end user side to pull this information together and review in a manner that is understandable and provides the ability to communicate the changes to other stakeholders. As a person that uses this process to obtain feedback from clinical staff for commenting, I believe creating a centralized system as proposed by the Kaizen participants would not only be helpful for me but would also further the engagement of outside stakeholders by making the review process less labor intense.	Joseph Kunisch, Memorial Hermann Health Care System	Thank you for your comment. NQF has identified short-term solutions to ensure our current IT infrastructure is more user friendly that progresses towards developing a more centralized system. NQF will advance a short-term initiative to aggregate information by grouping MAP measure recommendations and rationale issued each year into one comprehensive and filterable document, accessible from the existing MAP homepage on the NQF website. Similarly, NQF will work to consolidate existing information from CDP reports to make it easier for users to access measure information NQF will also strive to improve business rules around publishing timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in
the one used for commenting on the NQF's Common Formats for Patient Safety Data Document. This allows stakeholders to submit a single comment regarding a specific section of the document. I found this system to be very user friendly and significantly enhance the commenting process.		NQF's processes. Commenting opportunities will also be enhanced by increasing the character limit to 10,000 characters, real-time updates on comments forwarded to developers, and better regulated public comment periods during evaluation meetings.

Comment	Commenter	NQF Response
Comment Thank you for the opportunity to comment on NQF's planned and deferred changed. I applaud the proposed changes, particularly the addition of a methodological panel and the expansion to continuous commenting. It may be helpful for NQF to consider in future the value of providing feedback to measure stewards/developers at the Intent to Submit phase. If NQF could identify, at the intent stage, that a measure is unlikely to meet with a warm reception, it would be helpful to share this with measure developers. I am in favor of improving the measure technical reports, as these are often long and arcane. However, I urge NQF to consider test alternative formats for these reports, and not to rely solely on expert guidance on how to revise and restructure them. Much as we test measures to make sure they function as intended, it is equally important to test the efficacy of different ways of communicating measure-related information. Otherwise, we may still struggle to expand the "voices at the table" to those important stakeholders who don't wish to make a hobby of learning how to read measure reports. This need is also related to the barriers NQF notes for making a change to the endorsement decision process. Finding sufficient stakeholder perspectives for the standing committees may be more challenging without attention to their needs. Regarding the information exchange and access recommendations, I am reminded	Commenter Rikki Mangrum, American Institutes for Research	Thank you for your comment. NQF currently offers technical assistance to measure stewards and developers at any time. Stewards and developers do not have to wait until there is an active project to receive technical assistance. NQF welcomes measure stewards and developers to request technical assistance at the Intent to Submit phase. NQF has clarified the opportunity to receive technical assistance in the final report. NQF will continue to identify ways to improve the structure and format of the technical reports to capture the needs of all stakeholder perspectives. NQF has also identified short-term solutions to ensure our current IT infrastructure is more user friendly that progresses towards developing a more centralized system. NQF will advance a short-term initiative to aggregate information by grouping MAP measure recommendations and rationale issued each year into one comprehensive and
that Henry Wei of Google sat on the stage at the NQF annual conference back in April and told us all that combining data from disparate sources is now "a trivial matter." His co-panelists agreed wholeheartedly that this was true. I read about companies like Palantir that can rapidly bring together wildly different databases, including those that are entirely unstructured into dashboards that allow users to		filterable document, accessible from the existing MAP homepage on the NQF website. Similarly, NQF will work to consolidate existing information from CDP reports to make it easier for users to access measure information. Additionally, NQF will strive to improve business rules around publishing
including those that are entirely unstructured, into dashboards that allow users to access and combine data for new purposes. This makes me wonder whether a new system is truly beyond reach. Perhaps NQF has received input from the wrong experts. The problem may be the suggestion to build a big new centralized system. The answer may be in a smaller, more flexible infrastructure that allows transparent		timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in NQF's processes. Commenting opportunities will also be enhanced by

Comment	Commenter	NQF Response
interaction with decentralized systems.		increasing the character limit to 10,000 characters, real-time updates on comments forwarded to developers, and better regulated public comment periods during evaluation meetings.
We at C-TAC support the spirit and intent of the Kaizen recommendations, which are intended to facilitate efficiency and inclusive participation in the CDP. We also respect NQF's responses regarding the logistical implications for their staff and systems, and their response that some of these cannot be implemented immediately. Further, we emphasize the need for careful deliberation that leads to thoughtful implementation of key "driver measures" that encourage performance improvement but do not stifle innovation regarding how to accomplish the desired outcome. In short, the goal is to drive for the end-product of performance improvement, not dictate every step in the improvement process.	David Longnecker, Coalition to Transform Advanced Care (C-TAC)	Thank you for your comment. NQF appreciates your feedback on the proposed recommendations for the CDP Redesign.
<ul> <li>The American Academy of Family Physicians (AAFP) is in general support of the proposed changes and offers the following comments and suggestions:</li> <li><b>1) Increased opportunity for measures submission:</b> We agree with the changes. We would like a better understanding of which topic areas will be consolidated, with assurance that family medicine will continue to be represented in areas that impact primary care.</li> <li><b>2) Technical Review: Methods Panel</b>: We agree this aspect of measure evaluation is best addressed by statistical experts. We hope the change will free-up time to devote to measure alignment, duplication and identification of best-in-class measures, an important task that frequently gets less attention than it deserves in the current process.</li> <li><b>3) Measure Evaluation Technical Report:</b> We agree with the proposed changes.</li> </ul>	Sandra Pogones, American Academy of Family Physicians	Thank you for your comment. Increased opportunity for measure submissions: NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures, while distributing measures to committees with the needed expertise to conduct an evaluation. As a result, many of the smaller portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more cross-cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care.

Comment	Commenter	NQF Response
<ul> <li>4) Public Commenting Period with NQF Member Expression of Support: We support the changes.</li> <li>5) CSAC Role in Endorsement Decisions and Appeals: The AAFP agrees that standing committees are in the best position to make the final endorsement decisions. However, it is not clear if this change would require that workgroup seats previously filled by clinicians would instead be filled with consumers and purchasers. We would support adding one or two seats to workgroups for consumers and purchasers, but would oppose heavy weighting of workgroups with public members. Clinicians' professional work, payments, patient care and safety are significantly impacted by measures, and the endorsement process must remain scientific. It is also critical that enough seats be available for professional members to ensure cross-specialty evaluation, endorsement, and acceptance by the medical community. We support having CSAC function as the Appeals Board and agree with disbanding the separate appeals board. We also suggest that CSAC be more involved in identifying potential gaps in measures, an area where consumer input would be very valuable.</li> <li>6) Enhancing Training and Education: We support increased training and education for those involved in CDP and for all stakeholders. We also encourage NQF to offer</li> </ul>		<ul> <li>appreciates your suggestion on the composition of the standing committees. Given current resources and other important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. If or when this change occurs, NQF will consider your feedback on the approach.</li> <li>Enhancing Training and Education: Thank you for the suggestion. As we develop our training and education plan, we will consider your recommendation.</li> </ul>
<ul> <li>training opportunities to inexperienced professionals to help groom such professionals for measures evaluation work. We've found opportunities for this type of training are limited, and suggest that each workgroup offer a limited number of "observational" seats (1-2) to be filled by inexperienced members that wish to gain experience in the process.</li> <li>7) Improvement in Information Exchange and Access: We support eliminating duplicative information sources and centralizing information in one location,</li> </ul>		
accessible via a user-friendly tool.		
The American Academy of Family Physicians wishes to append our prior comments on the proposed Methods Review process, considering recent and important	Sandra Pogones, American	Thank you for your additional comment. Although NQF staff or the Scientific Methods Panel will review and rate the

Comment	Commenter	NQF Response
feedback we have received from our members. While we see the value of having statistical expertise available for review of reliability and validity of measures, we are concerned that non-clinicians may not be able to identify certain issues that are apparent to clinicians in their daily practice. For example, different registries or EHRs may not equally measure certain specifications due to clinical or technical features and logic, which will impact reliability. A statistician may not have identified such differences because they are not actually using the EHR and may make assumptions about commonality that do not exist. We are concerned that once a measure "passes" the hurdles for scientific acceptability and a recommendation is made to the committee, the process will become a rubber stamp approval, and due consideration of reliability and validity will not be performed by the committee. We believe there are committee members that are skilled enough to handle scientific acceptability review, although not all members may feel comfortable with this. We would not oppose having statistical experts review the measures and participate in discussion of scientific reliability with the committee, but prefer they withhold making prior recommendations. We feel it is important for all committee members to hear and participate in the full discussion of scientific acceptability, as such discussion spurs questions, enhances member understanding of the measures, and improves overall effectiveness of members in reviewing all measure criteria.	Academy of Family Physicians	reliability and validity of the measure, standing committees may raise concerns with the specifications of the measure or with potential threats to validity (e.g., selection of variables for risk adjustment model) and can therefore overturn the staff or Scientific Methods Panel rating. As part of its ongoing education efforts, NQF will provide clear guidance to standing committees regarding the circumstances wherein an overturn of the rating would be permissible.
The American Academy of Neurology (AAN) an association of more than 28,000 neurologists and neuroscience professionals appreciates the opportunity to comment on the 2017 Kaizen Consensus Development Process. The AAN is grateful of NQF's efforts to improve the Consensus Development Process (CDP). Several questions arise from the NQF's plan to limit the number of measures to be reviewed twice yearly. It is anticipated that for many standing committees more than eight new measures will be submitted in a year. How will NQF prioritize measures in this	Amy Bennett, American Academy of Neurology	NQF appreciates your comment. NQF will prioritize measures based on the measure maintenance schedule and the submissions of the Intent to Submit forms. NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures,

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situation, and how will NQF ensure endorsement review occurs in a timely manner?		while distributing measures to committees with the needed
How will the 22 current topical areas be reduced to 16, and will the public have input		expertise to conduct an evaluation. As a result, many of the
on these future groupings? It is anticipated that there will be situations where		smaller portfolios have been consolidated into cross-cutting
measure developers disagree with the NQF staff or external methods panel		topics with a broader range of experience. In addition, some
determination of low or insufficient ratings. What recourse is available when a		clinical groupings of committees were made to reflect more
developer disagrees? Will there be an appeal process through the external methods		cross-cutting clinical areas, such as primary care and chronic
panel or the standing committee?		illness care, pediatrics and geriatrics and palliative care. NQF
		will not formally open a public comment period on the
The AAN is concerned the open comment period could result in confusion, and		consolidated list of topical areas. However, NQF welcomes
potentially standing committee members would not receive or review comments		your feedback on the list. If you have input, feel free to email
within the meaningful timeframe for action on the comments. The AAN notes there		NQF at NQFKaizen@qualityforum.org.
is a need for increased training and education, but there is little no discussion on		
how NQF will evaluate the effectiveness of training. The NQF may benefit from		The process includes a two-week process for measure
analyzing the effectiveness of standing committees, and developing a plan to		steward or developer to respond to the ratings. In addition,
address situations when a standing committee is not operating efficiently (e.g., poor		the standing committees can still discuss relevant issues,
direction from committee chairs, questions to NQF staff are unanswered, etc.). The		such as risk adjustment. Given the opportunity for more
AAN would also encourage NQF to make improvements in the information exchange		frequent submission, measures may need to be move to the
and access. Submission to NQF is an arduous process taking no less than 40 hours for		next review cycle to address methodologic concerns.
one submission in a large part due to lack of smart forms and the required		
resubmission of duplicative information.		Standing committees will receive all comments submitted
		during the measure review process. Comments submitted up
		to one week prior to the committee evaluation meeting will
		be included in the meeting materials for discussion during
		the evaluation meeting. All comments submitted after the
		evaluation meeting through the end of the public comment
		period will be included into the meeting materials for the
		committee discussion on the post-comment call.
		NQF currently surveys standing committee members on
		their experience and solicits feedback on ways to improve

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		their involvement in the CDP. NQF intends to assess the effectiveness of the education and training program. NQF is working to identify solutions to enhance our current IT infrastructure to provide a more user-friendly experience when submitting a measure for endorsement consideration.
Thank you for the opportunity to comment on the National Quality Forum's proposed changes. The National Viral Hepatitis Roundtable (NVHR) is a coalition of approximately 500 member organizations working to fight, and ultimately end, the hepatitis B and C epidemics in the United States. NVHR believes this goal can be achieved by addressing stigma and health disparities, removing barriers to prevention, care, and treatment, and ensuring respect and compassion for all affected communities. We would like to express concern with the proposal to only consider NQF member input on measures under consideration. We think it is important that stakeholders with subject matter expertise continue to be allowed to provide input and feedback for measures under consideration. We would like to encourage NQF to continue to seek input and comments from all relevant stakeholders and not just those who	Ryan Clary, National Viral Hepatitis Roundtable	Thank you for your comment. All stakeholders, regardless if the individual or organization is an NQF member, can submit comments and feedback on the measures during the 12- week comment period as well as provide comments during the committee evaluation meetings. The option to express support or non-support for the measures under consideration would only be limited to NQF members.
have paid membership dues to NQF. Recommendation: NQF should expand input on measures under consideration to the general public.		
NQF's role in facilitating the Measure Applications Partnership (MAP) is to serve as a voluntary consensus standards body, which requires openness to stakeholder participation and input in line with Circular A-119 (please see note with additional details below). While NQF's proposal to only consider NQF member input on measures under consideration may pass a low bar for "openness," we generally		

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believe that th	e openness criterion was not intended to mean receptivity to input		
from just those	e stakeholders who paid membership dues to NQF.		
Note: OMB Cir	cular A-119 defines voluntary consensus standards bodies as		
"domestic or in	nternational organizations which plan, develop, establish, or		
coordinate vol	untary consensus standards using agreed-upon procedures. For		
purposes of th	is Circular, "voluntary, private sector, consensus standards bodies," as		
cited in Act, is	an equivalent term. The Act and the Circular encourage the		
participation o	f federal representatives in these bodies to increase the likelihood		
that the standa	ards they develop will meet both public and private sector needs. A		
voluntary cons	ensus standards body is defined by the following attributes:		
(i)	Openness.		
(ii)	Balance of interest.		
(iii)	Due process.		
(iv)	An appeals process.		
(v)	Consensus, which is defined as general agreement, but not		
	necessarily unanimity, and includes a process for attempting to		
	resolve objections by interested parties, as long as all comments		
	have been fairly considered, each objector is advised of the		
	disposition of his or her objection(s) and the reasons why, and the		
	consensus body members are given an opportunity to change their		
	votes after reviewing the comments.		
On behalf of th	e American Society of Nephrology (ASN), thank you for the	Eleanor Lederer,	Thank you for your comments.
opportunity to	provide comment on the National Quality Forum (NQF) 2017 Kaizen	American Society	
Consensus Dev	elopment Process Proposed Redesign draft report. ASN represents	of Nephrology	Technical Review: Methods Panel: Because updated
nearly 17,000	physicians, scientists, nurses, and other health professionals dedicated		reliability and validity testing is not required for maintenance
to treating and	studying kidney diseases to improve the lives of people with kidney		measures, NQF staff will review previous testing results for
diseases. ASN	s a not-for-profit organization dedicated to promoting excellence in		complex maintenance measures and attest to the adequacy
			of prior testing. If prior testing is inadequate, updated

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kidney care and ensuring access to optimal patient-centered quality care, regardless of socioeconomic status, geographic location, or demographic characteristics. ASN appreciates the opportunity to provide public comment on renal measures under consideration and regarding the annual Measures Application Partnership (MAP) process. The society values NQF's efforts to enhance healthcare value, make patient care safer, and achieve better outcomes, and commends NQF for undertaking this redesign initiative at this time. In general, ASN is supportive of the proposed changes outlined in the draft report, particularly efforts to make the consensus development cycle more rapid. The proposal suggests a greater role for the NQF staff in guiding the measure consideration and endorsement process. ASN supports this shift, and believes more engagement and leadership roles by the staff will benefit the organization and help it achieve its goal of more efficient processes. The society offers a few additional comments and questions for consideration that we hope are helpful as NQF finalizes and implements the report.		testing is provided, or NQF staff deems an external review necessary, the measure will be submitted to the external Scientific Methods Panel to evaluate the reliability and validity of the measure. The standing committees will not determine which measures will be sent to the Scientific Methods Panel. Upon submission of the Intent to Submit form, NQF will assess whether the measure will be reviewed by the Scientific Methods Panel. The process includes a two-week process for measure steward or developer to respond to the ratings. In addition, the standing committees can still discuss relevant issues, such as risk adjustment. Given the opportunity for more frequent submission, measures may need to be move to the next review cycle to address methodologic concerns.
<ul> <li><u>Technical Review: Methods Panel</u></li> <li>ASN applauds the proposal to create a separate technical advisory panel tasked with conducting methodological reviews of complex measures. This change would have several benefits, including ensuring a group of experts in this complex arena have a dedicated mission of assessing aspects that may not receive the optimal amount of attention or expertise they warrant in the current system. Additionally, it may create more consistency in the statistical validity of all measures across the NQF portfolio. In addition to having the Methods Panel assess the measures that the NQF staff categorize as "complex measures", ASN would also recommend that the Panel assess:</li> <li>All complex measures undergoing maintenance review for which there are performance data and/or when there are existing, new or updated testing</li> </ul>		Public Commenting Period with NQF Member Expression of Support: Thank you for your comment. NQF will continue to make every effort to lengthen the public commenting period on the measures under consideration (MUC) list. However, this is contingent on the release of the MUC by CMS. Endorsement Decision: Thank you for the suggestion. As we develop our training and education plan, we will consider your recommendation.

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data.		
• Any measure for which a standing committee member moves to request a		
review by the methodology panel.		
ASN also suggests that NQF develop a process to adjudicate situations where		
measure developers may disagree with the determination by NQF staff or external		
methods panel regarding low or insufficient ratings.		
Public Commenting Period with NQF Member Expression of Support		
NQF's proposal to create one continuous comment period directly addresses an area		
of concern for ASN, and the society strongly supports this recommendation.		
Providing a longer period for public comment will both allow more stakeholders to		
share input and to ensure that commenters have adequate time to consider the		
often very complex and technical issues that are under consideration—thereby		
increasing the value of the feedback.		
Related to this positive change, ASN would also encourage NQF to provide more		
time for public comment regarding the annual MAP process. The society recognizes		
that NQF is also working with other stakeholders (such as the Department of Health		
and Human Services) and thus the timeline may not be fully within NQF's control,		
but anything that the organization can do to lengthen the amount of time for public		
comment on the MAP would make it possible to provide more thoughtful,		
meaningful input.		
Endorsement Decision		
ASN concurs with the draft report recommendation that NQF not prioritize efforts to		
switch final endorsement decisions from the Consensus Standards Approval		
Committee (CSAC) to the standing committees. Many other changes, outlined in this		
report, would have higher value and it is important to get those modifications right.		
In the future, while ASN supports the concept of encouraging participation of more		
patient and consumer voices, the society would have concerns about keeping the		

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level of scientific discourse at an appropriately expert level. In general, the society		
would encourage NQF to consider providing more advance training and education		
about some of the more technical aspects of measure development to the lay		
persons participating in the CSAC (or any other NQF panels) in advance of the		
meetings to maximize their ability to contribute.		
Enhancing Training and Education		
ASN applauds the proposals to raise awareness about NQF's current training and		
education opportunities and to expand those efforts in the future. Offering		
developer-focused sessions that would allow developers—or those considering		
entering the measure development arena— to talk with NQF experts in an informal		
setting to pose questions and discuss opportunities and challenges about their area		
of interest would be of immense value.		
Again, thank you for the opportunity to provide comment on the draft report. ASN		
would be pleased to discuss these comments with NQF if it would be helpful. To		
discuss ASN's comments, please contact ASN Director of Policy and Government		
Affairs Rachel Meyer at (202) 640-4659 or at <u>rmeyer@asn-online.org</u> .		
Overall, the summary document is comprehensive and the recommendations are	Anne Leddy,	Thank you for your comment. NQF appreciates your
well conceived.	Member,	feedback on the proposed recommendations for the CDP
	Endocrine	Redesign.
#1. Increased opportunities for measure submission – excellent having two measure	Standing	
submission cycles per year. I support limiting the number of measures to 12 per	Committee	
cycle, 8 being routine maintenance of endorsed measures, and 4 new measures.		
Reduction of topical areas to 16 also good. Also encourage that active standing		
committees have not only virtual web meetings but in-person meeting yearly. I		
understand there is limited funding.		
#2. Intent to submit requirement for new measures – I support this. Should facilitate		

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consideration of new measures		
#3. Technical review: Methods Panel - Excellent recommendation. I am a clinician. I		
have a rudimentary grasp of statistics. I have spent countless hours boning up on the		
methods needed to review the material essential to determining reliability and		
validity. Would suggest that the "Methods Panel" provide a brief synopsis of their		
statistical review of each proposed measure and that be provided with other usual		
material for the standing committees. This is truly a "value add".		
#4. Measure evaluation technical report – Encourage that these changes be		
implemented as soon as possible.		
#5. Public Commenting Period – Implement as soon as feasible.		
#6. Endorsement process – I approve of the suggestion that the standing committees		
make final endorsement decisions without ratification by the CSAC. I also		
understand that it will take time to bolster the membership of the standing		
committees with more representation by consumers and producers so their		
stakeholder perspectives remain part of the endorsement process.		
#7. Adjudication of Appeals – I support tasking the CSAC with adjudicating appeals		
rather than being the endorsement body. Having a separate Appeals Board is		
redundant.		
#8. Enhancing Training and Education – Bravo. This is absolutely essential. I am sure		
that the NQF will work toward providing these resources within the limits of the		
budget.		
Preface	John Bott,	Thank you for your comment.
The following are comments from Consumer Reports regarding the NQF draft report	Consumer	

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titled "2017 Kaizen Consensus Development Process: Proposed Redesign", which	Reports	Technical Review: Scientific Methods Panel:
appears at the following link:		Maintenance measures can be complex measures. Because
http://www.qualityforum.org/2017 Kaizen Comment.aspx		updated reliability and validity testing is not required for
If you have any questions about these comments please contact John Bott at		maintenance measures, NQF staff will review previous
jbott@consumer.org, Doris Peter at <u>dpeter@consumer.org</u> , or Lisa McGiffert at		testing results for complex maintenance measures and attest
Imcgiffert@consumer.org.		to the adequacy of prior testing. If prior testing is
Technical Review: Methods Panel		inadequate, updated testing is provided, or NQF staff deems
To summarize the draft NQF report a key component of the external methods panel		an external review necessary, the measure will be submitted
(on page 4, last paragraph), it appears to propose the following for the defining and		to the external Scientific Methods Panel to evaluate the
processing three types of measures:		reliability and validity of the measure.
[1] Newly submitted: Complex measures		Due to volume and capacity concerns, all submitted
Measure types: risk adjusted outcomes, composites, cost		measures cannot be reviewed by the Scientific Methods
Body: external methods panel		Panel. NQF staff has the appropriate expertise to review the
Charge: provide recommendation to standing committees		non-complex measures. NQF will train and provide resources
		to the Scientific Methods Panel to ensure consistency in
[2] Newly submitted: Non-complex measures		applying the testing information submitted to the measure
Measure types: such as process and structural measures		evaluation criteria.
Body: NQF staff		
Charge: provide recommendation to standing committees		Upon submission of the Intent to Submit form, NQF will
		assess whether the measure will be reviewed by the
[3] Currently endorsed: Measure maintenance		Scientific Methods Panel. No matter the classification of the
Measure types: endorsed measures reviewed in maintenance		measure (complex or non-complex), the review by NQF staff
Body: NQF staff		or the Scientific Methods Panel will not add additional time
Charge: attest to adequacy of prior testing		to the review process. All measures that are ready for
		committee review, will be sent to the committee with
Note regarding the above: Because measures in maintenance (i.e. #3) are called out		adequate time for the committee to review prior to the
separately from complex and non-complex measures (i.e. #1 and #2 respectively) as		committee evaluation meeting.
to use of an external methods panel, it appears that the references to complex and		In the final report, NQF has clarified the definition of a

Comment	Commenter	NQF Response
non-complex measures are in regard to newly submitted measures. Consumer Reports provides the following recommended changes, recommended attributes and concluding comments: Recommended changes We recommend to use an external panel for measures noted above in #1, #2 and #3 above (vs. NQF staff for some measures and an external panel for others). Rationale: Having a set of NQF staff evaluate some measures for Scientific Acceptability while an external methods panel evaluates other measures increases the likelihood of using differing standards to vet the measures against and a differing bar that the measures must meet to be deemed acceptable. The above #3 is silent on the process for currently endorsed measures with changes to the measure as it relates to the Scientific Acceptability differ for maintenance measures, staff would review testing results for maintenance measures and attest to the adequacy of prior testing." What occurs with measures where substantial changes are made, and thus pointing to the prior testing is irrelevant? Is the proposal that measures that are largely unchanged (and the measure steward attests to adequacy of prior testing) are reviewed by NQF staff, and measures that substantially change are reviewed by the external methods panel? If NQF adopt a framework where measures in maintenance with changes are channeled to the external methods panel, and measures without changes go to NQF staff, steps are added to the process to review and sort the measures to channel them accordingly. Such added steps have the		<ul> <li>complex measure. The following types of measures will be considered complex and therefore may require an evaluation by the Scientific Methods Panel:</li> <li>Outcome measures, including intermediate clinical outcomes <ul> <li>Instrument-based measures (e.g., PRO-PMs)</li> <li>Cost/resource use measures</li> <li>Efficiency measures (those combining concepts of resource use and quality)</li> <li>Composite measures</li> </ul> </li> <li>Additionally, NQF has also provided additional information on the composition and disclosure of interest of the Scientific Methods Panel in the final report. The new NQF Scientific Methods Panel will consist of 15 to 25 nominated statisticians, epidemiologists, psychometricians, economists, performance measure methodologists and individuals with expertise related to eMeasures and disparities. All nominees will complete an annual general disclosure of interest (DOI) form, as well as measure-specific disclosures to identify recusals from specific measures. NQF will assign measure review based on identified conflicts of interest, relevant expertise, and availability of panel members. All reviews provided by the Scientific Methods Panel will be shared not only with the committee but also with the steward/developer and the public. Furthermore, the Scientific Methods Panel's charge will include providing</li> </ul>
consequence of: a) adding time, b) consuming resources and c) creating the		expertise for methods/testing-related issues for NQF and

Comment	Commenter	NQF Response
opportunity of inappropriately sending measures to the incorrect group.		advance NQF's guidance on these issues.
The definitions of the #1 and #2 groupings are: a) not mutually exclusive, b) unclear, and c) not encompassing of all measures. For example, there are risk adjusted process measures and there are outcome measures that are not risk adjusted. Sorting measures between "complex" and "non-complex" (where some measures go to NQF staff and others to the external methods panel) will add steps the process. In turn, this consumes resources and time, and increases the likelihood of misclassifying measures. Thus, the result is inappropriately sending some measures to down the "complex" path and "non-complex" path. As noted above regarding #3, use of an external methods panel streamlines the review process as all measures are channeled to the panel. The result is reducing waste, which is a principle of Kaizen, as well as standardizing the process for evaluating scientific information.		Enhancing Training and Education: Improving the training and education for the standing committee members will assist in ensuring consistency across all 15 committees when applying the measure evaluation criteria. Endorsement Decision: NQF appreciates your suggestion on the composition of the standing committees. Given current resources and other important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. If or when this change occurs, we will consider your feedback on the approach. Increased staff training and education will further ensure NQF procedures are adhered to during measure evaluation
<ul> <li>Recommended attributes</li> <li>The following are noted as recommended attributes vs. changes for the draft report is silent on a number of aspects of the proposed methods panel. The following are attributes we recommend that are used in building the framework for the external methods panel: <ul> <li>a) The external methods panel has a majority of consumers and purchasers.</li> <li>b) The external methods panel meetings are open to the public, the same as standing committees (SC) are. This process needs to be transparent; transparency is another reason why NQF staff should not be making any decisions re measures (e.g., #3 above)</li> <li>c) The external methods panel members are subject to review for conflict of interests. As a result of this review, qualifying panel members will be free from conflicts of interest.</li> </ul> </li> </ul>		process. Furthermore, the CSAC also provides oversight on the consensus development process and performs a final review of the process prior to making an endorsement decision.

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d) NQF will vet the external methods panel nominations for sufficient		
competencies in the areas evaluated in the Scientific Acceptability criterion.		
e) Draw on the recent National Academy of Medicine (NAM) report titled		
"Vital Directions for Health and Health Care" as it relates to the concept of		
creating a health care performance measurement parallel to the Financial		
Accounting Standards Board (FASB). Related to the FASB concept, we		
suggest to charge the external methods panel with responsibilities aimed at		
improving the standardization of measure review for Scientific		
Acceptability. For example, the panel's scope could state they are charged		
with standardizing across all measures being evaluated:		
<ul> <li>How measures are reviewed for Scientific Acceptability;</li> </ul>		
The acceptable minimum threshold a measure must pass for		
Scientific Acceptability		
Concluding comments		
As it currently stands, 16 NQF SCs are reviewing measures as to Scientific		
Acceptability. Needless to say, such a large number of bodies evaluating measures		
against this criterion increases the likelihood of inconsistent standards applied to		
measures being reviewed. This risk for inconsistency is exacerbated by the following		
existing NQF attributes, which are noted in the NAM's "Vital Directions" report:		
<ul> <li>NQF criteria are not evaluated in a strict quantitative sense;</li> </ul>		
<ul> <li>The NQF does not define specific validity tests for different types of measures;</li> </ul>		
<ul> <li>NQF does not require a minimum bar for reporting a measure's validity and reliability;</li> </ul>		
<ul> <li>NQF does not define specific thresholds for validity and reliability for endorsement</li> </ul>		
The "Vital Directions" report goes onto provide recommendations that relate to this		

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particular proposal in the draft NQF report, and suggests the potential for NQF to be part of the solution. Specifically, "Vital Directions" notes: "Policy-makers could create an inde-pendent body to write standards for healthcare performance measures One option would be to build on NQF The entity charged with this work ideally would be a private, nongovernment self-regulating organization"		
Through this recommendation for one external methods panel on Scientific Acceptability, NQF can move a step closer to the NAM's vision for NQF.		
Endorsement Decision The NQF draft report indicates the Kaizen process recommended that: "standing committees should make the final endorsement decisions, without ratification by the CSAC."		
Consumer Reports generally supports the above stated proposal; however we strongly agree that final endorsement decisions should not move to the SCs until the membership of each SC is reconstituted to have a simple majority of consumers/purchasers.		
<u>NQF staff ensuring procedures are adhered to during SC process</u> A current role of the CSAC is to ensure the proper protocols were adhered to in the review and voting on the measures by the SC. Such review of appropriate adherence to procedures should occur concurrently during each step of the endorsement process. Thus, NQF staff should fulfill this role during the SC's work as well as the work of the external methods panel.		
Rationale: Putting the procedural review on the back end of the endorsement process is illogical. Identifying if there were procedural issues after the fact is		

Comment	Commenter	NQF Response
inefficient and runs counter to a stated Kaizen process. Specifically, the NQF draft reports notes one of the aims of changes to the endorsement process is: <i>"reducing cycle time of the CDP"</i> . <u>Improvement in Information Exchange and Access</u> We strongly agree with the recommendations of the Kaizen participants to create <i>"a centralized information system that would allow for a comprehensive and longitudinal view of a measure. This system would allow staff, developers, and the public to access information, including both MAP and CDP data, as the information is updated in real-time."</i> We encourage NQF to move toward such improvements in the future.		
We thank the National Quality Forum (NQF) for the opportunity to comment on the NQF Kaizen Draft Report. First, we note that a key process change discussed at the Kaizen event is not reflected in the current draft report. To avoid steering committees wasting time on applications that are unclear or for which the NQF staff and developer views on the technical content differ, NQF staff and the developer will reach agreement on the application of NQF criteria to the measure (or articulate differences where necessary) <u>before</u> the measure goes to the steering committee. This would ensure that the developer does not have to separately rebut NQF staff application of NQF guidelines post-hoc in front of the committee where there are differences. Second, we provide comments on the NQF Kaizen Draft Report in the table below.	Anouk Lloren, Yale-CORE	We appreciate your feedback on the recommendations. NQF has incorporated the steward/developer review process of the preliminary analysis of the measure in the final report. Two measure submissions per year: The timing of review for the maintenance measures will depend on when the measures are scheduled to undergo maintenance. NQF will not prioritize measures based on the type of measure evaluation meeting. NQF will limit 12 measures per topical area to regulate increased workload for the standing committees (and may
<ul> <li><u>Two measure submissions per year; 6 months each</u></li> <li>We generally support this approach but would like to better understand: <ul> <li>How will NQF prioritize which measures will go to the in-person vs. webinar session?</li> <li>What will NQF do if the submitted number of measures exceeds 8</li> </ul> </li> </ul>		include one or two measures as deemed appriopate). The combination of maintenance and new measures may vary depending on number of measures submitted, opportunities for related and competing measure review, and measure prioritization. Any Intent to Submit forms that are submitted once capacity has been reached for a particular cycle, NQF

Comment	Commenter	NQF Response
maintenance and 4 new measures?		will notify the steward/developer and provide the date of
• Our preference would be an approach that accommodates all submitted measures in timely way.		the next cycle in which there is availability. Intent to Submit: Steward/developers can submit prior
Developer signals intent to submit Requiring a developer to signal intent two months before submission deadline (3 months before review starts) is reasonable For maintenance measures (i.e. measures that are currently NQF-endorsed), the		testing data for maintenance measures as long as it continues to meet the measure evaluation criteria. However, if the steward/developer re-tested the maintenance measure, it is expected that the steward/developer would provide the updated testing data.
report indicates that "measures must indicate if new testing data will be available." We request clarification on what NQF means by "new testing data." It is most helpful when NQF describes this in terms of the specific sections of the submission or testing forms that, if updated, require notification to NQF staff.		Technical Review: Scientific Methods Panel The Scientific Methods Panel will review all measures deemed as complex. The following types of measures will be considered complex and therefore may require an evaluation by the Scientific Methods Panel:
The draft report does not reflect that, at the Kaizen, we mapped out that NQF staff would review submissions and rate them against criteria prior to NQF's applications going to committees. Specifically, we discussed to have any disagreements adjudicated and resolved before the measure moves forward to the committee, so that the staff-developer differences are adjudicated in advance of the committee instead of in front of the committee. This is a critical step that would address a pain point, so it needs to be made explicit.		<ul> <li>Outcome measures, including intermediate clinical outcomes</li> <li>Instrument-based measures (e.g., PRO-PMs)</li> <li>Cost/resource use measures</li> <li>Efficiency measures (those combining concepts of resource use and quality)</li> <li>Composite measures</li> </ul>
<u>New methodological panel</u> The draft captures the Kaizen conclusions fairly well, but it is unclear on the scope of the methods panel's review. Will the methodological panel always review reliability/validity as well as risk adjustment modeling for "complex measures?" If so, we support this approach.		Additionally, NQF has provided information on the Scientific Methods Panel composition and disclosure of interest process in the final report. The new NQF Scientific Methods Panel will consist of 15 to 25 nominated statisticians, epidemiologists, psychometricians, economists,
We recommend more clarity about the type of experts NQF will seek for the		performance measure methodologists and individuals with expertise related to eMeasures and disparities. All nominees

Comment	Commenter	NQF Response
method's panel and how that panel will reach decisions about recommendations to		will complete an annual general disclosure of interest (DOI)
the committee assuming that all conclusions will not be unanimous.		form, as well as measure-specific disclosures to identify
		recusals from specific measures. NQF will assign measure
Also, the report indicates that "standing committees may raise concerns with the		review based on identified conflicts of interest, relevant
specifications of the measure or with potential threats to validity (e.g., selection of		expertise, and availability of panel members. All reviews
variables for risk adjustment model)." We assume the intention of this statement is		provided by the Scientific Methods Panel will be shared not
to make clear that clinical and other content experts on the committee, who may		only with the committee but also with the
not be methodology experts, should be able to raise concerns about appropriateness		steward/developer and the public. Furthermore, the
of risk variables, cohort definitions, and the like. However, we recommend that		Scientific Methods Panel's charge will include providing
committees receive clear guidance for overturning the recommendations of the		expertise for methods/testing-related issues for NQF and
methods panel and that someone from the panel be available during committee		advance NQF's guidance on these issues.
discussions and voting.		
		Continuous commenting: NQF has provided clarification
Continuous commenting		regarding the developer's role in responding to comments
As under the current process, developers should have an opportunity to respond to		during the commenting period. NQF will ensure the measure
comments. It is unclear how NQF will manage this input in a continuous comment		developer receives the submitted committees in order to
process.		prepare for the measure evaluation meeting. Measure
		developers will not be required to provide written responses
It will be important for the developers to also receive all comments submitted one		to the comments received prior to the measure evaluation
week prior to the in-person meeting so that they will be prepared to discuss the		meeting. The committee will review any comments received
comments at the meeting. We recommend that NQF not require developers to		after the committee evaluation meeting during the post-
prepare written responses to comments prior to the in-person meeting as this short		commenting period call. All submitted comments during this
response window would put an undue burden on the developers.		time will receive written responses from the standing
		committee, measure developers, and/or NQF, as
We also assume, although not explicitly stated, that developers would receive		appropriate.
comments at the close of the period (30 days after the posting of the committee		
report) and have some period of time to prepare written responses for the		Endorsement Decision: Given current resources and other
committee's consideration as is the current practice.		important strategic considerations, NQF will not be able to
		implement a change of this magnitude at this time.

Comment	Commenter	NQF Response
Simplifying technical report		
We support this but only if it is accompanied by better on-going public access of the		
findings of NQF's review of measures and the specifications of endorsed measures		
(i.e., availability of materials on NQF's website).		
Steering committees make final endorsement decision rather than CSAC;		
CSAC adjudicates appeals rather than Board		
We appreciate the effort to streamline processes and recognize that the CSAC often		
just follows the recommendations of the steering committee. However, we feel it is		
very valuable to have a single standing body with experience and diverse members		
confirm committee decisions. This is an opportunity to ensure some consistency in		
approach and guidance to committees from stakeholders. The steering committees		
have less experience with endorsement processes and therefore this change could		
lead to even more inconsistent results from committee to committee. We have		
experienced adjudication from the CSAC of issues that helped to clarify and		
standardized approaches across committees. We think the CSAC role as a central		
endorsement committee remains very valuable.		
Enhancing training and education		
Support		
Improvements in information exchange and access		
Support		
The Federation of American Hospitals ("FAH") appreciates the opportunity to	Jayne Hart	Thank you for your comment. We appreciate your feedback
comment on the proposed changes to the consensus Development Process (CDP).	Chambers,	on the proposed recommendations for the CDP Redesign.
We are encouraged that the National Quality Forum continues to work to improve	Federation of	NQF hosted the Kaizen event in collaboration with CMS to
the CDP, but we do not believe that the proposal in its current form provides	American	inform the CDP redesign. CMS, as the funder of this
sufficient detail to all us to completely understand the changes and their	Hospitals	initiative, has asked NQF to solicit public comment on the
implications. The FAH also is concerned that a two-week commenting period does		proposed recommendations and provide a final report

Comment	Commenter	NQF Response
not permit adequate time for input from the NQF membership.		outlining the new CDP by July 1, 2017. Thus, NQF had to limit the amount of time NQF members and the public had to
The following are initial questions about the various changes, particularly to ensure		provide feedback. However, as NQF continues to plan for
that the changes are consistent with the National technology Transfer and		implementation of the new CDP, additional feedback is
Advancement Act of 1995 and the Office of Management and Budget (OMB) Circular		welcomed.
A-119 as updated in October 2012 (12). In addition, the FAH believes that any		
improvements made to the CDP must further reinforce the Standing Committees'		Overarching questions and comments: To allow more
and the NQF staff's ability to easily and transparently ensure the evaluation process		frequent measure submissions, committees will convene
and criteria are applied consistently across and within projects.		more often. Additionally, there will be increased
		opportunities for NQF membership to engage in the process.
Overarching questions and comments		As a result, NQF emphasizes the importance of stakeholder
What is the impact and expected scope of the proposed increase in the number of		education.
topics to be considered twice a year. From the proposal, it is unclear what impact		
this would have for membership and committees. In any one year, how many		After the standing committee completes its measure review,
additional measures and reports might be released for member and public		a summary of the committee's recommendations – or "draft
comment? How does this compare to the current public comment periods,		report" – will be posted on the NQF website for the public
measures and projects?		and NQF membership to review and comment. Because
		there are more review cycles, NQF will revise the content
To what degree will the measure submissions and comment periods be staggered?		and structure of the report to highlight key elements of
A graphic that depicts what the revised CDP would look like once all the proposed		interest. These elements are included in the final report.
changes are implemented would be helpful. It is difficult to track projects in the		
current process. How will the new process make it easier to be thoughtful		To assist in planning and minimize burden for the measure
participants? Will the underlying computer systems be enhanced to support faster		stewards/developer, committee members and NQF,
and more accurate searches? How will NQF members be able to easily track the		submission deadlines will be staggered. A graphic that
projects? The final document needs to pull together more clearly every proposal so		outlines the new process is included in the final report.
that members can see the entire process from start to finish.		
		Scheduling/Frequency: The process includes a two-week
Scheduling/Frequency		process for measure steward or developer to respond to the
The FAH supports NQF providing an opportunity to allow more frequent submissions		ratings. In addition, the standing committees can still discuss
		relevant issues, such as risk adjustment. Given the
Comment	Commenter	NQF Response
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and reviews. The proposed approach still uses a "batching" methodology that will force decisions on measures within a certain timeframe and project. This process is fine for measures that are well developed and with little controversy. However, the proposed process till does not permit an iterative process where disagreements and concerns can be addressed before a final decision is made. The FAH believes there are way to create a process where measures can enter at any time, go to a committee for review when they are deemed ready and then go through multiple comment cycles or committee reviews until there is general agreement that the measure is ready to be endorsed. While the twice-yearly submission process meets some of those needs, it does not address the true need for the ability to submit and review a measure at any time nor the ability to achieve true consensus. • The proposed change states that 22 topic areas are merged/reduced to 16, but additional detail is not provided. What are the 16 topic areas? How does this impact the number and composition of the Standing Committees, particularly for topics that are retired and the ones that are combined? How does this merging impact the limit of 12 measures per cycle since the number of measures in some topic areas may increase with the proposed	Commenter	opportunity for more frequent submission, measures may need to be move to the next review cycle to address methodologic concerns. The consolidated topic areas are included in the final report. For larger topic areas that include multiple conditions or cross-cutting areas, NQF will utilize technical experts as needed. NQF will offer two measure submission opportunities for each topic area each year, limiting the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This was determined given that approximately 80% of the measures submitted for endorsement consideration are maintenance measures. The combination of maintenance and new
		measures may vary depending on number of measures submitted, opportunities for related and competing measur review, and measure prioritization efforts. Per NQF's maintenance of endorsement policy, measures are due for reassessment every three years.
<ul> <li>to submit" requirement below?</li> <li>How will NQF prioritize which measures will be reviewed if there are more than four new measures in a cycle? Would it be first submitted or some other criterion? The FAH notes that prioritization must be balanced and not be partial to any one group (e.g., federal agencies vs. private sector developers).</li> </ul>		Intent to Submit: NQF will remind measure stewards and developers of scheduled measure maintenance review several months prior to the review and notify each of their assigned review cycle. Technical review: Methods Panel: NQF has provided additional information on the composition and disclosure of

Comment	Commenter	NQF Response
		interest of the Methods Panel in the final report. The new
ntent to submit		NQF Scientific Methods Panel will consist of 15 to 25
How will this step be used for maintenance measures? Will NQF notify		statisticians, epidemiologists, psychometricians, economists,
developers when maintenance is due? As drafted, it appears that the onus		performance measure methodologists and individuals with
is on the developer regardless of whether the measure is new or		expertise related to eMeasures and disparities. All nominees
maintenance.		will complete an annual general disclosure of interest (DOI)
• This section of the proposal also does not describe what NQF will do if it		form, as well as measure-specific disclosures to identify
receives more than 12 measures in a review cycle? How does NQF decide		recusals from specific measures. NQF will assign measure
which measures get moved to the next cycle? How will NQF prioritize		review based on identified conflicts of interest, relevant
which measures will be reviewed in a given cycle? How quickly will NQF		expertise, and availability of panel members. All reviews
inform developers if their measure(s) cannot be reviewed (e.g. a month		provided by the Scientific Methods Panel will be shared not
before the submission deadline, after the submission deadline.)?		only with the committee but also with the
		steward/developer and the public. Furthermore, the
Technical review: Methods panel		Scientific Methods Panel's charge will include providing
The creation of this panel has the potential to strengthen the evaluations for		expertise for methods/testing-related issues for NQF and
scientific acceptability and may be able to improve consistency across measures and		advance NQF's guidance on these issues.
projects. The FAH has several questions around the panel itself, the roles and		NQF will provide standard guidance on assessing the
responsibilities of this panel, NQF staff and Standing Committees, and the actual		Scientific Acceptability criterion for a measure, using the
process and guidelines around these reviews.		current decision algorithm used from the measure
Who would be on the panel? Will it function similarly to a Standing		evaluation criteria. As part of its ongoing education efforts,
Committee with terms? What level of education/training will be provided		NQF will provide clear guidance to standing committees
to its members if needed? If these questions are not answered up front, the		regarding the circumstances wherein an overturn of the
same challenges experienced with the current Standing Committee		rating would be permissible.
members reporting difficulty in completing their reviews may still occur.		
Why does the creation of this panel allow for additional participation by		For both complex and non-complex measures, when the
consumers, patients, and purchasers? This statement implies that		preliminary analysis is complete, NQF staff will send the
determinations of reliability and validity of measures are moved from the		preliminary analysis to developers by email for review. The
committee to NQF staff. This seems contrary to a process that is designed		process includes a two-week process for measure steward or
to achieve consensus across stakeholder groups. What is the process for		developer to respond to the ratings. In addition, the standing

Comment	Commenter	NQF Response
<ul> <li>assuring the scientific soundness of the review for technical specifications?</li> <li>Will the reviews for scientific acceptability be completed via email or will there be calls open to the membership and the public? What opportunity will developers have to provide additional information or clarify questions about their submissions? The FAH is concerned and sees the potential for this process to be a "black box" and not meet the openness/transparency and due process components of OMB circular A-119.</li> <li>Will Standing Committee members be able to "overrule" the methods panel</li> </ul>	Commenter	committees can still discuss relevant issues, such as risk adjustment. The rating from the methods review—whether generated by NQF staff or the Scientific Methods Panel—will be used to rate the <i>Scientific Acceptability</i> of the measure. However, standing committees may raise concerns with the specifications of the measure or with potential threats to validity (e.g., selection of variables for risk adjustment model) and can therefore overturn the staff or Scientific
<ul> <li>or NQF staff and change the rating on reliability and/or validity? The language implies that the Standing Committee will only be able to raise concerns with the specifications or potential threats to validity. Are these the only items that committees will be able to address? If they do raise concerns, what happens to the preliminary ratings? Will it require a vote by the Committee?</li> <li>The proposal states that: "Generally, NQF will not forward measures with a "low" or "insufficient" rating from the methods review to the committee for further evaluation." What criteria will be used to determine if a measure with those ratings did or did not move forward to a committee? As the FAH understands the current process, if a measure fails on one of the 4 key criteria, it cannot move forward: i.e., if reliability or validity is low, then that is the an automatic Stop-Now indicator.</li> </ul>		Methods Panel ratings. NQF will assess each measure based on the measure evaluation criteria outlined in guidance documents for both developers and committee members. Measures rated by NQF staff or the Scientific Methods Panel as " <i>Low</i> " or " <i>Insufficient</i> " for reliability or validity will be removed from the current evaluation cycle, allowing time for any additional testing, clarification or NQF technical support, or review prior to consideration of the measure in a future cycle. NQF always welcomes measure stewards/developers to request technical assistance prior to the submission deadline.
<ul> <li>Currently, staff preliminary analyses are inconsistent across the various committees and can be too prescriptive (e.g., testing is marked as insufficient because Kappa statistics are not provided; yet, percent agreement is near or at 100% and a Kappa statistic would not be meaningful). Or, assessment is too lax (e.g. SDS submissions with inadequate conceptual analyses are not rated as insufficient). A good indicator of the committee agreement (or lack thereof) with staff analyses is the number of times a committee's final decision aligns with the staff</li> </ul>		Public Commenting Period: The committee will review any comments received after the committee evaluation meeting during the post-commenting period call. All submitted comments during this time will receive written responses from the standing committee, measure steward/developers, and/or NQF, as appropriate. The standing committee may revise its recommendations in response to a specific comment or series of comments submitted. These changes will be communicated broadly prior to the CSAC's review.

Comment	Commenter	NQF Response
<ul> <li>recommendation NQF should evaluate the degree of concordance and determine what the reasons for any lack of agreement may be. This exercise may be useful regardless of which group (i.e., NQF staff, methods panel, Standing Committee) is the one best able to perform the necessary preliminary analysis.</li> <li>The proposal calls for reviews not to move forward for non-complex measures that achieve low or insufficient ratings. The FAH has seen these low and insufficient ratings provided on maintenance measures that did not provide new testing. What would happen in those instances? What are the situations in which previous testing on maintenance measures would not be accepted? Would these measures not be put forward to the committee for evaluation? Would NQF just remove endorsement without any evaluation and adjudication by a committee?</li> <li>Because of this inconsistency, it would be preferable to have the methods panel review ALL measures or still ask the Standing committees to evaluate the less complex measures, while the methods panel reviews the complex measures. Since the process is designed to achieve consensus, it is unclear how having NQF staff serve as the arbiter/decision maker in the proposed process achieves the goal of consensus. The FAH is concerned about measures that are rated as low or insufficient by NQF staff, which means the review would stop at that point. The multi-stakeholder committee would not see the measures. This process seems to be fraught with potential to not meet the balance and due process components of the OMB circular.</li> <li>Measure Evaluation Technical Report</li> <li>The proposed changes seem reasonable and responsive to feedback provided by the FAH and other members.</li> </ul>		Endorsement Decision: Given important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. NQF is committed to implementing a plan to identify and solicit ongoing engagement and participation opportunities from the consumer and purchaser stakeholder perspective. Depending on the outcome of this initiative, NQF could potentially implement this proposed change at a later time. Improvements in Information Exchange and Access: NQF wi adopt a two-fold approach to addressing recommendations from Kaizen participants. Some aspects of the recommendations are resolvable through short-term solutions and adaptations of existing platforms. Other recommendations will be addressed through a long-term product development approach. NQF will solicit stakeholde input through this process as appropriate.

Comment	Commenter	NQF Response
Public Commenting Period with NQF member Expression of Support		
It appears that at least one or two steps are missing in this process. The proposed		
change would eliminate the voting step, and it is not clear how the indications of		
support would be determined or how differences in opinion across the membership		
will be identified and/or adjudicated.		
What happens to comments that are submitted after the committee		
evaluation meeting? Will there be a follow-up conference call to review the		
comments and consider revising recommendations on measures based on		
member submissions? If the recommendations change, will this		
information be posted for members in an easily found site? Will members		
be given an opportunity to change their indications of support? Currently, it		
is not clear what the process is after the initial meeting to evaluate the		
measure and how differences of opinion will be adjudicated.		
<ul> <li>How are these indications of support then used? Will it be the same</li> </ul>		
process that is currently used for voting with results provided to the CSAC?		
Will other members and the public be able to see the comments and		
indications of support throughout the process? The FAH recommends		
ensuring that information is transparent to anyone who participates in the		
process.		
<ul> <li>Additional information is needed on how these changes would be</li> </ul>		
implemented and what the actual process steps would be before NQF		
moves forward with this proposed change. The membership needs more		
opportunity to discuss and provide input after the basic questions are		
answered. The FAH is concerned that there is potential to lose the		
consensus-based process in this step or at least weaken the consensus if		
this step is not handled carefully. The FAH is concerned that this step, in		
particular, may not meet the OMB circular guidance.		
Endorsement Decision		

Comment	Commenter	NQF Response
The FAH is concerned about moving the final endorsement decision to Standing		
Committees. This change assumes that all committees evaluate measures in the		
same manner and are consistent in their decision-making. Removing an oversight		
body such as the CSAC seems premature. As a long-time NQF member, the FAH		
wants to see the system carefully specified and tested before such a drastic step is		
taken.		
Adjudication of Appeals		
The staff recommendation not to change the role of CSAC to be the arbiter of		
appeals appears reasonable at this point. The appeals board is new and whether this		
revised process works effectively or not must still be determined.		
Enhancing training and Education		
The proposal calls for additional steps to ensure that developers, committee		
members and staff are adequately trained. This is a positive step and should be		
undertaken no matter what happens to the overall proposal.		
Improvements in Information Exchange and Access		
The NQF is limited in the changes it can make at this time, but is making this public		
commitment to working on short-term solutions. The FAH strongly encourages NQF		
to solicit input from stakeholders involved in the CDP and MAP process. The two-		
week comment period on this broad and sweeping proposal to change the entire		
basis on which measures are endorsed is too limited for the import of the changes		
being put forth in this document. Many details have yet to be answered. It would		
be preferable for NQF to take the time to solicit input from the various stakeholders		
involved in the CDP and MAP processes to ensure that their issues of greatest		
significance are captured and addressed. Any finalized CDP process changes must		
ensure that the membership is fully on board. This short two-week comment		
period, which happens to fall right in the middle of the heaviest federal public		

Comment	Commenter	NQF Response
commenting period on payment rules for the next fiscal year, is concerning. It is highly unlikely that NQF members will feel they have been truly engaged with these proposed major changes to the CDP process.		
The American Medical Association (AMA) appreciates the opportunity to comment	Koryn Rubin,	Thank you for your comment. We appreciate your feedback
on the proposed changes to the Consensus Development Process (CDP). We support	American	on the proposed recommendations for the CDP Redesign.
the ongoing efforts to continuously improve the CDP process and are encouraged by	Medical	NQF hosted the Kaizen event in collaboration with CMS to
the possibility of submitting measures more frequently for review but are highly	Association	inform the CDP redesign. CMS, as the funder of this
concerned with the lack of detailed information provided in the draft document.		initiative, has asked NQF to solicit public comment on the
The brevity paired with the extremely quick turnaround time for review and		proposed recommendations and provide a final report
comment does not allow adequate time or sufficient information to provide		outlining the new CDP by July 1, 2017. Thus, NQF had to limit
substantive comments. Rather, we have outlined our general thoughts and		the amount of time NQF members and the public had to
questions for each of the proposed changes. Any changes to the CDP and		provide feedback. However, as NQF continues to plan for
application of the measure evaluation criteria must be consistent within and across		implementation of the new CDP, additional feedback is
projects.		welcomed. Additional details have been included in the final
		report for clarity.
The following are questions and comments on the proposed changes:		
General Questions/Comments		General Questions/Comments: To allow more frequent
		measure submissions, committees will convene more often.
<ul> <li>It would be useful to understand what the increase in the number of topics twice a year would mean for NQF membership and committees. How many</li> </ul>		Additionally, there will be increased opportunities for NQF
additional measures and reports might be released in a year for member		membership to engage in the process. As a result, NQF
		emphasizes the importance of stakeholder education.
and public comment compared to now? Currently, there are too many		To assist in planning and minimize burden for the measure
competing projects including measures under review and frameworks,		stewards/developer, committee members and NQF,
which result in an extremely low member comment response rate, and the		submission deadlines will be staggered. A graphic that
inability of members to adequately evaluate, review and comment on NQF activities.		outlines the new process is included in the final report.
To what degree will the measure submissions and comment periods be		After the standing committee completes its measure review,
staggered?		a summary of the committee's recommendations – or "draft
A graphic or visual that shows what the revised CDP would look like once all		report" – will be posted on the NQF website for the public

Comment	Commenter	NQF Response
of these changes are implemented would be helpful. There is nothing in		and NQF membership to review and comment. Because
this document that pulls everything together so that you can see the entire		there are more review cycles, NQF will revise the content
process from start to finish.		and structure of the report to highlight key elements of
		interest. These elements are included in the final report.
Scheduling/Frequency		
The AMA supports the opportunity for more frequent submissions and reviews		Scheduling/Frequency: The process includes a two-week
during the NQF CDP process. However, the proposed approach still uses a "batching"		process for measure steward or developer to respond to the
methodology that will force decisions to be made on measures within a certain		ratings. In addition, the standing committees can still discuss
timeframe and project. This process is fine for measures that are well developed		relevant issues, such as risk adjustment. Given the
and generate minimal controversy. What the process still does not allow for is an		opportunity for more frequent submission, measures may
iterative process where disagreements and concerns can be addressed before a final		need to be move to the next review cycle to address
decision is made. NQF should develop a process through which measures can enter		methodologic concerns.
at any time, be sent to a committee for review when they are deemed ready and		
then go through multiple comment cycles or committee reviews until there is		The consolidated topic areas are included in the final report.
general agreement that the measure is ready to be endorsed. The twice-yearly		Topic areas were consolidated with the goal of reassessing
submission process meets some of those needs, but it does not address the true		and balancing NQF's library of measures, while distributing
needs – the ability to submit a measure at any time and the ability to achieve true		measures to committees with the needed expertise to
consensus. For instance, the process should incorporate some sort of tabling		conduct an evaluation. As a result, many of the smaller
mechanism—where a controversial measure can be sidelined to allow issues to be		portfolios have been consolidated into cross-cutting topics
worked through, and brought back for review when ready. The AMA has experience		with a broader range of experience. In addition, some clinical
in convening approval and evaluation processes and would be happy to explain how		groupings of committees were made to reflect more cross-
the CPT and RUC processes handle such issues.		cutting clinical areas, such as primary care and chronic illness
		care, pediatrics and geriatrics and palliative care. NQF will
While the document says that 22 topic areas are merged into 16 topics it lacks		offer two measure submission opportunities for each topic
sufficient detail. The AMA has the following outstanding questions and issues that		area each year, limiting the number of measures evaluated
must be clarified before a new CDP can be finalized:		by the standing committees in each cycle to a maximum of
• 16 Topic Areas: What are the 16 topic areas and what topics will sunset		12 (up to eight measures undergoing maintenance review
and/or merge? How did NQF define and arrive at the 16 topic areas? How		and up to four new measures). This was determined given
does this impact the number and composition of the Standing Committees,		that approximately 80% of the measures submitted for

	Comment	Commenter	NQF Response
parti	icularly for topics that are retired and combined?		endorsement consideration are maintenance measures.
• Capp	bing CDP at 12 Measures Per Cycle: We are concerned with capping the		However, the combination of maintenance and new
num	ber of measures at twelve per cycle and the potential ramifications of		measures may vary depending on number of measures
this a	arbitrary cap do not appear to have been considered. It is unclear how		submitted, opportunities for related and competing measure
NQF	arrived at the number and why only a maximum of eight measures		review, and measure prioritization efforts. Per NQF's
unde	ergoing maintenance and up to four new measures can be considered		maintenance of endorsement policy, measures are due for
per o	cycle. It is possible that NQF will receive more than twelve measures		reassessment every three years. NQF will remind measure
for o	one cycle. For instance, MIPS requires a physician to report on six		stewards and developers of scheduled measure
meas	sures, one of which must be an outcome or high priority measure.		maintenance review several months prior to the review and
Ther	efore, we envision measure developers would put forward a suite of		notify each of their assigned review cycle.
meas	sures in a clinical topic area, but the revised process may not allow for		
revie	ew of multiple measures in one clinical area during a single cycle.		Technical review: Methods Panel: NQF has provided
Ther	efore, we are concerned that the arbitrary cap on new measures (up to		additional information on the composition and disclosure of
four	per cycle) may impact MIPS compliance and the transition to more		interest of the Scientific Methods Panel in the final report.
inno	vative and meaningful measures by limiting the number of new		The new NQF Scientific Methods Panel will consist of 15 to
meas	sures that can undergo review. We, also request further clarification		25 statisticians, epidemiologists, psychometricians,
on th	he proposal to cap measure reviews at twelve per cycle:		economists, performance measure methodologists and
	<ul> <li>How will NQF prioritize measures? What if there are more than</li> </ul>		individuals with expertise related to eMeasures and
	four new measures in a cycle? Would priority be given to the first		disparities. All nominees will complete an annual general
	measures submitted or be based on some other criterion? We		disclosure of interest (DOI) form, as well as measure-specific
	would note that prioritization must be balanced and not partial to		disclosures to identify recusals from specific measures. NQF
	any one group (e.g., CMS).		will assign measure review based on identified conflicts of
	<ul> <li>Will there be any flexibility on the maximum number of</li> </ul>		interest, relevant expertise, and availability of panel
	maintenance and new measures that can be included in any one		members. All reviews provided by the Scientific Methods
	cycle? If so, what are the parameters by which the numbers in		Panel will be shared not only with the committee but also
	each category may change? How will maintenance measures be		with the steward/developer and the public. Furthermore,
	selected for a cycle, especially given the "intent to submit"		the Scientific Methods Panel's charge will include providing
	requirement?		expertise for methods/testing-related issues for NQF and
	<ul> <li>How does capping the sixteen topic areas impact the limit on</li> </ul>		advance NQF's guidance on these issues.

Comment	Commenter	NQF Response
twelve measures per cycle since the number of measures in some		
of the topic areas may increase with this shift?		NQF will provide standard guidance on assessing the
<ul> <li>It is possible that maintenance measures may not be reviewed</li> </ul>		Scientific Acceptability criterion for a measure, using the
within the 3-year cycle given the limited number of measures		current decision algorithm used from the measure
within each cycle?		evaluation criteria. As part of its ongoing education efforts,
		NQF will provide clear guidance to standing committees
Intent to Submit		regarding the circumstances wherein an overturn of the
How will this step be used for maintenance measures? Will NQF notify		rating would be permissible.
developers when maintenance is due? As proposed, it appears that the		
onus is on the developer regardless of whether it is a new measure or		For both complex and non-complex measures, NQF will sen
maintenance measure.		the preliminary analysis to developers by email for review.
• This section also does not say what NQF will do if they get more than 12		The process includes a two-week process for measure
measures to be submitted in a cycle review? How will NQF decide which		steward or developer to respond to the ratings. In addition,
measures get moved to the next cycle? How will NQF prioritize which		the standing committees can still discuss relevant issues,
measures will be reviewed in a given cycle? How quickly will NQF let		such as risk adjustment. The rating from the methods
developers and the public know if that occurs (e.g., a month before the		review—whether generated by NQF staff or the Scientific
submission deadline, after the submission deadline)?		Methods Panel—will be used to rate the Scientific
		Acceptability of the measure. However, standing
Technical Review: Methods Panel		committees may raise concerns with the specifications of the
The creation of this panel has the potential to strengthen the evaluations for		measure or with potential threats to validity (e.g., selection
scientific acceptability and may improve consistency across measures and projects.		of variables for risk adjustment model) and can therefore
However, there are several questions around the panel itself, the roles and		overturn the staff or Scientific Methods Panel ratings.
responsibilities of this panel along with NQF staff and Standing Committees, and the		
actual process and guidelines around these reviews. The AMA requests more detail		NQF will assess each measure based on the measure
on who would be on the Technical Review panel. Specifically, we seek more		evaluation criteria outlined in guidance documents for both
information on how the Technical Review panel will operate. Will the Technical		developers and committee members. Measures rated by
Review panel operate similar to a Standing Committee with terms, level of		NQF staff or the Scientific Methods Panel as "Low" or
education/training? The level of education, training and standardization will have a		<i>"Insufficient"</i> for reliability or validity will be removed from
big impact on the success of the Technical Review panel. Otherwise, the same		the current evaluation cycle, allowing time for any additional
Page		testing, clarification or NQF technical support, or review

Comment	Commenter	NQF Response
challenges which occurred with the Standing Committee, such as members reporting		prior to consideration of the measure in a future cycle. NQF
difficulties completing scientific and statistical reviews, may still occur. We are also		always welcomes measure stewards/developers to request
concerned that this may create a diminished role for clinical perspective and		technical assistance prior to the submission deadline.
expertise.		
		Public Commenting Period: The committee will review any
It is not clear how the creation of this panel would allow for additional participation		comments received after the committee evaluation meeting
by consumers, patients, and purchasers. This statement implies that determinations		during the post-commenting period call. All submitted
of reliability and validity of measures are moved from the committee to NQF staff.		comments during this time will receive written responses
This seems contrary to a process that is designed to achieve consensus across		from the standing committee, measure steward/developers
stakeholder groups.		and/or NQF, as appropriate. The standing committee may
		revise its recommendations in response to a specific
We also seek further clarification on the operations of the Technical Review panel:		comment or series of comments submitted. Any decisions
• Will the scientific acceptability reviews be completed via email or will there		will be communicated broadly prior to the CSAC's review.
be calls open to the membership and public? What opportunity will developers have to provide additional information or clarify questions?		Endorsement Decision: Given important strategic
There is the potential for this step in the process to be a "black box" and		considerations, NQF will not be able to implement a change
decrease transparency with the CDP.		of this magnitude at this time. NQF is committed to
• Will Standing Committee members be able to "overrule" the methods panel		implementing a plan to identify and solicit ongoing
or NQF staff and change the rating on reliability and/or validity? As drafted,		engagement and participation opportunities from the
the proposal implies that Standing Committees will only be able to raise		consumer and purchaser stakeholder perspective.
concerns with the specifications or potential threats to validity. Are these		Depending on the outcome of this initiative, NQF could
the only issues that committees will be able to address? If they do raise		potentially implement this proposed change at a later time.
concerns, what happens to the preliminary ratings? Will it require a vote by		
the Committee?		Improvements in Information Exchange and Access: NQF wil
• The document states that, "Generally, NQF will not forward measures with		adopt a two-fold approach to addressing recommendations
a 'low' or 'insufficient' rating from the methods review to the committee for		from Kaizen participants. Some aspects of the
further evaluation". What criteria would be used to determine if a measure		recommendations are resolvable through short-term
with those ratings did or did not move forward to a committee?		solutions and adaptations of existing platforms. Other
<ul> <li>Currently, staff preliminary analyses are incredibly inconsistent across the</li> </ul>		recommendations will be addressed through a long-term

	Comment	Commenter	NQF Response
	various committees and can be too prescriptive (e.g., testing is marked as		product development approach. NQF will solicit stakeholder
	insufficient because Kappa statistics are not provided; yet, percent		input throughout this process as appropriate.
	agreement is near or at 100% and a Kappa statistic would not be		
	meaningful) or too lax (e.g., SDS submissions with inadequate conceptual		
	analyses are not rated as insufficient). A good indicator of the committee		
	agreement (or lack thereof) with staff analyses is the number of times a		
	committee's final decision aligns with the staff recommendation.		
	Therefore, NQF should look at the degree of concordance and determine		
	the reasons for lack of agreement. Our recommended exercise may be		
	useful regardless of which group (i.e., NQF staff, methods panel, Standing		
	Committee) is best able to perform this preliminary analysis.		
٠	Furthermore, it is proposed that if the reviews for non-complex measures		
	are low or insufficient, the measures would not move forward. Based on		
	our experience with the CDP, we have seen these low and insufficient		
	ratings provided on maintenance measures that did not provide new		
	testing. What would happen in those instances? What are the instances in		
	which previous testing on maintenance measures would not be accepted?		
	Would these measures not be put forward to the committee for		
	evaluation? Would NQF remove endorsement without any evaluation and		
	adjudication by a committee?		
•	Due to the potential continued inconsistency, it may be preferable to have		
	the methods panel review ALL measures or to continue to have the		
	Standing Committees evaluate the less complex measures, while the		
	methods panel reviews the complex measures. Because the process is		
	designed to achieve consensus, it is unclear how having NQF staff serve as		
	an arbiter/decision maker in the process achieves that goal; particularly, if		
	measures that are rated as low or insufficient by NQF staff would not go		
	forward to a multi-stakeholder committee.		

Comment	Commenter	NQF Response
Measure Evaluation Technical Report		
<ul> <li>These changes seem reasonable and responsive to feedback provided by</li> </ul>		
the AMA and other members over the years.		
Public Commenting Period with NQF Member Expression of Support		
As proposed, it appears there are steps missing in the revised process because the		
proposed changes would eliminate voting. It is also unclear how the indications of		
support would be determined or how differences in opinion across membership will		
be adjudicated. We request clarification on the following questions related to public		
commenting and voting:		
What happens to comments that are submitted after the evaluation		
meeting? Will there be a follow-up conference call to review and consider		
revising recommendations on measures based on what is submitted? If the		
recommendations change, will this information be posted and members		
given an opportunity to change their indication of support? As proposed, it		
is not clear what the process is to evaluate a measure after the initial		
meeting and how differences of opinion will be adjudicated.		
<ul> <li>How are the indications of support then used? Will it be the same process</li> </ul>		
as is currently used for voting now with results provided to the CSAC?		
<ul> <li>Will other members and the public be able to see the comments and</li> </ul>		
indications of support throughout the process? It will be important to		
ensure that this information is transparent to anyone who participates in		
the process.		
Additional information is also needed on how these changes would be implemented		
and what steps would be taken before NQF moves forward with this proposed		
change. There is significant potential that these changes could reduce the		
consensus-based nature of the process.		

Comment	Commenter	NQF Response
Endorsement Decision		
We are concerned with moving the final endorsement decision(s) to Standing		
Committees. The change assumes that all committees evaluate measures in the		
same manner and are consistent in their decision-making. Removing an oversight		
body such as the CSAC seems premature.		
Adjudication of Appeals		
The staff recommendation not to change the role of CSAC to be the arbiter of		
appeals right now is very reasonable. The appeals board is new and whether this		
revised process works effectively or not must still be determined.		
Enhancing Training and Education		
The additional steps to ensure that developers, committee members and staff are		
adequately trained are very positive. Additional detail on exactly what will be		
provided (e.g., schedule of events) would be helpful.		
Improvements in Information Exchange and Access		
Since NQF is limited in the changes that can be made at this time but commits to		
working on short-term solutions, it would be preferable if they solicit input from		
various stakeholders involved in the CDP and MAP as they move forward in this		
effort to ensure that what is most important to membership is prioritized.		
For questions or to discuss the AMA's comments further, please contact Koryn		
Rubin, Assistant Director, Federal Affairs at 202-789-7408 or koryn.rubin@ama-		
assn.org.		
The Council of Medical Specialty Societies (CMSS) is pleased to comment on the	Norman Kahn,	Thank you for your comment. NQF appreciate your feedback
proposed changes to the endorsement process of the National Quality Forum (NQF).	Council of	on the proposed recommendations for the CDP Redesign.
CMSS is a long-time member of NQF and serves on the National Quality Partners	Medical	
Leadership Council.	Specialty	Endorsement Decision: NQF appreciates your suggestion on

Comment	Commenter	NQF Response
On at least two occasions in the past decade, CMSS has requested that NQF	Societies	the composition of the standing committees. Given current
streamline its endorsement process, and make it more "user-friendly." We are		resources and other important strategic considerations, NQF
pleased that NQF has been responsive to such feedback in the past. At present, it		will not be able to implement a change of this magnitude at
appears that NQF's Kaizen process promises to modernize the critical measure		this time. If or when this change occurs, we will consider
endorsement function that NQF plays in the US health system.		your feedback on the approach.
CMSS supports NQF's proposed changes, with the following comments:		Improvement in Information Exchange and Access:
<ul> <li>We agree with NQF's proposed changes, but caution against replacing</li> </ul>		NQF is working to identify solutions to enhance our current
professional clinicians with consumers and payers on the workgroup		IT infrastructure to provide a more user-friendly experience
committees (if CSAC is no longer responsible for final endorsement). While		when submitting a measure for endorsement consideration.
we support adding seats for the public on workgroups, we would oppose		NQF will advance a short-term initiative to aggregate
heavily weighting workgroups with consumers and payers, as we feel this		information by grouping MAP measure recommendations
may put at risk the scientific nature of workgroups. It is critical that enough		and rationale issued each year into one comprehensive and
seats be available for professional members to ensure cross-specialty		filterable document, accessible from the existing MAP
evaluation, endorsement, and acceptance by the medical community.		homepage on the NQF website. Similarly, NQF will work to
We encourage NQF to make improvements in information exchange and		consolidate existing information from CDP reports to make it
access. Submission to NQF is an arduous process taking no less than 40		easier for users to access measure information. In addition,
hours for one submission, in large part due to lack of smart forms and the		NQF will advance a short-term initiative to improve business
required resubmission of duplicative information.		rules around publishing timelines and meeting materials, to
When several disciplines are lumped into one topic, a limitation of 4 new		ensure developers, committee members, and members of
measures may be too restrictive. As an example, ENT, ophthalmology, and		the public are more aware of opportunities to participate in
optometry are all in the same topic. This would restrict three disciplines to 4		NQF's processes. Commenting opportunities will also be
new measures.		enhanced by increasing the character limit to 10,000
• Twice yearly cycles are good, but may very well result in a burden on the		characters, real-time updates on comments forwarded to
committee members. Each committee meeting tends to be two days plus		developers, and better regulated public comment periods
conference calls, which is a significant work burden for volunteers.		during evaluation meetings.
In addition, CMSS has a few questions as NQF proposes changes:		Scheduling and Frequency: NQF has conducted an analysis of
How will NQF prioritize measures in a situation where more than eight new		the number and types of measures (maintenance and new)
		submitted for endorsement consideration for each topic

Comment	Commenter	NQF Response
<ul> <li>measures are submitted in a year, and how will NQF ensure endorsement review occurs in a timely manner for these extra measures received?</li> <li>How will the 22 current topical areas be reduced to 16, and will the public have input on these future groupings?</li> <li>It is anticipated that there may be situations where measure developers</li> </ul>		area over the last several years. In addition, NQF used a decision logic to inform which topical areas can be combined to create a comprehensive topical area portfolio. NQF will monitor the submissions closely to ensure developers have the opportunity to submit measures in each discipline within
may disagree with the determination by NQF staff or external methods panel of low or insufficient ratings. What recourse is available when a		the combined topical areas.
developer disagrees? Will there be an appeal process through the external methods panel or the standing committee?		NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to
CMSS appreciates efforts on the part of NQF to modernize and streamline the process of measure endorsement. We hope that our feedback will serve to		four new measures). NQF may add one or two additional measures as deemed appropriate. However, the
continually improve the process. Sincerely,		combination of maintenance and new measures may vary depending on number of measures submitted, opportunities
Norman Kahn MD Executive Vice-president and CEO		for related and competing measure review, and measure prioritization efforts. Any Intent to Submit forms that are
		submitted once capacity has been reached for a particular cycle, NQF will notify the steward/developer and provide the date of the next cycle in which there is availability.
		NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included
		in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures,
		while distributing measures to committees with the needed expertise to conduct an evaluation. As a result, many of the
		smaller portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more

Comment	Commenter	NQF Response
		cross-cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care. NQF will not formally open a public comment period on the consolidated list of topical areas. However, NQF welcomes your feedback on the list. If you have input, feel free to email NQF at <u>NQFKaizen@qualityforum.org</u> .
		Technical Review: Methods Panel: The process includes a two-week process for measure steward or developer to respond to the ratings. In addition, the standing committees can still discuss relevant issues, such as risk adjustment. Given the opportunity for more frequent submission, measures may need to be move to the next review cycle to address methodologic concerns.
NCQA was pleased to participate in the recent Kaizen representing both the measure	Bob Rehm,	Thank you for your feedback on the Improvements in
development community, but also as a measure implementer. Kudos for NQF to	National	Information Exchange and Access comments. We concur
address CDP availability, management and expedited endorsement timeframe.	Committee for	that there are achievable short-term goals that will enhance
	Quality	usability. NQF will group measure recommendations &
Scheduling/Frequency: We want to thank NQF for demonstrating flexibility in	Assurance	rationale issued each year into one comprehensive and
shifting maintenance measures to later (or earlier) CDP projects. Often measure		filterable document, accessible from the existing MAP
developers are in the midst of a re-evaluation cycle when NQF schedules a CDP		homepage. We expect this will address the challenge in
review. Having the flexibility to complete that internal work prior to engaging with		tracking historical recommendations you highlighted.
NQF results in better measures and a more efficient process. A win-win. We agree		
with the reduction in topical areas and would recommend that NQF publish those		
for public comment.		
During the recent Kaizen there was feedback that standing committees are		
structured to have a variety of experts, and so sometimes a measure will apply to		
only some of the committee members. This means that committee members may		

Comment	Commenter	NQF Response
be reviewing measures for which they have limited experience/expertise. This might		
become an even bigger issue with the consolidation of topic areas. NQF might		
consider how to handle this so that committee members are discussing/voting on		
measures that are within their area of interest/expertise. One option would be the		
introduction of an "abstain" option in voting for panel members who do not feel		
they are in a position to vote on a specific criteria or suitability for endorsement.		
NQF could clarify this option during panel orientation and at the CDP endorsement		
meeting.		
CDP Meetings: We agree with the proposal to alternate in-person and virtual		
meetings. However, we would like to ensure that quorum for any meeting is		
reached and that voting occur at the same time as the measures are presented and		
discussed. Voting off-line by committee members not present during discussion of		
the measure is inappropriate.		
While not referenced in your proposal, we also recommend that public comment		
during standing committee meetings occur prior to voting on a given measure.		
Intent to Submit: We strongly recommend that NQF (not the measure		
developer/steward) initiate this request for Maintenance Measures. NQF currently		
prepopulates the submission form and asks developers/stewards to verify that there		
are no significant changes to the measure. If none, then NQF incorporates the		
measure into the relevant CDP process. The current process for maintenance		
measures does not need to be changed. Measure developers/stewards should not		
have to "tee-up" measures already endorsed.		
For New Measures, we support the new process and timing for intent to submit.		
Technical Review: Methods Panel: We would like to emphasize the importance of		

Comment	Commenter	NQF Response
NQF sharing the "initial staff review" with the measure developer/steward. This will		
ensure that any corrections, misinterpretations or inadvertent errors are addressed		
<i>prior</i> to distribution to the standing committee.		
NQF should identify the criteria used to determine if a measure will undergo an NQF		
staff review of methods, or if that function will be performed by a technical advisory		
panel or ad hoc group composed of other convened standing committee members.		
Developers/stewards should be provided an opportunity to provide feedback on that criteria.		
NQF should clarify that findings of the methods review will be presented to the		
standing committee as "recommendations" and that guidance be provided to a		
panel for over-riding these recommendations. Will developers have an opportunity		
to appeal the finding of the methods review? If so, NQF should provide guidance on		
that process. Will the standing committee continue to discuss and vote on scientific		
acceptability? Or will the vote be on the recommendation from the methods review?		
Will the same rules of voting still apply (< 40% - criteria not met; > 40 and < 60 -		
consensus not reached; > 60 - criteria met)? Is there time built in for the developer		
or steward to address issues that emerge from the methods review? We would like		
to recommend that NQF consider an option, short of a vote on endorsement, for the		
measure to be placed 'on hold' and allowing developers to come back to a future		
methods panel review for potential resolution. This alternate pathway may be more		
efficient and expedient for all concerned.		
Due to the scale of changes put forth by NQF, we would recommend that a pilot		
project be initiated for a subset of CDP projects. Results of that pilot may be helpful		
in addressing issues and identifying strategies that work for NQF, the standing		
committees and the measure developers/steward, alike.		

Comment	Commenter	NQF Response
Measure Evaluation Technical Report – Content and Structure: We support		
additional streamlining. We also stress the importance and the value of providing measure developers/stewards an opportunity to review the technical report prior to posting?		
Extended Public Comment Period: Can NQF clarify for maintenance measures, is this		
pre-committee evaluation public comment for the newly submitted maintenance		
measure or the maintenance measure from the last time it was reviewed/went		
through annual update? It seems it would be most helpful if the public was		
commenting on the newly submitted maintenance measure. This would avoid		
confusion, especially in cases where measures have been updated.		
NQF should clarify the expectations for responses to public and member comment		
by the developer/steward during this "rolling" public comment period. This has the		
potential to add to, not reduce confusion, for all parties.		
NQF should time stamp public and member comments; so they can be considered		
within the context of key events in the CDP process (e.g., comment received before		
materials published, comment received following standing committee meeting and		
vote). Commenters should attest that they have read and reviewed all available materials.		
Endorsement Decision: We support CSAC's oversight role and agree that the		
current process of CSAC ratification serves the field well.		
Adjudication of Appeals: We support the continued role of the Appeals Board and		
would recommend that NQF allow <i>all</i> measures the right of a formal appeal.		
Currently only measures recommended as suitable for endorsement by the standing		
committee and ratified by the CSAC can be appealed. We would recommend that		

Comment	Commenter	NQF Response
the Appeals Board have a broader charge including review of measures not		
recommended for endorsement and not ratified by CSAC. We recommend that		
appeals continue to be considered when the measure developer/steward or other		
parties believe that NQF criteria was not applied correctly or the process was not		
followed. An option for NQF to consider: the Appeals Board could overturn decisions		
and send the measure back to consideration in the next cycle of the standing		
committee without the loss of endorsement. This would actually streamline the		
process and eliminate unnecessary repeat steps in the CDP process.		
Enhancing Training and Education: We agree that further training and resources are		
needed for those engaged in the CDP process, including NQF staff, standing		
committee members, supporting advisory panels, developers/stewards, NQF		
members and the public. We would also recommend that new members to standing		
committees receive more concentrated support including holding of "dry runs" that		
can simulate the review process and expose them to a range of scenarios that test		
their understanding of the CDP process and interpreting the NQF evaluation criteria.		
Often times, this occurs during the first reivew of a measure by the panel, which is		
less than optimal. We would be delighted to support NQF in this effort.		
Improvements in Information Exchange and Access: We support this innovative		
approach to integrating measure information across projects. The field will benefit		
and in the process become more knowledgeable about measure development. In the		
meantime, as NQF works towards this ideal state, are there any immediate small		
successes that can be achieved? For example, creating an index that shows which		
MAP reports contain recommendations for each measure (so that users of this		
information do not have to search each report separately)?		
Additional Comments: Intention to Submit-Review Process: The proposal includes		
the following language, NQF staff will assess whether the measures will require a		

Comment	Commenter	NQF Response
<ul> <li>methodologic review based on a set of criteria. While we support this approach, NQF</li> <li>should make this set of criteria available for feedback before finalization. NQF should</li> <li>ensure that reviewers have sufficient training, experience and oversight to perform</li> <li>this important assessment and share findings with the measure developer/steward,</li> <li>allowing 3 business days to respond.</li> <li>Econometrica, Inc. appreciates the opportunity to comment on the 2017 Kaizen on</li> </ul>	Mark Stewart,	Thank you for your comment.
redesigning the Consensus Development Process. While we look forward to a more streamlined process with more frequent opportunities to submit measures to the Standing Committees, we have several questions and concerns regarding the proposed changes. First, will there be an opportunity to review or provide recommendations on the change from 22 to 16 Standing Committees? It is difficult to assess the impact of this change on the CDP without knowing what topic areas each Committee will cover.	Econometrica, Inc.	Scheduling/Frequency: NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures, while distributing measures to committees with the needed expertise to conduct an evaluation. As a result, many of the smaller
It seems likely that a Standing Committee will receive Intent to Submit for more than four measures in any given review cycle, particularly in the period immediately following the implementation of the new process, because many developers and stewards have been waiting for existing projects to open a new review cycle. How will the Standing Committees select which measures they will review when they receive notification regarding more than four new measures? Will the selection be based solely upon the timing of submission of the Intent notice, or will more than 4 measures be allowed to proceed to methodological review first? In the case that one or more of the measures do not end up being submitted to the Standing Committee, will there be a waiting list or some other process to allow the queue to remain		portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more cross- cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care. NQF will not formally open a public comment period on the consolidated list of topical areas. However, NQF welcomes your feedback on the list. If you have input, feel free to email NQF at <u>NQFKaizen@qualityforum.org</u> .
filled? As a developer working on multiple measures simultaneously, we also wonder whether any attempt will be made to keep multiple measures submitted to a single Standing Committee by the same developer or steward together? If multiple		NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). The combination of maintenance and new measures may vary depending on number of measures

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measures for a single program are split up based on limiting each review cycle to 4 new measures, this will limit the potential benefit to shortening the process by requiring developer/steward presence at an additional round of Committee meetings, and by potentially delaying implementation of measures while the steward waits for the remaining measures to enter the endorsement process. In terms of a methodological review panel, Econometrica does not have specific objections to this change. We would like to encourage NQF to ensure that the training and guidance provided to this review panel is also provided to developers so that the panel's expectations and processes can be clearly understood. We applaud NQF for addressing the critical issue of improving the consensus development process. Overall, the changes seem to be a positive step in the right direction. We particularly support the addition of a dedicated methods review team that utilizes individuals with methodological expertise. This method will help standardize the endorsement process while reducing the burden of clinical reviewers. Furthermore, we support embedding key stakeholders such as consumers and purchasers from the CSAC within the standing committee. Doing so will not only improve the timeliness of committee approvals, but also involves these stakeholders in the critical work of assessing performance measures in a more direct and accountable manner.	Amir Qaseem, American College of Physicians	submitted, opportunities for related and competing measure review, and measure prioritization. NQF will notify the steward/developer and provide the date of the next cycle in which there is availability. Technical Review: Methods Panel: Enhancing training and education for the standing committee and scientific methods panel members, developers, NQF members and the public is a top priority for NQF. In the final report, NQF has provided additional details regarding the training and education plan for each audience. Thank you for your comment. CSAC Role in Endorsement Decisions and Appeals: We appreciate your feedback on the role of the standing committee as the final endorsement body, however given current resources and other important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. If or when this change occurs, we will consider your feedback on the approach. Enhancing Training and Education: Thank you for the
While we agree with the proposed changes, we encourage NQF to move towards pushing endorsement decisions to the standing committee. We also encourage NQF to make improvements in information exchange and access to include transparency in the process.		suggestion. As we develop our training and education plan, we will consider your recommendation.
On behalf of more than 52,000 members of the American Society of Anesthesiologists® (ASA), I welcome the opportunity to offer comments on the 2017 Kaizen Consensus Development Process: Proposed Redesign issued by the National	Jeffrey Plagenhoef, American Society	Thank you for your comment. We appreciate your feedback on the proposed recommendations for the CDP Redesign. Increased Opportunities for Measure Submission: NQF has

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Quality Forum (NQF). ASA looks forward to these and future NQF improvements to	of	consolidated the 22 measure review topical areas into 15
the measure endorsement process.	Anesthesiologists	topical areas. (The list of the topical areas is included in the final report). NQF will limit the number of measures
ASA supports continuous improvement to the NQF Consensus Development Process. ASA supports NQF's commitment to simplifying the measure development and endorsement process and ensuring stakeholders, including specialty societies, have ample opportunity to engage in the Consensus Development Process. We support several components of the proposed redesign, especially those aimed at simplifying the current process and providing a more transparent, streamlined process for submitting measures for endorsement to NQF and inclusion in federal payment programs.		evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted; opportunities for related and competing measure review, and measure prioritization. NQF will make every effort to standardize how both in- person meetings and web meetings are conducted to ensure
ASA supports increased opportunities for measure submission to NQF.		consistency in the Committee's measure review and evaluation process.
ASA supports instituting two measure submission periods per year for each topic area, as it will allow for stakeholders to have standard expectations of when measures may be submitted for endorsement. Increasing measure submission opportunities throughout the year will improve continuity of measure development for stakeholders and reduce dormancy previously experienced by standing committees.		Consensus Standards Approval Committee: NQF appreciates the comments received on the recommendation of this endorsement body. However, given important strategic considerations, NQF will not be able to implement a change of this magnitude at this time. Currently, the CSAC is comprised of a simple majority of consumers and purchasers. In order to ensure those two stakeholder perspectives are a key part of the endorsement process, NQF
opportunities.		will need to make certain there is adequate representation of these groups on each standing committee. NQF is
The proposed twelve measure limit for each topic area during each measure		committed to implementing a plan to identify and solicit
submission period is limiting and will slow the measure development process for		ongoing engagement and participation opportunities from
dense topic areas such as in surgical care. This cap would stall the measure		these stakeholder groups. Depending on the outcome of this
development process for new measures, as NQF has proposed that only eight (8)		initiative, NQF could potentially implement this proposed

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new measures will be considered in each topic area each year. NQF should consider		change at a later time
expanding the number of measures to review during each submission period or, at a		
minimum, allow for flexibility with this cap. Additionally, ASA recommends the close		Technical Review: Methods Panel: The new NQF Scientific
monitoring of this cap, to ensure it is appropriately meeting stakeholder demand.		Methods Panel will consist of 15 to 25 statisticians,
		epidemiologists, psychometricians, economists,
ASA supports increased frequency of standing committee meetings throughout the		performance measure methodologists, and individuals with
Consensus Development Process.		expertise related to eMeasures and disparities. NQF will
		solicit and identify nominees through NQF's standard
With increased measure opportunities throughout the year, standing committees		nominations process.
should meet more frequently to discuss submissions. While ASA supports more		
frequent convening of standing committees, NQF should closely monitor both		
review cycles to ensure measures reviewed in in-person meetings versus virtual web		
meeting are held to the same rigorous standards. When choosing standing		
committee members, NQF must strike an appropriate balance between clinicians		
who will be assessed on a majority of endorsed measures, with other stakeholders.		
NQF should remove Consensus Standards Approval Committee (CSAC) authority to		
overturn standing committee decisions.		
Removing the CSAC's authority to overturn decisions from standing committees will		
eliminate an unnecessary layer to the Consensus Development Process that is rarely		
used. Standing committees possess the subject matter expertise related to the		
measures in which they review, and the CSAC often defers to each standing		
committee for recommendations. ASA recommends NQF focus more energy on		
ensuring standing committees have diverse stakeholder representation and receive		
final endorsement authority.		
ASA supports the use of methodologists and staff experts to conduct "methods		
review" and make recommendations on "complex" measures.		

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ASA supports shifting the responsibility of "methods review" from each standing		
committee to NQF staff experts or an external methods panel. This will ensure a		
standardized methodological review process for each measure under consideration		
and allow experts to make recommendations to standing committees in a consistent		
manner. Additionally, ASA supports the engagement of the external methods panel		
to review measures requiring complex methodological analysis, such as risk-adjusted		
outcome and composite measures. ASA recommends that NQF engage both		
physicians and non-physicians in the "methods review" as part of both expert staff		
or the external methods panel to ensure a well-rounded analysis from clinical and		
methodological perspectives.		
NQF should allow stakeholders to vote on individual measures and not require		
voting on all measures within a measure set.		
The current process requires stakeholders to declare support for an entire suite of		
measures, even if only one measure applies to their interests. Stakeholders should		
have the opportunity to indicate "Support," "Do Not Support," "Abstain" or "Not		
Voting" for individual measures within a report. A la carte voting will allow		
stakeholders to vote for or against specific measures within their area of specialty		
and expertise. In previous cycles, an NQF member who voted "Abstain" had their		
vote, for all intents and purposes, count against the measure. ASA appreciates NQF's		
effort to combine comment periods into one continuous public comment period		
throughout each measure cycle as this will reduce redundancy, compared to the		
current process of two comment periods.		
ASA supports increased training and education for stakeholders engaged in the		
Consensus Development Process.		
Education and training are essential to ensure continued success in measure		
development and endorsement processes. The ASA thanks NQF for their routine		

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education and training webinars related to the Consensus Development Process. These activities equip measure developers with tools to efficiently and effectively develop measures suitable for NQF endorsement with the end desire of inclusion in federal payment programs. Thank you for the opportunity to submit comments for your consideration. ASA looks forward to continued work with NQF in the future and appreciates its effort to improve the Consensus Development Process and improve opportunities for measure submission and endorsement. If you have any questions or would like to discuss any of our comments further, please contact Matthew Popovich, Ph.D., ASA Director of Quality and Regulatory Affairs at 202-591-3703 or Leslie Kociemba, M.P.H., ASA Quality Associate at 847-268-9266. They may also be reached at qra@asahq.org.		
The American Psychiatric Association (APA), the leading psychiatric organization in the world, represents more than 37,000 members involved in psychiatric practice, research, and academia representing the diversity of the patients for whom they care. We applaud the National Quality Forum's (NQF) newest effort to "streamline its measure endorsement process and encourage greater participation by consumers, patients, and payers on standing committees" as specified in the "2017 Kaizen Consensus Development Process: Proposed Redesign" draft report. The APA supports the proposed changes included within this redesign. However, we do have some questions about the details involved in some of these potential changes. We applaud the effort made by NQF to provide more than one measure submission opportunity per year to measure developers/stewards. This change will promote a more rapid response to the developers/stewards waiting to learn the status of their measure. Moreover, we welcome the new <i>Intent to Submit</i> period, required of developers/stewards to engage in, before measures are officially submitted into the official endorsement review process. Not only will this process allow NQF to gain insight on what to expect during the official measure submission	Samantha Shugarman, American Psychiatric Association	Thank you for your comment. Scheduling/Frequency: NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). NQF will not formally open a public comment period on the consolidated list of topical areas. However, NQF welcomes your feedback on the list. If you have input, feel free to email NQF at <u>NQFKaizen@qualityforum.org</u> . NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures) and may include one or two additional measures as deemed appropriate. This may vary depending on number of measure submitted; opportunities for related and competing measure review, and measure prioritization

Comment	Commenter	NQF Response
process, but it safeguards developers by ensuring that their measures meet conditions that promote endorsement. Without this step, developers/stewards learn late in the process that their measures didn't include necessary details, only after not being recommended for endorsement by the standing committee. This step creates an efficiency of time that the developers did not previously have. They can either fix the measure or opt out of the endorsement process. Thankfully due to this new step, they will not have to wait years before being able to resubmit for endorsement.		efforts. NQF will notify the steward/developer and provide the date of the next cycle in which there is availability. NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures, while distributing measures to committees with the needed
The draft report states that the number of measures subject to endorsement in each phase is 8 maintenance measures and 4 new measures. Given these limitations on the number of measures subject to the endorsement process, how will NQF prioritize additional new measures, should more than eight be submitted in a year? Given the submission of extra measures, how will NQF ensure that the endorsement review occurs in a timely manner? APA is also interested in learning how the 22 current topical areas will be reduced to 16. Will the public have an opportunity to provide input on these new topical area groups?		expertise to conduct an evaluation. As a result, many of the smaller portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more cross-cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care. Technical Review: Methods Panel: The process includes a two-week process for measure steward or developer to
The APA supports the suggestion to shift responsibility for assigning the degree to which candidate measures meet the NQF scientific acceptability to "NQF staff or an external methods panel, as needed, given their expertise." We hope that this change will implement a high degree of standardization and objectivity for this crucial aspect of the endorsement process. However, with this shift, we anticipate that measure developers/stewards might disagree with the NQF staff or external methods panel on the determination of low or insufficient test ratings. How will the appeal process, or the chance for developers/stewards to make the case for the strength of the test results, be implemented?		respond to the ratings. In addition, the standing committees can still discuss relevant issues, such as risk adjustment. Given the opportunity for more frequent submission, measures may need to be move to the next review cycle to address methodologic concerns. Endorsement Decision: We appreciate your suggestion on the composition of the standing committees. Given current resources and other important strategic considerations, NQF will not be able to implement a change of this magnitude at
Considering that the standing committee mainly consists of clinical experts and the Consensus Standards Approval Committee (CSAC), primarily includes consumers and		this time. If and when this change occurs, we will consider your feedback on the approach.

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payers, the APA supports the NQF's decision to maintain the process by which the		
endorsement decision is currently made. We do agree that consumer and payer		Enhancing Training and Education: Improving the training
perspective, informed by the standing committee expert opinions, have great value		and education for the standing committee members will
on the determination to endorse measures. We would support removing the CSAC,		assist in ensuring they are applying the measure evaluation
should seats for consumers and payers become available on the standing		criteria appropriately when reviewing measures.
committees. However, we would oppose heavily weighting the standing committees		
with consumers and payers, as we feel this could threaten the scientific nature of		
the NQF endorsement process. It is critical that enough seats be available for		
professional members to ensure multi-stakeholder evaluation, endorsement, and		
acceptance by the medical community.		
Lastly, the APA suggests that the NQF develop a better process for standing		
committee members to employ when evaluating "candidate measures" (measures		
under review for endorsement) against the criteria in the NQF Guidance on		
Evaluating Importance to Measure and Report. Though candidate measures are		
currently reviewed based on steps diagramed in a decision tree, the criteria		
described in the decision tree is subjectively interpreted by each standing committee		
member. Clear and consistent criteria for the data elements that standing		
committee participants apply in this process would ensure a more uniform review of		
candidate measures. Unfortunately, some candidate measures have failed to		
achieve endorsement largely because the experts appointed to the standing		
committee had varying interpretations of the NQF Guidance criteria. Standardizing		
this process and providing advance education to standing committee members		
would strengthen the NQF endorsement process and avoid unnecessary rejection of		
meaningful quality measures, and contribute to increasing the NQF compendium of		
measures.		
The APA appreciates the opportunity to comment on these proposed changes, and		
looks forward to the implementation of a more efficient and effective quality		

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measure endorsement process.		
<ul> <li>Thank you for the opportunity to submit comments on behalf of Swain Eng and Associates, LLC (SEA), a healthcare quality measurement and improvement organization focused on helping organizations to act as the catalysts to improvement the quality of care that patients receive. SEA has the privilege to help medical specialty societies and others with the development and testing of quality measures as well as assistance with advancing measures through the NQF consensus development process.</li> <li>We applaud NQF for continually seeking improvements on the endorsement process to make it more nimble, coordinated and faster. SEA would ask that NQF consider answering/clarifying the following questions and comments: <ul> <li>In order to move forward with the suggested changes to the CDP process, will NQF need to seek additional funding (that could delay its implementation)? What does the timeline look like for the phased implementation of the new recommendations to the CDP process?</li> <li>How and who will be involved with the change from 22 to 16 topics areas? Will stakeholders be given the opportunity to provide feedback?</li> <li>As several other commenters have noted, the limitation of 4 new measures per period/8 new measures in a 12 month period may create significant problems. How will you decide/prioritize which 4 new measures to review each period? How will measure sets with &gt;4 measures on a specific topic be reviewed-will they be able to be reviewed in the same period or need to be split up? This change in theory could take longer than the old process when there are greater than 4/8 measures waiting for review pet topic area.</li> </ul> </li> </ul>	Rebecca Swain- Eng, American Academy of Allergy, Asthma and Immunology	Thank you for your comment. We appreciate your feedback on the proposed recommendations for the CDP Redesign. Scheduling/Frequency: NQF's ability to solicit measures for endorsement consideration is contingent upon receiving federal funding. In the final report, NQF has communicated the implementation time frame for each of the proposed recommendations. NQF plans to implement the frequency of submission, the Intent to Submit form and the Scientific Methods Panel, to name a few, by the end of the year. Given the magnitude of some recommendations, such as the endorsement decision and a centralized IT system, implementation will occur at a later date. NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). Topic areas were consolidated with the goal of reassessing and balancing NQF's library of measures, while distributing measures to committees with the needed expertise to conduct an evaluation. As a result, many of the smaller portfolios have been consolidated into cross-cutting topics with a broader range of experience. In addition, some clinical groupings of committees were made to reflect more cross-cutting clinical areas, such as primary care and chronic illness care, pediatrics and geriatrics and palliative care. NQF will not formally open a public comment period on the consolidated list of topical areas. However, NQF welcomes

Comment	Commenter	NQF Response
<ul> <li>"NQF staff will assess whether the measures will require a methodological review based on a set of criteria." What are these criteria? Will stakeholders be able to comment on the criteria?</li> <li>Who and how many individuals do you envision will be on the external methods panel? How will you assess whether they have adequate knowledge/training to provide the necessary expertise?</li> <li>Enhanced Training and Education: "NQF will work to better promote available education offerings to ensure all stakeholders are fully aware of available resources." How are you going to measure that you are improving knowledge about NQF resources and that said resources are stronger?</li> <li>SEA agrees with the Kaizen participants that it is very important that NQF harmonize their CDP submission process with MAP to create a centralized information system that would allow for a comprehensive and longitudinal view of a measure. A user-friendly tool would be a significant benefit to the medical specialty society members of NQF as well as many other stakeholders. SEA understands that this would take significant investment of resources from NQF, however the NQF membership and other stakeholders would greatly benefit from and find great value from a user-friendly centralized system. Given the resource limitations, SEA would encourage the NQF to consider incremental changes or "short term advancements" to move towards this centralized information system as soon as possible.</li> </ul>		<ul> <li>your feedback on the list. If you have input, feel free to email NQF at NQFKaizen@qualityforum.org.</li> <li>NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). NQF may include one or two additional measures as deemed appriopate This may vary depending on number of measures submitted; opportunities for related and competing measure review, and measure prioritization efforts.</li> <li>Technical Review: Methods Panel: NQF will make every effort to standardize how both in-person meetings and web meetings are conducted to ensure consistency in the Committee's measure review and evaluation process. The measure steward/developer will need to notify NQF of their plan submission date. The timing of review for the maintenance measures will depend on when the measures are scheduled to undergo maintenance.</li> <li>In the final report, NQF has clarified the definition of a complex measure. The following types of measures will be considered complex and therefore may require an evaluation by the Scientific Methods Panel:</li> <li>Outcome measures, including intermediate clinical outcomes</li> <li>Instrument-based measures (e.g., PRO-PMs)</li> <li>Cost/resource use measures</li> </ul>
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		Efficiency measures (those combining concepts of
		resource use and quality)
		Composite measures
		Additionally, NQF has also provided additional information
		on the composition and disclosure of interest of the
		Scientific Methods Panel in the final report. The new NQF
		Scientific Methods Panel will consist of 15 to 25 statisticians,
		epidemiologists, psychometricians, economists,
		performance measure methodologists and individuals with
		expertise related to eMeasures and disparities. All nominees
		will complete an annual general disclosure of interest (DOI)
		form, as well as measure-specific disclosures to identify
		recusals from specific measures. NQF will assign measure
		review based on identified conflicts of interest, relevant
		expertise, and availability of panel members. All reviews
		provided by the Scientific Methods Panel will be shared not
		only with the committee but also with the
		steward/developer and the public. Furthermore, the
		Scientific Methods Panel's charge will include providing
		expertise for methods/testing-related issues for NQF and
		advance NQF's guidance on these issues.
		Enhancing Training and Education: NQF currently surveys
		standing committee members on their experience and
		solicits feedback on ways to improve their involvement in
		the CDP. NQF intends expand the audience of the survey (to
		include developers, NQF members and the public) to assess
		the effectiveness of the education and training program.

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		Improvement of Information Exchange and Access: NQF is working to identify short-term solutions to ensure our current IT infrastructure is more user friendly as we progress towards developing a more centralized system. NQF will advance a short-term initiative to aggregate information by grouping MAP measure recommendations and rationale issued each year into one comprehensive and filterable document, accessible from the existing MAP homepage on the NQF website. Similarly, NQF will work to consolidate existing information from CDP reports to make it easier for users to access measure information. NQF will also advance a short-term initiative to improve business rules around publishing timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in NQF's processes. Commenting opportunities will also be enhanced by increasing the character limit to 10,000 characters, real-time updates on comments forwarded to developers, and better regulated public comment periods during evaluation meetings.
On behalf of the Health Resources and Services Administration, we appreciate that	Marlene	Thank you for your comment. We appreciate your feedback
National Quality Forum held a kaizen focused on the consensus development	Matosky, Health	on the proposed recommendations for the CDP Redesign.
process and applaud National Quality Forum for inviting feedback on the	Resources and	
recommendations. We recently submitted measures for review and experience	Services	
many of the challenges presented in the recommendations. We highly encourage	Administration	
and fully support National Quality Forum to follow through in making changes to the		

Comment	Commenter	NQF Response
consensus development process. Thank you.		
<ul> <li>consensus development process. Thank you.</li> <li>Overall, I think the proposed refinements will be beneficial. In particular, offering more frequent opportunities for submission is likely to lead measure developers to wait until their measure is fully mature before submitting. This will make committee reviews more efficient.</li> <li>Regarding the intent to submit plan: what will be the approach taken if more than 12 measures are expected for a particular committee review cycle? How will NQF decide which measures to defer to a later cycle? I think this bears considerations.</li> <li>Regarding the addition of a methods review by a separate body: based on my experience, you do have some deep methods expertise on your existing committees, even if only a few individuals on each committee possess this. I would recommend that (1) your methods review panel include experienced NQF committee members from across different topic areas who possess methods expertise and are very familiar with its application to NQF's CDP. And (2) that you ask those members of each committee who do possess methods expertise to review the recommendations of either NQF staff and/or the methods advisory panel to assess the recommendation in the context of the particular topic area and measure. There may be times that a committee would want to be either more flexible (ie, need for innovation in measurement) or more stringent (ie, many existing measures) in the methods review.</li> <li>I agree that an annual cross-cutting report looking at measure trends, gaps and priorities across top areas would be extremely valuable.</li> <li>Regarding change in the role of CSAC: I would very much encourage NQF to integrate</li> </ul>	Ellen Schultz, American Institutes for Research	Thank you for your comment. Increased Opportunities for Measure Submission: NQF has consolidated the 22 measure review topical areas into 15 topical areas. (The list of the topical areas is included in the final report). NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures), however may include one or two additional measures as deemed appriopate. This may vary depending on number of measures submitted; opportunities for related and competing measure review, and measure prioritization.
consumer and purchaser perspectives more thoroughly into the standing		
committees so that these perspectives are reflected throughout measure review.		

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Given that CSAC rarely overturns the committees' recommendations, relegating these important perspectives to a rubber-stamp role at the end of the CDP does a disservice to both consumers and purchasers.		
Educational materials and support for patient, family and consumer participants in NQF's work is vitally important. I am thrilled to hear that NQF wants to take a leadership position in this important area.		
I agree with the Kaizen participants that there is much need to make information on a measure easier to find. A central repository the links endorsement submissions and MAP would be fantastic. As an early step in this direction, is it possible to at least list where on NQF site a particular measure is referenced, with a link out to relevant reports, excel files, etc?		
Thank you for the opportunity to comment on the 2017 Kaizen Consensus Development Process: Proposed Redesign.	Jane Lucas, Quality Insights	Thank you for your comment. Intent to Submit:
Quality Insights has been developing measures since 2007 and has successfully presented measures to NQF for initial and maintenance endorsement. We were pleased to review the proposed redesign and share our observations and comments. The proposed process of offering two measure submission opportunities for each topic area seems to apply better to new measure endorsement rather than maintenance.		Measure stewards/developers will need to notify NQF at least three months prior to the measure submission deadline to prepare for the committee's review in the upcoming cycle. This will allow NQF to adequately plan for measures in the pipeline and maintenance measures ready for evaluation in the various topic areas. NQF will continue to schedule maintenance measure evaluations based on a three-year cycle from its last endorsement review.
The majority of Quality Insights measures' are submitted under maintenance review and have been reviewed every 3 years. The schedule you are proposing seems to be open in that the measure developer notifies NQF of readiness to submit rather than NQF informing the developer of the assigned project and subsequent timeline. How will "intent to submit" be implemented? Also, it was not apparent in the document if		

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the maintenance cycle will continue to be on the 3 year cycle.		
Quality Insights supports the concept of having trained staff review the Scientific Acceptability of the measure, the proposed change to the public comment process, and the proposed change for the standing committee to determine the endorsement status.		
Quality Insights appreciates the guidance the project staff has provided over the years and we welcome and encourage the proposal of increased training and resources to be included in the redesign.		
Thank you for your time and consideration.		
The Society of Thoracic Surgeons (STS) was pleased to send Mark Antman, STS Senior Manager, Quality Metrics & Initiatives to participate in the recent Kaizen meeting,	Jane Han, Society of Thoracic	Thank you for your comment. Increased Opportunities for Measure Submission: NQF
and we appreciate the opportunity to comment on the draft " <i>Proposed Redesign</i> " report. Thank you in advance for your consideration of our comments.	Surgeons	recognizes the limitations in stakeholder engagement during an in-person versus a web meeting. NQF will make every
Increased Opportunities for Measure Submission		effort to standardize how both in-person meetings and web
We strongly support the proposed redesign of the Consensus Development Process		meetings are conducted to ensure consistency in the
(CDP) to allow for two measure submission opportunities for each topic area per		Committee's measure review and evaluation process.
year. We are also pleased to see that you have reduced the number of topical areas		Endorsement Decision: Thank you for the suggestion. Given
for measure review (and corresponding standing committees) from 22 to 16; we look		current resources and other important strategic
forward to seeing specific information on how the current topical areas have been		considerations, NQF will not be able to implement a change
reorganized. However, we have some concerns related to the proposal to convene		of this magnitude at this time. If and when this change
one measure review cycle per year via an in-person meeting and the other cycle via		occurs, we will consider your feedback on the approach.
virtual web meeting. Measure developer participation in the review process and our ability to respond directly to questions on our measures is enhanced greatly by the		
opportunity to engage directly with standing committee members at the in-person		Improvements in Information Exchange and Access: NQF will
meeting. Substituting a web-based meeting for one review cycle per topic area per		advance a short-term initiative to aggregate information by
Comment	Commenter	NQF Response
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year may introduce some limitations to the full engagement of measure developers and may lead to subtle disparities in standing committee recommendations from one review cycle to the other. The STS recognizes the challenges created by limited resources, but we encourage NQF to use the same meeting format and methodology for <u>all</u> standing committee meetings and review cycles. <u>Intent to Submit</u> We support the proposal to require an "intent to submit" notification from measure developers prior to the measure submission deadline. <u>Technical Review: Methods Panel</u> We support the proposal to move the <i>Scientific Acceptability</i> assessment of		IterIncr Responsegrouping MAP measure recommendations and rationale issued each year into one comprehensive and filterable document, accessible from the existing MAP homepage on the NQF website. Similarly, NQF will work to consolidate existing information from CDP reports to make it easier for users to access measure information.NQF will also work to improve business rules around publishing timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in NQF's processes. Commenting opportunities will also be enhanced by increasing the character limit to 10,000 characters, real-time updates on comments forwarded to developers, and more regulated public comment periods during evaluation meetings.
measures outside of the standing committees to NQF staff or an external methods panel, as appropriate. This change in the CDP will not only accommodate the lack of statistical expertise among standing committee members and encourage greater participation by consumers, patients, and purchasers, it will also promote greater consistency in <i>Scientific Acceptability</i> ratings across measure topic areas and review cycles.		
Measure Evaluation Technical Report – Content and Structure We support the proposal to reduce the amount of measure information included in the technical report, with other information provided either in an appendix or on NQF's public web site.		
<u>Public Commenting Period with NQF Member Expression of Support</u> We support the proposal to consolidate the public comment periods and to eliminate the separate member voting period, in favor of an expanded opportunity for NQF members to comment and express support/non-support for measures.		

Comment	Commenter	NQF Response
Endorsement Decision		
The STS recognizes that elimination of the CSAC role in all measure endorsement		
decisions may not be feasible due to the absence of some stakeholder perspectives		
in the standing committees, as they are typically constituted. However, we		
encourage NQF to consider the more conservative change recommended by Kaizen		
participants: to allow CSAC members to review a list of measures recommended for		
endorsement by the standing committees and to select those they wish to discuss by		
exception. This "consent calendar" approach is analogous to that used by the		
Measure Applications Partnership (MAP) Coordinating Committee and would		
provide an opportunity for the perspectives of the consumer and purchaser		
representatives on the CSAC to be heard and accommodated, without requiring a re-		
adjudication of <u>all</u> endorsement recommendations.		
Adjudication of Appeals		
We agree that shifting the adjudication of all submitted appeals to the CSAC and		
disbanding the Appeals Board is not a feasible recommendation to implement at this		
time.		
Enhancing Training and Education		
We strongly support the proposed enhancements in training and education for all		
stakeholders engaged in the CDP. We are particularly pleased to see the		
recommendation for routine meeting facilitation training for NQF staff and standing		
committee co-chairs, which we agree is needed to improve consistency in the		
measure review process and in the endorsement recommendations made across		
projects.		
Improvements in Information Exchange and Access		
It is certainly understandable that NQF cannot proceed at this time with		
development of a new, centralized information system to link CDP and MAP		

Comment	Commenter	NQF Response
processes – an "ideal-state" and resource-intensive recommendation from Kaizen		
participants. It will be disappointing, however, if NQF does not explore the various,		
smaller-scale innovations that were also proposed at the Kaizen event, such as:		
<ul> <li>using measure information from the CDP to auto-populate data fields and</li> </ul>		
forms for MAP review of the same measures;		
<ul> <li>consolidating duplicative comment periods; and</li> </ul>		
<ul> <li>identifying junctures in CDP and MAP processes at which measure</li> </ul>		
information and status updates can be shared.		
In general, we hope that NQF will take advantage of all opportunities to eliminate		
redundancies in data submission and measure evaluation for any measures that are		
reviewed both for endorsement and for implementation in federal programs.		
Again, thank you for the opportunity to comment on the draft "Proposed Redesign"		
report. If you would like any additional input from STS related to our comments		
above, please contact Mark Antman, DDS, MBA, Senior Manager, Quality Metrics		
and Initiatives, at 312-202-5856 or mantman@sts.org.		
The PCPI is pleased to comment on the National Quality Forum's (NQF) 2017 Kaizen	PCPI Foundation	Thank you for your comment.
Consensus Development Process: Proposed Redesign draft report. While we support		
the refinement of the CDP and the approach NQF has taken in developing this		Frequency/Scheduling: NQF has consolidated the 22
report, we respectfully submit the following comments.		measure review topical areas into 15 topical areas. (The list
		of the topical areas is included in the final report). Topic
Increased Opportunities for Measure Submission		areas were consolidated with the goal of reassessing and
The PCPI shares many of the concerns regarding the current CDP process, including		balancing NQF's library of measures, while distributing
the operational aspects addressed in this draft report as well as timeliness of the		measures to committees with the needed expertise to
endorsement process and the length of time between measure cycles. We are		conduct an evaluation. As a result, many of the smaller
pleased that NQF is taking steps to provide continuous and predictable measure		portfolios have been consolidated into cross-cutting topics
submission opportunities and to condense the measure endorsement process.		with a broader range of experience. In addition, some clinical
		groupings of committees were made to reflect more cross-
NQF plans to consolidate the measure topic areas from 22 to 16 but does not		cutting clinical areas, such as primary care and chronic illness

Comment	Commenter	NQF Response
provide detailed information on what the resulting topic areas will comprise. NQF committees will be convened each year via in-person meeting for one cycle and via a web-based meeting for the second. It is our experience that committee members tend to be more engaged during in-person meetings. We are interested in learning what steps NQF will take to ensure the same level of committee member engagement and discussion is given to measures regardless of meeting type. During each measure submission phase, standing committees will review a maximum of 12 measures – up to 8 measures undergoing maintenance review and up to 4 new measures. PCPI seeks clarification on the distinction between maintenance and new measures. The PCPI has submitted maintenance measures that have undergone considerable review similar to new measures. Furthermore, we are interested in hearing more about how measure prioritization will take place in cases where the number of either maintenance or new measures exceeds these limits but the total number of measures remain within the limit of 12.		<ul> <li>care, pediatrics and geriatrics and palliative care. Individual standing committees that will no longer convene for the following topical areas include:</li> <li>Since 80% of the measures submitted for endorsement consideration are maintenance measures, NQF determined that eight of the 12 measures in each cycle would be maintenance measures.</li> <li>NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures). This may vary depending on number of measures submitted; opportunities for related and competing measure review, and measure prioritization.</li> </ul>
Intent to Submit Currently, measure developers are informed of upcoming measure submission opportunities and provided a list of maintenance measures that are relevant to the topic area and eligible for submission. For maintenance measures, the redesigned process would require substantial coordination and communication between NQF staff and measure stewards/developers. The PCPI strongly recommends that NQF staff indicate which maintenance measures are eligible for submission especially given the consolidation of topic areas. Rather than needing to provide an intent to submit for maintenance measures, stewards/developers can focus on providing the required information for new measures. Endorsement Decision We share the perspective that standing committees should make the final		Technical Review: Methods Panel: NQF will make every effort to standardize how both in-person meetings and web meetings are conducted to ensure consistency in the Committee's measure review and evaluation process. Intent to Submit: NQF will schedule the evaluation of maintenance measures and notify measure stewards and developers in advance. Measure stewards/developers will need to notify NQF at least three months prior to the measure submission deadline to prepare for the committee's review in the upcoming cycle. An intent to submit will signal to NQF of the measure stewards/developers' plan and readiness to submit measures for endorsement consideration.

Comment	Commenter	NQF Response
endorsement decisions without ratification by the CSAC. The standing committees		
are in the best position to make the final endorsement decisions as they have first-		Endorsement Decision: NQF appreciates the comments
hand knowledge of the deliberations that took place for each measure. The PCPI		received on the recommendation of this endorsement body.
encourages NQF to fully consider this proposed change.		However, given important strategic considerations, NQF will
		not be able to implement a change of this magnitude at this
Technical Review: Methods Panel		time. Currently, the CSAC is comprised of a simple majority
Stakeholders recommended removing the detailed technical review and evaluation		of consumers and purchasers. In order to ensure those two
of measures from the standing committee responsibilities. NQF staff or an external		stakeholder perspectives are a key part of the endorsement
methods panel would undertake the technical review depending on measure		process, NQF will need to make certain there is adequate
complexity. The PCPI supports the rigorous yet consistent review of all measures		representation of these groups on each standing committee.
regardless of complexity. We request clarification on how NQF will ensure the		NQF is committed to implementing a plan to identify and
consistent application of the measure evaluation criteria when two groups will		solicit ongoing engagement and participation opportunities
undertake the review independently of each other. We recommend that an external		from these stakeholder groups. Depending on the outcome
methods panel be involved in the review of all measures submitted for consideration		of this initiative, NQF could potentially implement this
of endorsement. Furthermore, we recommend that the external methods panel		proposed change at a later time.
have the expertise to review the technical specifications and feasibility of electronic		
clinical quality measures.		Technical Review: Methods Panel: The new NQF Scientific
		Methods Panel will consist of 15 to 25 statisticians,
Additionally, the PCPI strongly supports the inclusion of the clinical perspective in all		epidemiologists, psychometricians, economists,
aspects of the measure development and consideration of endorsement. Therefore,		performance measure methodologists, and individuals with
we recommend that the standing committee have an opportunity to review and		expertise related to eMeasures and disparities. NQF will
provide input on the external methods panel recommendations before making any		solicit and identify nominees through NQF's standard
endorsement decisions. The measure developer should also be provided the		nominations process. Much like guidance for standing
opportunity to review the recommendations of the external methods panel and		committees, NQF will provide standard guidance on
clarify any questions or concerns regarding the testing methodology and results.		assessing the Scientific Acceptability criterion for a measure,
		using the current decision algorithm from NQF's Measure
Public Commenting Period with NQF Member Expression of Support		Evaluation Criteria. To ensure impartiality, three panel
The PCPI considers the opportunity for public comments an integral aspect in		members will independently evaluate each measure
measure development and endorsement. A continuous public commenting period		undergoing an external panel review. The majority

Comment	Commenter	NQF Response
will provide stakeholders and the general public with ample opportunity to provide input. We have found that we generally receive little pre-meeting comments. If others have similar experiences and receive minimal pre-meeting comments, NQF may wish to revisit the need for this commenting phase.		recommendation will serve as the overall assessment of reliability and validity. NQF will share all evaluations with the measure steward/developer.
Enhancing Training and Education We are very pleased that NQF plans to expand educational and training		
opportunities for standing committee members, NQF staff and measure developers/stewards. Enhanced training and education will help ensure that the NQF evaluation criteria are applied rigorously and consistently across all topic areas and projects. We look forward to seeing this plan in action.		
Improvements in Information Exchange and Access We appreciate NQF's consideration of the recommendation to establish a centralized and comprehensive measure information system. We certainly make use of NQF measure information systems and would support easier access and navigation to find complete measure information across all aspects of the NQF structure. We understand that long-term solutions take time and resources but would support any short-term solutions to enhancing information exchange and access.		
The Joint Commission appreciates the opportunity to comment on the 2017 Kaizen Consensus Development Process: Proposed Redesign. Increased Opportunities for Measures Submission: The Joint Commission supports these changes. We would like a better understanding of the consolidated topic categories. We would also like clarification around maintenance measures and whether the bi-annual consideration includes both annual review as well as the 3 year re-endorsement. Technical Review: Methods Panel: The Joint Commission agrees that the Scientific	JohnMarc Alban, The Joint Commission	Thank you for your comment. Increased Opportunities for Measures Submission: NQF will offer two measure submission opportunities for each topic area each year. However, because there would be more opportunities for submission, NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This was

Comment	Commenter	NQF Response
Acceptability section of the measure submission should be reviewed by statistical		determined given that approximately 80% of the measures
experts with the knowledge and expertise to base a determination on the reliability		submitted for endorsement consideration are maintenance
and validity of the measure. We would suggest that the Methods Panel should be a		measures. The combination of maintenance and new
standing committee across topic areas to promote consistency of interpretation, as		measures may vary depending on number of measures
well as, the feedback provided to measure developers. Furthermore, we are		submitted, opportunities for related and competing measure
recommending that the measure submission process should be amended to allow		review, and measure prioritization efforts. Per NQF's
for submission of the Specifications and Scientific Acceptability sections of the		maintenance of endorsement policy, measures are due for
submission first prior to completion of the entire measure submission form. For		reassessment every three years. NQF will remind measure
measures rated by the Methods Panel as "high" or "moderate" respecting Scientific		stewards and developers of scheduled measure
Acceptability, the measure developer could then complete and submit the other		maintenance review several months prior to the review and
sections of the submission (i.e., Importance, Feasibility, Use and Usability, Related or		notify each of their assigned review cycle.
Competing Measures). These sections could then be scheduled for review by the		
standing committee during the next cycle period. Division of the measure		NQF's portfolio of measures have been consolidated from 22
submission process in this fashion would remove the potential for unnecessary work,		topical areas to 15 topical areas. Topic areas were
conserve measure developer resources, and make the process more user-friendly.		consolidated with the goal of reassessing and balancing
		NQF's library of measures, while distributing measures to
Measure Evaluation Technical Report: The Joint Commission supports the proposed		committees with the needed expertise to conduct an
changes.		evaluation. As a result, many of the smaller portfolios have
		been consolidated into cross-cutting topics with a broader
Public Commenting Period with NQF Member Expression of Support: The Joint		range of experience. In addition, some clinical groupings of
Commission supports the proposed changes.		committees were made to reflect more cross-cutting clinical
Followers and Provide and Advantage of Associate The United Computations and		areas, such as primary care and chronic illness care,
Endorsement Decision and Adjudication of Appeals: The Joint Commission supports		pediatrics and geriatrics and palliative care.
the proposed changes.		
Enhancing Training and Education and Improvements in Information Exchange and		Technical Review: Methods Panel: NQF staff will assess
Access: The Joint Commission supports the proposed changes. We agree with the		whether a measure is sufficiently 'complex' to require a
Kaizen recommendation to create a more consistent, transparent, and user-friendly		methodological review by the Scientific Methods Panel,
tool for submitting, reviewing and analyzing measures and comments.		based on a set of criteria (details below). Because the newly
toor for submitting, reviewing and analyzing measures and comments.		formed Scientific Methods Panel will evaluate the Scientific
		Acceptability of new (and some previously endorsed)

Comment	Commenter	NQF Response
Other Comments The Joint Commission would like clarification on whether evidence criterion is to remain the first vote in the proposed process or if scientific acceptability would be the first pass. The Joint Commission suggests the current submission form process be amended and simplified into a single form format. The current process of completing 3 separate forms includes redundancies that create confusion and consume unnecessary resources on the part of the developers.		<ul> <li>complex measures, measure stewards/developers must submit measure specifications and testing information along with the Intent to Submit form at least three months prior to the measure submission deadline.</li> <li>Other Comments: The <i>Evidence</i> criterion will remain as the first must pass criterion for review during the measure evaluation process.</li> <li>Thank you for the suggestion. As we develop ongoing IT solutions to enhance and improve the measure submission form, we will consider your recommendation.</li> </ul>
<ul> <li>The Association of American Medical Colleges (AAMC or the Association) welcomes the opportunity to comment on the National Quality Forum (NQF)'s 2017 Kaizen Consensus Development Process (CDP): Proposed Redesign. The AAMC represents all 147 accredited U.S. medical schools, nearly 400 major teaching hospitals and health systems, and more than 80 academic and scientific societies. Through these institutions and organizations, the AAMC represents 160,000 faculty members, 83,000 medical students, and 115,000 resident physicians.</li> <li>General questions/comments         <ul> <li>The AAMC has concerns that the NQF is proposing significant changes to the CDP without sufficient opportunity for stakeholders to comment. In addition to a comment period extension, we believe that changes should be made through an iterative process in which NQF staff frequently seek feedback from stakeholders to improve the measure review and</li> </ul> </li> </ul>	Janis Orlowski, Association of American Medical Colleges	Thank you for your comments. We appreciate your feedback on the proposed recommendations for the CDP Redesign. NQF hosted the Kaizen event in collaboration with CMS to inform the CDP redesign. CMS, as the funder of this initiative, has asked NQF to solicit public comment on the proposed recommendations and provide a final report outlining the new CDP by July 1, 2017. Thus, NQF had to limit the amount of time NQF members and the public had to provide feedback. However, as NQF continues to plan for implementation of the new CDP, additional feedback is welcomed. Increased Opportunities for Measure Submission: Standing Committees will meet more often to allow for more frequent
<ul> <li>NQF should provide additional details on its website to allow stakeholders to more easily understand the specifications and history of any single</li> </ul>		measure submissions. However, the time needed for committee reviews remains unchanged. NQF has consolidated the 22 measure review topical areas into 15

Comment	Commenter	NQF Response
quality measure. On the NQF-Endorsed Standards section of the website,		topical areas (The list of the topical areas is included in the
users should be able to clearly see which project the measure is currently		final report). Topic areas were consolidated with the goal of
placed in, the full measure specifications and developer discussion notes,		reassessing and balancing NQF's library of measures, while
whether the measure was reviewed during the trial period, and other		distributing measures to committees with the needed
pertinent information.		expertise to conduct an evaluation. As a result, many of the
• NQF should take steps to improve the email notification process. Members		smaller portfolios have been consolidated into cross-cutting
should have the opportunity to receive emails for specific projects, and all		topics with a broader range of experience. In addition, some
of the distribution lists should be easy to find on the website. Reminders for		clinical groupings of committees were made to reflect more
upcoming comment period deadlines should also be an option for		cross-cutting clinical areas, such as primary care and chroni
members.		illness care, pediatrics and geriatrics and palliative care
Increased Opportunities for Measure Submission		Due to the increased workload for the standing committees
• AAMC supports greater opportunity for developers and stakeholders to		NQF cannot accept more than 12 measures per cycle per
submit and review measures. We have concerns, however, that the CDP		topical area, however may include one or two additional
changes may force the committees to make final decisions on a measure		measures as deemed appropriate. Any Intent to Submit
without sufficient review. NQF should ensure that there is an iterative		forms that are submitted once capacity has been reached f
process where disagreements and concerns can be addressed before a final		a particular cycle, NQF will notify the steward/developer ar
decision is made.		provide the date of the next cycle in which there is
• NQF plans to decrease the topic areas from 22 to 16 – however, no detail is		availability.
provided on these 16 topic areas or how this will impact the number and		
composition of the Standing Committees. In addition, how does this impact		Intent to Submit: NQF will schedule the evaluation of
the proposed 12 measures per cycle limit since the number of measures in		maintenance measures and notify measure stewards and
some of the topic areas may increase with this shift?		developers. Any Intent to Submit forms that are submitted
		once capacity has been reached for a particular cycle, NQF
Intent to Submit		will notify the steward/developer and provide the date of
What will be the process for measure maintenance? Will NQF notify		the next cycle in which there is availability.
developers when maintenance is due?		Tasknisel Daview Matheda Davak The new NOS Scientific
• What will be the process when NQF receives more than 12 measures during		Technical Review: Methods Panel: The new NQF Scientific
a cycle review period? Who will decide which measures get moved to the		Methods Panel will consist of 15 to 25 statisticians,
		epidemiologists, psychometricians, economists,

Comment	Commenter	NQF Response
next cycle?		performance measure methodologists, and individuals with
		expertise related to eMeasures and disparities. NQF will
Fechnical review: Methods Panel		solicit and identify nominees through NQF's standard
• AAMC supports the creation of the technical review panel, however, we		nominations process. Much like guidance for standing
have some questions we ask NQF to address:		committees, NQF will provide standard guidance on
• Who will be on this panel?		assessing the Scientific Acceptability criterion for a measure
<ul> <li>Will it be similar to how the Standing Committees operate?</li> </ul>		using the current decision algorithm used from the measure
<ul> <li>Will there be one methods panel or many panels?</li> </ul>		evaluation criteria. To ensure impartiality, three panel
<ul> <li>What opportunity will developers have to provide additional</li> </ul>		members will independently evaluate each measure
information or clarify questions?		undergoing an external panel review. The majority
<ul> <li>Will Standing Committee members be able to "overrule" the</li> </ul>		recommendation will serve as the overall assessment of
methods panel or NQF staff and change the rating on reliability		reliability and validity. NQF staff will send the preliminary
and/or validity?		analysis to developers for review prior to finalizing and
<ul> <li>The document states that, "Generally, NQF will not forward</li> </ul>		sending to the standing committee. If developers disagree
measures with a 'low' or 'insufficient' rating from the methods		with the staff or Scientific Methods Panel review or ratings,
review to the committee for further evaluation". What criteria		they can use the two-week review period to provide
would be used to determine if a measure with those ratings did or		additional clarification, which can be considered by staff
did not move forward to a committee?		when finalizing the preliminary analysis. Developers will also
		have the opportunity to introduce their measures during th
Public Commenting Period with NQF Member Expression of Support		committee evaluation meeting and answer questions from
The AAMC requests clarification on how comments submitted after the		the committee during the discussion.
evaluation meeting be incorporated into the measure recommendation		
decision. If recommendations are revised due to these comments, how will		Measures will be rated by the Scientific Methods Panel and
that be reflected?		NQF staff against the measure evaluation criteria. Standing
		committees may raise concerns with the specifications of the
Indorsement Decision		measure or with potential threats to validity (e.g., selection
AAMC has concerns with allowing the standing committees to make the		of variables for risk adjustment model) and can therefore
final decision on measure endorsement, since the standing committees may		overturn the staff or Scientific Methods Panel rating. As par
be inconsistent in their evaluation process. The AAMC agrees with keeping		of its ongoing education efforts, NQF will provide clear

Comment	Commenter	NQF Response
the CSAC in place as an oversight body for the time being.		guidance to standing committees regarding the
		circumstances wherein an overturn of the rating would be
Improvements in Information Exchange and Access		permissible.
• Since NQF is limited in the changes that can be made at this time but		
commits to working on short-term solutions, it would be preferable if they		Public Commenting Period with NQF Member Expression of
solicit input from the various stakeholders involved in the CDP and MAP as		Support: NQF has provided clarification regarding the
they move forward in this effort.		developer's role in responding to comments during the
		commenting period. NQF will ensure the measure developer
		receives the submitted committees in order to prepare for
		the measure evaluation meeting. Measure developers will
		not be required to provide written responses to the
		comments received prior to the measure evaluation
		meeting. The committee will review any comments received
		after the committee evaluation meeting during the post-
		commenting period call. All submitted comments during this
		time will receive written responses from the standing
		committee, measure developers, and/or NQF, as
		appropriate. The standing committee may revise its
		recommendations in response to a specific comment or
		series of comments submitted during this phase of the
		process.
		Endorsement Decision: Kaizen participants recommended
		that standing committees make the final endorsement
		decisions, without ratification by the CSAC. Participants
		noted that the CSAC rarely overturns the measure
		recommendations of the committee. NQF appreciates
		comments on the recommendation of this endorsement
		body. However, given important strategic considerations,
		NQF will not be able to implement a change of this

Comment	Commenter	NQF Response
		magnitude at this time. Improvements in Information Exchange and Access: NQF will advance on short-term initiatives to improve business rules around publishing timelines and meeting materials, to ensure developers, committee members, and members of the public are more aware of opportunities to participate in NQF's processes.
On behalf of our nearly 5,000 member hospitals, health systems and other health care organizations, and our clinician partners – including more than 270,000 affiliated physicians, 2 million nurses and other caregivers – and the 43,000 health care leaders who belong to our professional membership groups, the American Hospital Association (AHA) appreciates the opportunity to comment on the National Quality Forum's (NQF) proposed redesign of the consensus development process (CDP). The measure endorsement process has been at the core of the NQF's work since its inception. Health care providers, consumers, and public and private payers engage with the CDP to help identify those quality measures that are sufficiently important, scientifically sound, useful and feasible. However, the execution of the CDP is a daunting task given the heterogeneity of health care, the high demand for measures in accountability applications, the lack of consistency in data infrastructure across the health care system, and the ever-evolving science of quality measurement. These factors, among many others, make it more challenging to achieve a timely, consistent CDP that permits the meaningful engagement that members want and that NQF's position as a voluntary national consensus-standards body demands.	Nancy Foster, American Hospital Association	<ul> <li>Thank you for your comment. We appreciate your feedback on the proposed recommendations for the CDP Redesign.</li> <li>This revised process is designed to allow more opportunities for public input and measure discussion and to ensure best practices in building consensus for performance measurement and standards-setting are put into place.</li> <li>While the final report provides descriptions of proposed processes to the extent possible, there are many details that may change as implementation continues. NQF will continue to keep all stakeholders informed and will solicit ongoing feedback as deemed appropriate.</li> <li>If you have specific input or suggestions, feel free to email NQF at <u>NQFKaizen@qualityforum.org</u>.</li> </ul>
The AHA appreciates NQF's commitment to improving its process, and we believe some of the CDP redesign ideas outlined in the draft report merit serious		

Comment	Commenter	NQF Response
consideration. In particular, we generally support NQF's efforts to streamline its		
technical report content, improve information access, and enhance education.		
However, we are concerned that the report's recommendations are far too		
narrowly focused on improving the timeliness of the CDP, and may undermine the		
CDP's consistency and the ability for stakeholders to engage. We also question the		
feasibility of using a single methodological panel to support the review of up to		
384 measures per year. Lastly, we are disappointed that the CDP redesign misses		
the opportunity to improve NQF's process for identifying and cultivating measures		
that will advance our understanding of quality and safety and its process for		
selecting best-in-class measures. The NQF's engagement on these issues is urgently		
needed as hospitals, other providers, and the public are drowning in overlapping,		
conflicting measurement that takes time away from what matters most –using		
measures to improve care.		
As we understand it, NQF would move to a model in which 16 standing committees		
would conduct two measure reviews per year, with up to 12 measures in each (i.e.		
up to 384 measures per year). In addition, public comment would be a continuous		
process paired with voting in which stakeholders could change their vote at any		
time. Lastly, each standing committee would be the final arbiter of endorsement,		
and the CSAC would become an "appeals board."		
We appreciate the desire among some stakeholders to get measures through the		
NQF endorsement process more quickly. However, we seriously question the ability		
for NQF members to track 16 committees at once and continually update their		
votes. Before embarking on such a change, NQF's website and communications must		
be enhanced to ensure members can select the committees and measures they care		
about, and receive sufficient notice about a measure being "active" in the review		
process to ensure they can comment. This infrastructure is especially important		

Comment	Commenter	NQF Response
because opportunities other than the NQF endorsement process for providing input		
on measures abound – including CMS proposed rules, measure developer requests		
for comment, the JIRA tool used to provide input on electronic clinical quality		
measures (eCQMs), and so forth. Implementing this new process without the		
needed underlying infrastructure could impede its success.		
Yet, even better infrastructure likely will be insufficient to enable all interested		
parties to follow the NQF endorsement activities in which they have an interest if		
there are nearly 400 measures being processed each year. The intended virtue of		
developing standards through a voluntary consensus development agency is that at		
the end of the review, all interested stakeholders are supposed to agree that the		
standard is the best in class approach available. Without sufficient engagement, that		
will not happen.		
We observed a couple of years ago that the routine level of engagement of		
organizations in commenting on and voting on most measures processed by NQF is		
extremely low. We worry that the implied expectation of continuous involvement in		
measure endorsement process envisioned by these proposals would prove too		
burdensome for even those of us who are routinely following NQF's work. Without		
the multi-stakeholder engagement that NQF was created to foster, the steering		
committees become just another technical expert panel with a different name. We		
urge NQF to think about ways in which to foster greater stakeholder		
involvement. We would be disappointed if these proposed changes in the CDP		
further diminished engagement.		
The AHA also believes the concept of using a "methods review" panel has merit,		
particularly for highly complex outcome measures whose submissions are		
accompanied by a large amount of testing information. However, given that the		
health care field is increasingly moving towards outcome measures, as well as		

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measures collected and reported using electronic health records, we fear that a		
single review panel could become a process bottleneck. Before fully supporting this		
idea, we urge NQF to provide more detailed information about the circumstances		
under which the panel will be used, what kinds of expertise will be needed on it,		
how it will be resourced, and how quickly they will be expected to complete their		
reviews.		
Lastly, the AHA strongly urges NQF to take more aggressive steps to select "best-		
in-class" measures. The endorsed measures are called voluntary national consensus		
standards that become broadly adopted to reduce the burden of data collection and		
to provide one "source of truth" about their relative performance of providers.		
However, there are numerous instances in which NQF has endorsed multiple		
measures assessing the same aspect of care, and reached the conclusion that one		
measure is not clearly superior to the others. While we agree that it is entirely		
plausible that such "competing measures" can have strengths and weaknesses that		
make it difficult to choose one based solely on merit, NQF must choose anyway.		
To endorse more than one measure of an aspect of care defeats the concept of a		
voluntary national consensus standard. According to the American National		
Standards Institute, voluntary consensus standards work behind the scenes to make		
everyday life work. They define the size, shape and information contained on bank		
cards so that they can be used at any ATM in the world. They describe the size and		
dimensions of the end of a light bulb so that it will fit into a socket made by any		
manufacturer. They define the spacing on railway tracks so that a train car made by		
any manufacturer will ride the rails smoothly. In the same way, the voluntary		
standards set by NQF should identify and define key aspects of measuring quality so		
that attention can be focused on improving quality.		
On behalf of over 18,000 board-certified orthopaedic surgeons, the American	Neha Agrawal,	Thank you for your comment.
Academy of Orthopaedic Surgeons (AAOS) would like to offer comments on the NQF	The American	

Academy of Orthopaedic Surgeons	Increased Opportunities for Measure Submission: NQF has consolidated the 22 measure review topical areas into 15 topical areas. The list of the topical areas is included in the final report. NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted; opportunities for related and competing measure review,
	topical areas. The list of the topical areas is included in the final report. NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted;
Surgeons	final report. NQF will limit the number of measures evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted;
	evaluated by the standing committees in each cycle to a maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted;
	maximum of 12 (up to eight measures undergoing maintenance review and up to four new measures). This may vary depending on number of measures submitted;
	maintenance review and up to four new measures). This may vary depending on number of measures submitted;
	vary depending on number of measures submitted;
	opportunities for related and competing measure review,
	and measure prioritization efforts.
	Public Comment Period: In place of two separate public
	commenting periods (14-day pre-meeting commenting and
	30-day post-meeting commenting), NQF will have one
	continuous public commenting period. This will allow
	sufficient time for the public and NQF membership to submit
	comments on measures under review. Comments received a
	week prior to the measure evaluation meeting will be
	submitted to the Committee for their consideration.
	Enhanced Training and Education: Thank you for the
	suggestion. As we develop our training and education plan to
	educate and inform measure developers, we will consider
	your recommendation.

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Technical Review: Methods Panel: The NQF proposes to conduct a "methods review"		
internally or via an external technical advisory panel, for the scientific acceptability		
section of a measure. The NQF staff and/or the methods panel would provide their		
review, ratings, and comments of the technical aspects of reliability and validity		
analyses and results, to the relevant standing committee. The standing committees		
will still ultimately make a recommendation. The AAOS agrees with this proposal, as		
a standing committee of physicians may not have the time or may not be as qualified		
to evaluate the scientific acceptability of a measure. Physicians on the standing		
committee can focus more on their area of expertise, as it relates to review of the		
measure. This change would be especially good for complex outcome measures,		
which require a more thorough evaluation of the scientific acceptability.		
Measure Evaluation Technical Report: The NQF proposes to reduce the amount of		
information provided in the technical report, with the remaining background		
information on the topic area to be included on NQF's public website. The AAOS		
agrees with this proposal, and in particular supports having an annual report which		
summarizes endorsement activities and identifies prioritized gaps in measurement		
across all topic areas.		
Public Comment Period: The NQF proposes to have one continuous public		
commenting period spanning 12 weeks, in place of two separate public comment		
periods spanning six weeks. The AAOS requests additional detail on how "earlier and		
more continuous expression of support/non-support from NQF members" will have		
a "more significant impact on the measure evaluation," per the proposed redesign.		
The AAOS also requests more detail on how increasing the public comment period		
from 6 weeks to 12 weeks contributes to the NQF goal of reducing overall		
endorsement time to about six months.		
Enhanced Training and Education: The NQF proposes to expand and strengthen the		

Comment	Commenter	NQF Response
current range of educational resources offered for staff, committee members, and		
measure developers, including on-demand virtual references, developer educational		
webinars, written guidance materials, consumer/patient-focused webinar training,		
and meeting facilitation training. The AAOS agrees with this recommendation and		
urges the NQF to keep to their schedules as best as possible, as there have been		
many monthly webinars cancelled at the last minute. The AAOS requests that the		
NQF identifies a staff point person for each measure steward/developer to improve		
communication between measure stewards/developers and the NQF. Currently the		
AAOS sends an e-mail to a general e-mail box and historically the technical support		
has been limited and vague. We urge the NQF to offer specific instructions and		
guidance on the submission process.		
Improvements in Information Exchange and Access: Kaizen participants		
recommended a centralized information system that would allow for a		
comprehensive and longitudinal view of a measure, including real-time updates and		
Measure Applications Partnership (MAP) and CDP data accessible by staff,		
developers, and the public. The NQF notes that it will not be able to implement this		
change at this time, given available resources and other strategic considerations. The		
AAOS agrees with this Kaizen recommendation and urges the NQF to prioritize this		
as a long-term goal as it would have significant positive benefits to both NQF and		
measure stewards/developers.		
Thank you for considering our comments on this important matter. If you have any		
questions, please do not hesitate to contact our Medical Director, William O. Shaffer,		
MD at (202) 548-4145 or shaffer@aaos.org.		
KCP appreciates the opportunity to comment on NQF's 2017 Kaizen Consensus	Frank Maddux,	Thank you for your comment.
Development Process Proposed Redesign Draft Report. KCP is a coalition of members	Kidney Care	
of the kidney care community that includes the full spectrum of stakeholders related	Partners (KCP)	Methods Panel for Technical Review: NQF appreciates your
to dialysis care—patient advocates, healthcare professionals, dialysis providers,		recommendations and have included these concerns within

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researchers, and manufacturers and suppliers—organized to advance policies that		the final report. For complex measures, the Scientific
improve the quality of care for individuals with both chronic kidney disease and end		Methods Panel will evaluate the measure's reliability and
stage renal disease. We commend NQF for undertaking this important work and		validity (or Scientific Acceptability criterion) and provide a
offer comment on one proposed CDP revision—creation of a technical advisory		preliminary recommendation to NQF staff and the standing
panel to assist in conducting methodological reviews of complex measures—and		committee. Because updated reliability and validity testing is
recommend one additional area we ask NQF to address in the CDP redesign.		not required for maintenance measures, NQF staff will
		review previous testing results for complex maintenance
Generally speaking, KCP agrees that the creation of a methodology panel would		measures and determine the adequacy of prior testing. If
enhance the NQF process and better align the consideration of statistically complex		prior testing is inadequate, updated testing is provided, or
measures that might otherwise be evaluated dissimilarly across different standing		NQF staff deems an external review necessary, the measure
committees with variable statistical expertise. We thus support NQF's proposal to		will be submitted to the external Scientific Methods Panel to
utilize a methods panel to assist in conducting methodological reviews of complex		evaluate the reliability and validity of the measure.
(e.g., risk-adjusted outcome, composite, cost) measures. However, we are deeply		Following the current process, NQF staff will perform a
concerned about the proposal that NQF staff would review the Scientific		preliminary analysis against all of the other evaluation
Acceptability criterion of all endorsement maintenance measures—including		criteria for both new and maintenance measures. For non-
complex measures. Under this scenario, previously endorsed complex measures		complex measures (e.g., structure and process measures),
would be held to a less rigorous statistical standard than similar or related newly		NQF staff will complete the preliminary analysis against all
submitted metrics, potentially creating methodological inconsistencies in the		measure evaluation criteria, including the Scientific
publically-reported and penalty-based systems within which such measures are		Acceptability criterion.
frequently deployed. Currently-endorsed measures that a knowledgeable		
methodology panel might readily identify as statistically-flawed or ineffective at		For both complex and non-complex measures, when the
identifying statistically significant and meaningful differences in performance might		preliminary analysis is complete, NQF staff will send the
"pass" a staff review of Scientific Acceptability requiring only an attestation as to		preliminary analysis to developers for review. Measures
"the adequacy of prior testing." To address this concern, we request that NQF		rated by NQF staff or the Scientific Methods Panel as "Low"
amend this proposal to also require review by the methodology panel of two		or "Insufficient" for reliability or validity will be removed
additional categories of measures:		from the current evaluation cycle, allowing time for any
1. All complex measures undergoing maintenance review for which there are		additional testing, clarification or NQF technical support, or
performance data and/or new or updated testing data.		review prior to consideration of the measure in a future

2. Any measure for which a standing committee member moves to request a

	cycle. NQF will continue to follow the same ad-hoc process. An ad hoc review may be carried out at the same time as an active measure review cycle. This will minimize committee and developer burden in managing various reviews under different schedules.
	hoc review may be carried out at the same time as an active measure review cycle. This will minimize committee and developer burden in managing various reviews under
lleen cKiernan, The win Group	Thank you for comment. Increased opportunities for measure submission: NQF will make every effort to standardize how both in-person
	meetings and web meetings are conducted to ensure consistency in the Committee's measure review and evaluation process. Intent to Submit: NQF will provide a schedule for all maintenance measures in advance to allow developers

Comment	Commenter	NQF Response
the dates by which reviews of the literature, updates to the specifications, and measure testing must be completed to present Steering Committees the most accurate and up-to-date evidence available during measure review. Lewin encourages NQF to ensure in-person and virtual reviews are as similar as possible to help standardize the evaluation of each criterion for measures discussed on site and via webinar; examples of some scenarios in which disparate reviews could occur include voting during the meeting vs. polling Standing Committee members after the webinar (the latter of which prevents some voters from having context for results or asking questions) and ensuring the meeting facilitation for in-person and virtual webinars is as similar as possible.		measures are due every three years from its last endorsement review. Developers are required to maintain their measure in accordance with the Measure Steward Agreement. Failure to do so may result in removal of endorsement, however NQF will continue to identify potential options with the developer prior to removing endorsement. Technical Report Content and Structure: Thank you for your comment.
<ul> <li>Intent to Submit: Lewin supports creation of an Intent to Submit form to pipeline measures that will come forward for review by a Standing Committee, moving forward. We encourage NQF staff to build in additional transparency to the timeline by which maintenance measures must be reviewed before losing endorsement to ensure that measure stewards have multiple CDP options to which they could submit.</li> <li>Technical Report Content and Structure: Lewin supports the streamlining of the evidence presented therein into a short, more usable document. Lewin also favors</li> </ul>		Methods Panel for Technical Review: The complexity of a measure, complex vs. non-complex, will be based on information provided in the <i>Intent to Submit</i> form. Measures that are considered complex may require an evaluation by the Scientific Methods Panel. For both complex and non-complex measures, NQF staff will send the preliminary analysis to developers for review prior to finalizing and sending to the standing committee. A flowchart illustrating this process is included in the final
preparation of a cross-cutting annual report in which themes from CDPs held each year and gaps identified by Standing Committees are summarized.		report. Standing committees may raise concerns with the
Methods Panel for Technical Review: Lewin supports the creation of a methods panel for review of complex scientific acceptability submissions. We would appreciate clarification on how and when the method panel's feedback will be built into the CDP process to help estimate the additional level of effort required by stewards/developers for this second level of review. We also encourage representatives from the methods panel to liaise directly with Standing Committees		specifications of the measure or with potential threats to validity (e.g., selection of variables for risk adjustment model) and can therefore overturn the staff or Scientific Methods Panel rating. NQF will provide the Scientific Methods Panel and standing committees education and training on changes to the process and expectations on their roles and updated written guidance documents. As part of

Comment	Commenter	NQF Response
(attending the in-person meetings/webinars for each measure's review) to answer questions posed by SC members. We recommend NQF prepare a Methods Panel guidebook, similar to the documents used by SCs during the CDP process, to help standardize the approach methods panel members take to reviewing <i>Scientific</i> <i>Acceptability</i> results. We also suggest NQF create an algorithm or flowchart to clarify how measures will be selected for review by the methods panel (vs. by NQF staff); this selection process should include discussion of the appropriate review body with the measure steward. Finally, we suggest NQF to stand up a process for Standing Committees to override the vote of a methods panel or NQF staff member if they interpret the reliability and/or validity findings for a measure differently than is recommended.		<ul> <li>its ongoing education efforts, NQF will provide standing committees clear guidance regarding the circumstances wherein an overturn of the rating would be permissible.</li> <li>Public Comment Period: NQF will make every effort to provide developers with public comments in real-time to the extent possible.</li> <li>Enhancing Training and Education: Thank you for your comment.</li> </ul>
Public Comment Period: Lewin supports the revised approach proposed by NQF to hold a single, continuous comment period during which members of the public could submit feedback on measures undergoing endorsement review. Lewin encourages NQF to provide as much time as possible to measure stewards and developers to prepare responses to public comments to ensure that we provide the most meaningful, well-thought-out feedback to comments as is possible; Lewin suggests delivering comments to measure stewards in near-real time (or as rapidly as is feasible) to maximize our response time.		
<b>Enhancing Training and Education:</b> Lewin agrees that NQF should provide as many public-facing education resources as possible, including increased promotion of technical assistance support for measure developers, on-demand and real-time webinars for those new to the measure development/NQF submission processes, and creation of orientation sessions for those less experienced in submitting measures to NQF. Lewin also supports standardization of NQF staff competencies through trainings on meeting facilitation and other shared operational skills that can apply across Committees.		

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Information Exchange and Access: It would be helpful if this section were separated into short term and long term actions rather than listing everything in this section collectively as not feasible. The revision to the submission form should be called out as a key recommendation. Overall this is a very concise summary of the key themes of the meeting. However, I would suggest adding a summary section that would detail how the proposed short- term changes would impact the overall endorsement timeline. A key theme/goal of the meeting was reducing the length of the process. It would be helpful to highlight if a reduction in length was achieved given the changes NQF is committed to implementing short term. Further, I believe separating into short term and long term actions throughout versus the statements concerning current feasibility would be more strategic.	Kyle N. Campbell, Health Services Advisory Group, Inc.	Thank you for your comment. We appreciate your feedback on the proposed recommendations for the CDP Redesign. Information Exchange and Access: NQF will adopt a two-fold approach to addressing recommendations from Kaizen participants. Some aspects of the recommendations are resolvable through short-term solutions and adaptations of existing platforms. Other recommendations will be addressed through a long-term product development approach. This is outlined in the final report. This revised process is designed to allow more opportunities for public input and measure discussion and to ensure best practices in building consensus for performance measurement and standards-setting are put into place. While the final report provides descriptions of proposed processes to the extent possible, there are many details that may change as implementation continues. NQF will continue to keep all stakeholders informed and will solicit ongoing feedback throughout this process.