Guidelines for Quality Measure Stewards and Developers to Test for SDOH/SES Effect

Special Needs Plans Described
SNPs are a subset of Medicare Advantage (MA) plans specifically authorized and designed to meet special care needs of Medicare beneficiaries. The plan types and subgroups include:

- **Chronic condition SNPs** (C-SNPs): serving persons with certain severe or disabling chronic conditions (e.g., HIV/AIDS, chronic heart failure, COPD, mental illness)
- **Institutional SNPs** (I-SNPs): serving persons residing in nursing homes or with comparable care needs in the community.
- **Dual eligible SNPs** (D-SNPs): serving persons covered by both Medicare and Medicaid.
- **Fully Integrated Dual Eligible SNPs** (FIDESNPs) and Medicare-Medicaid Plans (MMPs) – which are a specific type of D-SNP, providing both Medicare and Medicaid benefits, including long-term services and supports.

While SNPs are regulated, evaluated, and paid on the same basis as other MA plans, they are required to provide additional benefits and services to their target populations and to implement tailored care management according to unique Models of Care that serve every enrollee.

Quality Measures Background
**Quality Ratings are Tied to Payment**
In 2012, CMS began to implement the MA Star Rating system, which makes quality incentive payments to plans that obtain at least a 4-Star rating under a 5-Star rating system. Currently, plan ratings are based on 47 performance measures derived from HEDIS, CAHPS, and HOS instruments, and from CMS administrative data.

**Need for Recognition of Factors Affecting Health Outcomes**
The SNP Alliance supports pay-for-performance as a tool to improve care for Medicare beneficiaries. However, the current system ignores the reality that poverty, low levels of education, disability status, dual eligible status, and other social determinants of health (SDOH) affect outcomes and Stars.

**Mounting Evidence Supports Need for Risk Adjustment**
There is compelling new evidence from the Assistant Secretary for Planning & Evaluation (ASPE) 2016 Report to Congress and from the National Academies for Sciences, Engineering, and Medicine, that socio-economic status of enrollees affects Star measure outcomes and thus adversely impacts a health plan’s ability to achieve excellence under the Star Rating system.

In 2015 and 2016 the National Academies Committee on Accounting for Socioeconomic Status in Medicare Value-based Payment Programs conducted a thorough examination of socioeconomic and social determinant of health risk factors (called “social risk factors”) and found many impacted outcomes and should be taken into account in quality measurement and value-based payment. They released a series of 5 reports.

In December 2016, ASPE released their Report to Congress and found that dual-eligibility status was the most significant predictor of poor health outcomes as measured by the Medicare Star Ratings system. Furthermore, dual status, low income status, and disability status (used as proxies for SES and other social risk factors) impacted outcomes— independent of plan or provider behavior/actions.

Previously, findings from a CMS-commissioned RAND study provided evidence that a beneficiary’s dual-eligible status significantly lowered outcomes on 12 of 16 Star Rating measures. It also found that disability status significantly lowered outcomes on 11 of 16 measures. A 2015 Inovalon study found that characteristics of dual-eligible enrollees explained 70% or more of the disparity in outcomes compared to non-dual eligible enrollees on five of eight measures.

**Congress Urges CMS to Modify Stars to Account for SES**
Members of Congress have urged CMS to modify the Star Ratings system to better account for the clinical and socio-demographic risk factors that are out of a plan’s control, arguing that MA performance measurement should accurately reflect the challenges in caring for low-income, chronically ill people.

**CMS Agrees, but No Changes in Risk Adjustment or Measures or Measurement Approach are Proposed in 2017**
CMS recognizes that risk adjustment is needed “because the existence of risk factors before or during healthcare encounters may contribute to different outcomes, independently of the quality of care received” (Source: CMS, Blueprint for the CMS Measures Management System, Version 12.0, May, 2016).
However, the limited approach CMS used in 2016—applying a modest factor in just 6 of 47 measures—has had almost no effect. Only 19 plans out of several thousand had small changes in their Star ratings last year when the “Categorical Adjustment Index” was put in place by CMS.

**Problem Statement**
There are underlying problems with both the measures within Stars and with a lack of SES adjustment. Many of the measures have not been sufficiently tested in a low-income, diverse, high social risk factor population. Therefore the measures themselves may be problematic. On top of this, the measure adjustments for social risk factors are minimal. We suggest that a set of guidelines from CMS for measure re-testing and for measure developers would help address part of this problem.

**CMS Has Not Set Guidelines or Standards**
CMS has stated that it is the measure developer’s responsibility to determine if patient/individual SDOH factors should be accounted for to accurately compare plan performance. This agency also leaves it to the developers to determine how to apply these factors in their measurement specification.

While CMS offers guidance about attributes of risk adjustment models, there are no requirements for: (1) the sample size and definition, (2) a minimum data set and sources that accurately capture SDOH characteristics of patients, (3) a minimum set of SDOH variables to include in the risk adjustment testing/models, (4) specific analytic methods to be used, or (5) disclosure in a way that is useful to researchers to replicate (validate) the work and also in a way that is understood by the lay public.

**Limited Scope**
CMS guidance to measure developers and stewards only refers to selecting variables that are clinically relevant, where research has consistently shown that social determinants of health—such as income level/poverty status, dual eligible status, disability, living alone, housing transience, education level—may be equally important to achieving good health outcomes and should also be tested as risk adjustment factors in the models.

**Potential for Harm**
The SNP Alliance believes the lack of guidance around minimum requirements for testing these and other attributes creates an environment where models and testing may not be sufficiently rigorous and could lead to faulty conclusions regarding the measure and its adequacy for application in the field with certain populations, particularly those who have high social risk factor issues, and multiple chronic conditions. Unadjusted measures used in the field could therefore yield inaccurate results about providers and plans who serve a disproportionate number of individuals with these characteristics. These quality ratings can influence consumer behavior and are tied to payment. This may inadvertently negatively impact providers and plans serving the most vulnerable population subgroups.

**Solution**
Creating (and making public) one consistent set of standards with minimum parameters for sample size, variables, data definitions, data sources, and methods for testing the measures, would provide critical guidance needed so that key stakeholders can be assured that the developers/stewards have performed risk adjustment evaluation with due diligence and used a sound approach, and so that there is full transparency to the public. We have therefore reviewed the literature and worked with those in the field to craft a set of parameters and make recommendations—for consideration by CMS.

**Recommendation**
We encourage CMS to set a **minimum standard** for measure developers and stewards to **consistently test** their measures—especially among the dual subpopulation groups who are not like the majority Medicare population and who represent the highest cost, most complex Medicare beneficiaries—to ensure that the measure accurately portrays experience in the intended area of focus, and to reveal areas where adjustment is needed.

This should include the follow requirements:
- **Sampling** - a minimum sample size and diversity within the population used for testing that would include at a minimum younger disabled individuals (18-64) and older adults with multiple chronic conditions, as well as include subgroups reflecting the enrollment from a variety of health plan types.
- **Unit of Analysis** – utilize small geographic areas as units of analysis when testing measures. Studies show that variances are masked when 5-digit ZIP code data is used, since these areas are too large and heterogeneous and thus mask true disparities. On the other hand, neighborhood-level data has been shown to be highly predictive of individuals’ health outcomes.
- **Variables Tested** - a minimum set of SDOH factors to be tested, building from the criteria offered by ASPE or the National Academies Committee report. For the SNP population these would include: dual status, disability status, and factors with significant effect on outcomes, such as: living in a poor neighborhood, single person household size, limited social support, low education level, and limited English proficiency.
- **Accommodation and Sound Methods of Administration** - the measure stewards ensure that the survey methods and administration adequately accommodates low-income, diverse, non-English speaking beneficiaries. Methods should not require beneficiaries to have cell phones, computers or Internet. Surveys should be appropriately translated and interpreter support should be available.
- **Transparency** - the measure developers publish their scientific methods, data sources, and findings to provide a comprehensive technical report available to allow other scientists and analysts in the field to replicate results, as per standard scientific protocol.
- **Dissemination** - the measure developers provide a summary report for disclosure to the general public so that findings are widely available in a form that is accessible.