



MEASURE WORKSHEET

This document summarizes the evaluation of the measure as it progresses through NQF's Consensus Development Process (CDP). The information submitted by measure developers/stewards is included after the Brief Measure Information, Preliminary Analysis, and Pre-meeting Public and Member Comments sections.

To navigate the links in the worksheet: **Ctrl + click link to go to the link; ALT + LEFT ARROW to return**

Brief Measure Information

NQF #: [3211](#)

Measure Title: [Prescription of HIV Antiretroviral Therapy](#)

Measure Steward: [Health Resources and Services Administration - HIV/AIDS Bureau](#)

Brief Description of Measure: [Percentage of patients, regardless of age, with a diagnosis of HIV prescribed antiretroviral therapy for the treatment of HIV infection during the measurement year. A medical visit is any visit in an outpatient/ambulatory care setting with a nurse practitioner, physician, and/or a physician assistant who provides comprehensive HIV care.](#)

Developer Rationale: [Human immunodeficiency virus \(HIV\) is a communicable infection that leads to a progressive disease with a long asymptomatic period. Approximately 50,000 persons in the United States are newly infected with HIV each year. Without treatment, most persons develop acquired immunodeficiency syndrome \(AIDS\) within 10 years of HIV infection. HIV antiretroviral therapy delays this progression and increases the length of survival.](#)

[Current HIV treatment guidelines now recommend universal prescription of HIV antiretroviral therapy for sustained viral load suppression which in turn is directly related to reduction in disease progression and reduction in potential for transmission of HIV infection. Among persons in care, sustained viral load suppression represents the cumulative effect of prescribed therapy, ongoing monitoring, and patient adherence. The proposed measure will direct providers' attention and quality improvement efforts towards this important outcome.](#)

[In 2011, the HIV community saw the emergence of the HIV care continuum. This simple model outlines the sequential steps of medical care that people living with HIV go through from initial diagnosis to achieving the goal of viral suppression. The steps include diagnosis, linkage to care, retention in care, receipt of HIV antiretroviral therapy and viral suppression. This model has been incorporated into the National HIV/AIDS Strategy as it has focused all HIV prevention, care, and treatment efforts in the United States. As outlined in the model, all though there are five different steps, each step is dependent upon each other. For instance, you cannot become virally suppressed if you are not receiving HIV antiretroviral therapy or retained in medical care.](#)

[The most recent nationwide data from CDC dated 2014 estimates that although 86% of people living with HIV have been diagnosed, only 40% are engaged in care, 37% have been prescribed HIV antiretroviral therapy, and 30% have achieved viral suppression.](#)

[Right now, we are at a very special time and place. Many states and large metropolitan areas across the United States have developed plans to end the HIV epidemic in the communities. These jurisdictions have used the HIV care continuum and its steps as the framework by which they have developed their plans.](#)

Numerator Statement: [Number of patients from the denominator prescribed HIV antiretroviral therapy during the measurement year.](#)

Denominator Statement: [Number of patients, regardless of age, with a diagnosis of HIV with at least one medical visit in the measurement year](#)

Denominator Exclusions: [There are no patient exclusions.](#)

Measure Type: [Process](#)

Data Source: [Electronic Health Record \(Only\)](#)

Level of Analysis: [Facility](#)

IF Endorsement Maintenance – Original Endorsement Date: Most Recent Endorsement Date:

New Measure -- Preliminary Analysis

Criteria 1: Importance to Measure and Report

1a. Evidence

Maintenance measures – less emphasis on evidence unless there is new information or change in evidence since the prior evaluation.

1a. Evidence. The evidence requirements for a *process or intermediate outcome* measure is that it is based on a systematic review (SR) and grading of the body of empirical evidence where the specific focus of the evidence matches what is being measured.

The developer provides the following evidence for this measure:

- **Systematic Review of the evidence specific to this measure?** Yes No
- **Quality, Quantity and Consistency of evidence provided?** Yes No
- **Evidence graded?** Yes No

This measure is the new eMeasure version of NQF #2083. The information provided for Evidence and Opportunity for Improvement is identical to that submitted for NQF #2083. Measure #2083 will be discussed first – the ratings for evidence and opportunity for improvement will automatically be assigned to this eMeasure without further discussion.

Evidence Summary

- The developer provided a [diagram](#) outlining the sequential steps of medical care that people living with HIV go through from initial diagnosis to ultimately achieving viral suppression.
- [Evidence](#) and clinical guidelines state that Antiretroviral Therapy is recommended for all HIV-infected individuals in order to reduce morbidity and mortality. Evidence focuses on the percent of providers prescribing ART and the percent of patients with viral load suppression across those providers, the data suggests a positive correlation.
- As a whole, the general evidence suggests that prescription to ART for those infected with HIV will lead to viral suppression if treatment is maintained.
- The developer provided [multiple guidelines](#) for the administration of antiretroviral therapy and viral load monitoring intervals for adults, adolescents and pregnant women.
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Questions for the Committee:

- *What is the relationship of this measure to patient outcomes?*
- *How strong is the evidence for this relationship?*
- *Is the evidence directly applicable to the process of care being measured?*
 - *For possible exception to the evidence criterion:*
- *Are there, or could there be, performance measures of a related health outcome, OR evidence-based intermediate clinical outcomes, intervention/treatment?*

Guidance from the Evidence Algorithm

Process measure evidence based (Box3) → Empirical evidence without grading (BOX 7) → Possible related performance measures (Box 10) → (No exception) → Insufficient

Preliminary rating for evidence: High Moderate Low Insufficient

RATIONALE: Evidence provided does not directly assess the prescription of ART but rather the viral suppression associated with adherence to ART for HIV infected persons.

1b. Gap in Care/Opportunity for Improvement and 1b. Disparities
Maintenance measures – increased emphasis on gap and variation

1b. Performance Gap. The [performance gap](#) requirements include demonstrating quality problems and opportunity for improvement.

[Provider-level performance scores](#) for antiretroviral treatment (ART) for 2014 are presented below.

	2014	2013	2012	2011	2010
Rate	77.6	77.5	74.3	71.1	68.4
Pts w/ ≥1 medical visit (den)	316,087	327,618	335,408	327,744	324,455
Pts w/viral suppression (num)	255,342	249,436	234,505	214,650	200,584
Mean	78.0	77.5	73.4	70.1	65.9
Median	90.0	86.5	83.8	79.8	76.5
Standard Deviation	28.0	24.1	25.4	26.4	27.5
10th percentile	29.6	42.9	31.7	26.1	17.8
90th percentile	98.3	96.4	94.7	93.2	91.2
Min, Max	0.0, 100.0	0.0, 100.0	0.0, 100.0	0.0, 100.0	0.0, 100.0
Pts prescribed ART	245,400 (77.6)	253,972 (77.5)	249,094 (74.3)	233,132 (71.1)	221,908 (68.4)
# of facilities	813	823	816	811	846

Disparities

- The data for measure testing were collected via the Ryan White HIV/AIDS Program Services Report (RSR), which is HRSA HIV/AIDS Bureau's primary source of annual, client-level data collected from more than 2,000 funded grant recipients and subrecipients. Descriptive characteristics are provided by the developer in the table below. The full table can be found [here](#).

	2010		2011		2012		2013		2014	
	N	%	N	%	N	%	N	%	N	%
RACE/ETHNICITY										
American Indian /Alaska Native	1,473	0.5	1,366	0.4	1,371	0.4	1,414	0.5	1,272	0.4
Asian	3,382	1.1	3,598	1.2	3,980	1.2	3,835	1.2	3,791	1.2
Black/African American	146,460	47.3	149,834	47.8	150,974	47.2	146,056	47.0	142,746	46.9
Hispanic/Latino ^a	71,002	22.9	71,240	22.7	75,201	23.5	74,967	24.1	74,714	24.5
Native Hawaiian /Pacific Islander	627	0.2	710	0.2	575	0.2	510	0.2	442	0.2
White	83,854	27.1	83,061	26.5	83,820	26.2	78,953	25.4	75,931	24.9
Multiple races	3,177	1.0	3,716	1.2	4,238	1.3	4,899	1.6	5,651	1.9
GENDER										
Male	219,625	69.7	223,379	69.9	230,075	70.8	221,930	70.7	216,965	70.7
Female	93,266	29.6	93,687	29.3	92,186	28.4	89,212	28.4	87,071	28.4
Transgender	2,313	0.7	2,585	0.8	2,848	0.9	2,779	0.9	2,974	1.0

Questions for the Committee:

- Without data from the eMeasure as specified, do you agree that there is a quality problem with prescription of HIV antiretroviral therapy?
- Are you aware of evidence that other disparities exist in prescribing HIV antiretroviral therapy?

Preliminary rating for opportunity for improvement: High Moderate Low Insufficient

Committee pre-evaluation comments
Criteria 1: Importance to Measure and Report (including 1a, 1b, 1c)

1a. Evidence

*Evidence consistent with the NQF #2083 Measure = INSUFFICIENT

1b. Performance Gap

*Same as the NQF #2083 = MODERATE

Criteria 2: Scientific Acceptability of Measure Properties

2a. Reliability

2a1. Reliability [Specifications](#)

Maintenance measures – no change in emphasis – specifications should be evaluated the same as with new measures

2a1. Specifications requires the measure, as specified, to produce consistent (reliable) and credible (valid) results about the quality of care when implemented.

Data source(s):

- Electronic Health Records (only). This is an eMeasure

Specifications:

- HQMF specifications for the eMeasure are included in the document set on SharePoint. See [eMeasure Technical Advisor](#) review below.
- This measure is specified level of analysis is at hospital/facility/agency level

- Patients are included in the [numerator](#) if they were prescribed HIV antiretroviral therapy during the measurement year
- The [denominator](#) includes the number of patients, regardless of age, with a diagnosis of HIV with at least one medical visit in the measurement year
- There are no patient exclusions
- The [calculation algorithm](#) calculates a rate where a higher score is associated with better performance. The rate is calculated by dividing the numerator population by the denominator population and then multiplying by 100.
- The [value sets](#) needed to calculate the numerator and denominator are included in the specifications.

Questions for the Committee:

- Are all the data elements clearly defined? Are all appropriate codes included?
- Is the logic or calculation algorithm clear?
- Is it likely this measure can be consistently implemented?

eMeasure Technical Advisor(s) review (if not an eMeasure, delete this section):

Submitted measure is an HQMF compliant eMeasure	The submitted eMeasure specifications follow the industry accepted format for eMeasure (HL7 Health Quality Measures Format (HQMF)). HQMF specifications <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Documentation of HQMF or QDM limitations	N/A – All components in the measure logic of the submitted eMeasure are represented using the HQMF and QDM
Value Sets	The submitted eMeasure specifications uses existing value sets when possible and uses new value sets that have been vetted through the VSAC
Measure logic is unambiguous	Submission includes test results from a simulated data set demonstrating the measure logic can be interpreted precisely and unambiguously
Feasibility Testing	The submission contains a feasibility assessment that addresses data element feasibility and follow-up with measure developer indicates that the measure logic is feasible based on assessment by EHR vendors

2a2. Reliability Testing [Testing attachment](#)

Maintenance measures – less emphasis if no new testing data provided

2a2. Reliability testing demonstrates if the measure data elements are repeatable, producing the same results a high proportion of the time when assessed in the same population in the same time period and/or that the measure score is precise enough to distinguish differences in performance across providers.

SUMMARY OF TESTING

Reliability testing level Measure score Data element Both

Reliability testing performed with the data source and level of analysis indicated for this measure Yes No

- The [dataset](#) used for testing included 34 synthetic patients created in the [Bonnie testing system](#) simulating the year 2012. The developer tested the following [data elements](#) using the Bonnie testing tool to evaluate the measure logic:
 - Patient name
 - Date of birth

- Race
- Ethnicity
- Gender
- Payer
- Diagnosis
- Medication Orders
- Encounters
- The patient bundle’s demographics were designed to mimic the HIV/AIDS population, specifically drawing from the patient characteristics collected via the Ryan White HIV/AIDS Program Services Report (RSR).
- Data element validity testing was performed and will count for data element reliability – see validity testing section.
- The developer provided [reliability results from the chart-abstracted measure \(#2083\)](#) and stated, “Currently, there is no performance data available to test the eCQM. However, the chart-abstracted version of this measure has been in use in national quality reporting programs since as early as 2010.”

Questions for the Committee:

- *Is the test sample adequate to generalize for widespread implementation?*
- *Do the results from the Bonnie tool demonstrate sufficient reliability so that differences in performance can be identified?*
- *Do you agree that the reliability test results of the eMeasure will be comparable to the paper based measure (#2083)?*

Guidance from the Reliability Algorithm Precise specifications (Box 1) → Empirical reliability testing (Box 2) → Empirical validity testing of patient-level data (Box 3) → Refer to validity testing of patient-level data elements using Bonnie tool (Box 10 of the Validity algorithm) → Method appropriate for legacy eMeasures (Box 11) → Moderate is the highest possible rating

Preliminary rating for reliability: High Moderate Low Insufficient

2b. Validity

Maintenance measures – less emphasis if no new testing data provided

2b1. Validity: Specifications

2b1. Validity Specifications. This section should determine if the measure specifications are consistent with the evidence.

Specifications consistent with evidence in 1a. Yes Somewhat No

Question for the Committee:

- *Are the specifications consistent with the evidence?*

2b2. Validity testing

2b2. Validity Testing should demonstrate the measure data elements are correct and/or the measure score correctly reflects the quality of care provided, adequately identifying differences in quality.

SUMMARY OF TESTING

Validity testing level Measure score Data element testing against a gold standard Both

Method of validity testing of the measure score:

- Face validity
- Empirical validity testing of the measure score

Validity testing method:

- The [Bonnie testing tool](#), with 34 synthetic patient records were used to test the measure logic and data elements.
 - For each synthetic patients, an expected result was assigned to reflect an expected result of the measure. The synthetic patients were then run against the HQMF output loaded into Bonnie, which “calculates” a measure result for each patients and evaluates it against the expected result.
 - A patient is considered to pass Bonnie testing when the expected result matches the “calculated” result.
- The following testing was completed on the synthetic patients
 - [100% logic coverage](#): The bundle of synthetic patients collectively includes all data elements and conditions that are specified within the measure logic.
 - Clinical relevance. References cited within the chart abstracted measure specification were used to design clinically relevant, realistic patient profiles for the measure’s target population. This approach ensured the eCQM logic maintained alignment with the clinical intent of the chart abstracted measure.
 - [Edge case testing](#): Data elements that test the upper or lower boundary of measure logic conditions.
 - [Negative testing](#): Use of test cases that do not evaluate positively against the measure logic but are otherwise clinically relevant and realistic.
- The developer used references cited within the chart abstracted measure specifications to ensure the eCQM logic maintained alignment with the [clinical intent](#) of the chart abstracted measure.
- In addition to Bonnie testing, the measure specifications were reviewed independently by three eCQM experts to confirm the logic was syntactically correct, using appropriate and current versions of the eCQM standards and terminologies, and consistent with the intent of the chart-abstracted measure.

Validity testing results:

- The testing results from the Bonnie tool reached 100% coverage and confirmed there was a test case for each pathway of logic (negative and positive test cases).
- The measure had a 100% passing rate which confirmed that all the test cases performed as expected.

Questions for the Committee:

- *Is the test sample adequate to generalize for widespread implementation?*
- *Do the results demonstrate sufficient validity so that conclusions about quality can be made?*
- *Do you agree that the score from this measure as specified is an indicator of quality?*

2b3-2b7. Threats to Validity

2b3. Exclusions:

No exclusions

2b4. Risk adjustment: Risk-adjustment method None Statistical model Stratification

2b5. Meaningful difference (*can statistically significant and clinically/practically meaningful differences in performance measure scores can be identified*):

The Data represents variability across providers, In 2014, the bottom 10% of providers had ART prescription rates of 29.6% or lower; the top 90% of providers had rates of 98.3% or higher. These differences demonstrate the continued value of the measure in identifying sites based on poor performance relative to the top performers.

2017	2018	2019	2020	2021	2022	2023	2024	2025	2026	2027
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Question for the Committee:

- o Does the Committee agree the e-Measure will demonstrate similar results to the chart-abstracted measure?

2b6. Comparability of data sources/methods:

- o Not applicable

2b7. Missing Data

- Per the developer, “The HQMF standard specifies that if data are unknown or missing, they shall fail the criterion. This constraint embodies the notion that absence of evidence is evidence of absence, i.e. data not present in a structured field from which the measure draws will not be considered for measure calculation. In certain cases, missing data may have no impact on the measure outcome for a given patient. For example, a data element used in a series of OR statements will not impact the measure outcome if another data element in the OR statement is present and meets all other defined constraints.”
- All Bonnie synthetic patients with missing data performed according to the HQMF standard specification and as expected.

Guidance from the Validity Algorithm Specifications consistent with evidence (Box 1) → Some threats to validity addressed (Box 2) → Empirical validity testing (Box 3) → Face validity testing (Box 4) and empirical testing of data elements using Bonnie tool (Box 10) → Method appropriate for legacy eMeasures (Box 11) → Moderate is the highest possible rating

Preliminary rating for validity: High Moderate Low Insufficient

Committee pre-evaluation comments
Criteria 2: Scientific Acceptability of Measure Properties (including all 2a, 2b, and 2d)

2a1. Reliability Specifications

*All data elements are well defined. All codes are provided, while no descriptors are available other than the field definition All steps are clear. No concerns that this can be consistently implemented. - Due to Field testing and no Score testing, Reliability = MODERATE

*The data elements are clearly defined
 All appropriate codes included
 The logic or calculation algorithm is clear
 It is likely this measure can be consistently implemented"

2a2. Reliability Testing

*Reliability testing good. Measure is a legacy eCQM. Elements are "yes"/"no" responses based on data in the medical record and provide good reliability. Not a PRO-PM measure.

*The reliability test results of the eMeasure should be comparable to the paper based measure (#2083)

2b1. Validity Specifications

*None. Not a PRO-PM measure. No empirical validity testing. Validity = MODERATE

2b2. Validity Testing

*Adequate testing. Bonnie Testing Tool used which is a gold standard for data element testing. No empirical validity testing. Validity = MODERATE.

*The test sample is adequate to generalize for widespread implementation

The results demonstrate sufficient validity so that conclusions about quality can be made
It is not clear that medications prescribed are received by the patient."

2b3-7 Threats to Validity

*Missing data/responses are not considered in calculation of the rates. "absence of evidence is evidence of absent".

*2b.5 The e-Measure will demonstrate similar results to the chart-abstracted measure

"

Criterion 3. Feasibility

Maintenance measures – no change in emphasis – implementation issues may be more prominent

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

- The developer provided information on feasibility testing in the [eMeasure Feasibility Score Card](#). The developer did not identify the EHRs used for feasibility testing. Instead, the developer stated that the feasibility assessment was “conducted by consensus of a panel of MITRE clinical informatics, measure development, and eCQM standards experts”.
- The developer provided a summary of the latest publicly available data on Meaningful Use EHR capabilities and provider performance on objectives and measures related to the eCQM’s data elements:
 - CPOE – Meds
 - CPOE – Labs
 - Demographics
 - Problem List
 - Lab test results
- On a scale from 1 to 3 where 3 is the highest score, all but 3 of the data elements received a score of ‘3’.
 - Both ‘Encounter, Performed: Face to Face Interaction’ and ‘Patient Characteristic Payer’ scored a 2 on Data Standards.
 - The Score 2 definition for Data Standards is “terminology standards for this data element are currently available, but it is not consistently coded to standard terminology in the EHR, or the EHR does not easily allow such coding.”
 - The data element ‘Patient Characteristic Expired’ scored a 2 on Data Accuracy. Data accuracy looks at the correctness of the information contained in the data element and whether the data source and recorder are specified.
 - The Score 2 definition for Data Accuracy is “the information may not be from the most authoritative source and/or has a moderate likelihood of being correct”. The scorecard notes that this information is similar to “self-reporting of a vaccination”.
 - The developer notes that “The accuracy of this data element is dependent on full end-to-end interoperability across providers and between providers and public health agencies.”
- The developer indicates that on a scale from 0 to 100 percent, the measure is currently **98.33%** feasible and in one to two years, will be **98.89%** feasible.

Questions for the Committee:

- Are the required data elements routinely generated and used during care delivery?
- Is the data collection strategy ready to be put into operational use?
- Does the eMeasure Feasibility Score Card demonstrate acceptable feasibility in multiple EHR systems and sites?

Preliminary rating for feasibility: High Moderate Low Insufficient

RATIONALE:

Committee pre-evaluation comments

Criteria 3: Feasibility

3. Feasibility

*The required data elements are routinely generated/used in clinical care delivery. The data collection strategy is already in place. No concerns.

*The required data elements are routinely generated and used during care delivery

The data collection strategy is ready to be put into operational use
I was unable to access the eMeasure Feasibility Score Card "

Criterion 4: Usability and Use

Maintenance measures – increased emphasis – much greater focus on measure use and usefulness, including both impact /improvement and unintended consequences

4. Usability and Use evaluate the extent to which audiences (e.g., consumers, purchasers, providers, policymakers) use or could use performance results for both accountability and performance improvement activities.

Current uses of the measure

Publicly reported? Yes No

Current use in an accountability program? Yes No UNCLEAR

OR

Planned use in an accountability program? Yes No

Accountability program details

- Ryan White HIV/AIDS Program
 - Sponsor: Federal government
 - Geographic area: Nationwide
 - Accountable entities: Approximately 600 Ryan White HIV/AIDS Program grant recipients and their providers
 - Patients: Approximately 316,000 patients
- Physician Quality Report System (PQRS) and Value Based Modifier
 - Sponsor: Federal government
 - Geographic area: Nationwide
 - Accountable entities: Physicians and practitioners
 - Patients: Unknown
- National HIV/AIDS Strategy
 - Sponsor: Federal government
 - Geographic area: Nationwide
 - Accountable entities: Federal agencies and service providers
 - Patients: All people living with HIV in the United States

Improvement results

- The developer reports that the Ryan White HIV/AIDS Program has experienced a 10 + point increase in viral suppression from 65.9% in 2010 to 78.0% in 2014. Prescription of HIV antiretroviral therapy has increased across all demographic groups and subpopulations.

Unexpected findings (positive or negative) during implementation

- This measure has been adopted by Centers for Medicare and Medicaid measurement programs, Department of Health and Human Service Secretary as a one of the core HIV indicators, countless outpatient/ambulatory care settings, and health departments. National learning collaborates have used this measure to focus the improvement efforts of grant recipients and subrecipients. Additionally, prescription of HIV antiretroviral therapy is one of five stages of the HIV care continuum. This measure has become the standard when measuring prescription of HIV antiretroviral therapy.

Potential harms

- Not applicable

Vetting of the measure

- The developer reports that Health Resources and Services Administration worked diligently to release the annual data report in the same year it was collected which is publically available on the Health Resources and Services Administration website (<http://hab.hrsa.gov/data/data-reports>).
- Ryan White HIV/AIDS Program national partners has provided feedback regarding the timeliness, feasibility, and usability of the release of the Ryan White HIV/AIDS Program Annual Client-Level Data Report, supplemental reports, slide decks, fact sheets, and infographics. The national partners encourage the continued release of the data in all its formats.
- During the initial development of the chart-abstracted measure, formal feedback was gathered. The measures were modified during the development phase and have not been modified since. On an annual basis, the measures are reviewed for clinical relevance, change in scientific acceptability, and consistency with guidelines. The chart-abstracted measure has not been modified as a result of the annual reviews.

Feedback:

- The developer reports that feedback has been received from Ryan White HIV/AIDS Program grant recipients and subrecipients regarding the feasibility and usefulness of the data presented in the Ryan White HIV/AIDS Program Annual Client-Level Data Report. Significant feedback has been provided about the timeliness and expansions of the data release. Grant recipient report using the data for benchmarking their program, setting goals/targets, and gaining a fuller understanding of all aspects of the Ryan White HIV/AIDS Program. Grant recipients and subrecipients have also requested additional analyses.

Questions for the Committee:

- *How can the performance results be used to further the goal of high-quality, efficient healthcare?*
- *Do the benefits of the measure outweigh any potential unintended consequences?*
- *How has the eCQM been vetted in real-world settings by those being measured or others?*

Preliminary rating for usability and use: High Moderate Low Insufficient

Committee pre-evaluation comments
Criteria 4: Usability and Use

4. Usability and Use

*Data are currently reported publically and used to identify providers who need improvement in this area. It is also being used in a pay-4-performance programs. The results are also used as benchmarks for quality improvement work being done in clinics both inside and outside of the Ryan White HIV/AIDS Program.

Criterion 5: [Related and Competing Measures](#)

Related or competing measures

- 0405 HIV/AIDS: Pneumocystis Jiroveci Pneumonia (PCP) Prophylaxis
- 0409 HIV/AIDS: Sexually Transmitted Disease Screening for Chlamydia, Gonorrhea, and Syphilis
- 2079 HIV Medical Visit Frequency
- 2080 Gap in HIV Medical Visits
- 2082 HIV Viral Suppression
- 3210 HIV Viral Suppression
- 3010 HIV Medical Visit Frequency

Harmonization

Measure is fully harmonized to the extent possible according to the developer

Endorsement + Designation

The “Endorsement +” designation identifies measures that exceed NQF's endorsement criteria in several key areas. After a Committee recommends a measure for endorsement, it will then consider whether the measure also meets the “Endorsement +” criteria.

This measure is a candidate for the “Endorsement +” designation IF the Committee determines that it: meets evidence for measure focus without an exception; is reliable, as demonstrated by score-level testing; is valid, as demonstrated by score-level testing (not via face validity only); and has been vetted by those being measured or other users.

Eligible for Endorsement + designation: Yes No

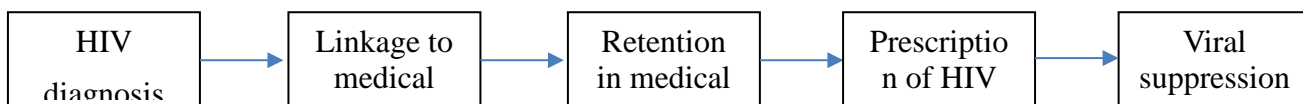
RATIONALE IF NOT ELIGIBLE: The measure is not eligible for Endorsement+ because empirical reliability and validity testing of the measure score was not conducted and the measure has not been vetted in real world settings by those being measured and other users.

Pre-meeting public and member comments

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Measure Title: Prescription of HIV Antiretroviral Therapy

1a.12 LOGIC MODEL



Although the above diagram outlines the sequential steps of medical care that people living with HIV go through from initial diagnosis to achieving the goal of viral suppression (also known as the HIV care continuum). For some patients, this is a linear path with sustained viral suppression for many years. For other patients, there may be years between diagnosis and linkage. Yet still for others, retention in medical care is not consistent, which results in missed visits, no prescription for or adherence to HIV antiretroviral therapy (ART), and lack of viral suppression.

1a.2 FOR OUTCOME MEASURES including PATIENT REPORTED OUTCOMES State the rationale supporting the relationship between the health outcome (or PRO) to at least one healthcare structure, process (e.g., intervention, or service).

Regularly attending medical visits (retention) is paramount to monitoring patient’s health status, screenings, and laboratory values. Providers need this information to make an informed decision in order to prescribe HIV antiretroviral therapy (ART). ART reduces HIV-associated morbidity and mortality by maximally inhibiting HIV replication (as defined by achieving and maintaining plasma HIV RNA (viral load) below levels detectable by commercially available assays). Durable viral suppression improves immune function and quality of life,

lowers the risk of both AIDS-defining and non-AIDS-defining complications, and prolongs life. Emerging evidence also suggests that additional benefits of ART-induced viral load suppression include a reduction in HIV-associated inflammation and possibly its associated complications.

1b.1. Briefly explain the rationale for this measure (e.g., how the measure will improve the quality of care, the benefits or improvements in quality envisioned by use of this measure)

Human immunodeficiency virus (HIV) is a communicable infection that leads to a progressive disease with a long asymptomatic period. Approximately 50,000 persons in the United States are newly infected with HIV each year. Without treatment, most persons develop acquired immunodeficiency syndrome (AIDS) within 10 years of HIV infection. HIV antiretroviral therapy delays this progression and increases the length of survival.

Current HIV treatment guidelines now recommend universal prescription of HIV antiretroviral therapy for sustained viral load suppression which in turn is directly related to reduction in disease progression and reduction in potential for transmission of HIV infection. Among persons in care, sustained viral load suppression represents the cumulative effect of prescribed therapy, ongoing monitoring, and patient adherence. The proposed measure will direct providers' attention and quality improvement efforts towards this important outcome.

In 2011, the HIV community saw the emergence of the HIV care continuum. This simple model outlines the sequential steps of medical care that people living with HIV go through from initial diagnosis to achieving the goal of viral suppression. The steps include diagnosis, linkage to care, retention in care, receipt of HIV antiretroviral therapy and viral suppression. This model has been incorporated into the National HIV/AIDS Strategy as it has focused all HIV prevention, care, and treatment efforts in the United States. As outlined in the model, all though there are five different steps, each step is dependent upon each other. For instance, you cannot become virally suppressed if you are not receiving HIV antiretroviral therapy or retained in medical care.

The most recent nationwide data from CDC dated 2014 estimates that although 86% of people living with HIV have been diagnosed, only 40% are engaged in care, 37% have been prescribed HIV antiretroviral therapy, and 30% have achieved viral suppression.

Right now, we are at a very special time and place. Many states and large metropolitan areas across the United States have developed plans to end the HIV epidemic in the communities. These jurisdictions have used the HIV care continuum and its steps as the framework by which they have developed their plans.

1a.4. CLINICAL PRACTICE GUIDELINE RECOMMENDATION

1a.4.1. Guideline citation (including date) and URL for guideline (if available online):

Panel on Antiretroviral Guidelines for Adults and Adolescents. Guidelines for the use of antiretroviral agents in HIV-1-infected adults and adolescents. Department of Health and Human Services Accessed November 15, 2016: <http://www.aidsinfo.nih.gov/ContentFiles/AdultandAdolescentGL.pdf>

World Health Organization (WHO). (2016). Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection: recommendations for a public health approach. Accessed November 15, 2016: http://apps.who.int/iris/bitstream/10665/208825/1/9789241549684_eng.pdf?ua=1

International Advisory Panel on HIV Care Continuum Optimization (IAPAC). (2015). IAPAC Guidelines for Optimizing the HIV Care Continuum for Adults and Adolescents. Accessed November 15, 2016. <http://www.iapac.org/uploads/IIAPAC-IAPAC-Guidelines-for-Optimizing-the-HIV-Care-Continuum-Supplement-Nov-Dec-2015.pdf>

Günthard HF, Saag MS, Benson CA, del Rio C, Eron JJ, Gallant JE, Hoy JF, Mugavero MJ, Sax PE, Thompson MA, Gandhi RT, Landovitz RJ, Smith DM, Jacobsen DM, Volberding PA. Antiretroviral Drugs for Treatment and Prevention of HIV Infection in Adults 2016 Recommendations of the International Antiviral Society–USA Panel. JAMA. 2016. <https://www.iasusa.org/content/antiretroviral-drugs-treatment-and-prevention-hiv-infection-adults-2016-recommendations>

1a.4.2. Identify guideline recommendation number and/or page number and quote verbatim, the specific guideline recommendation.

Guidelines for the Use of Antiretroviral Agents in HIV-1-Infected Adults and Adolescents:

Initiation of Antiretroviral Therapy (page E-1)

- Antiretroviral therapy (ART) is recommended for all HIV-infected individuals, regardless of CD4 T lymphocyte cell count, to reduce the morbidity and mortality associated with HIV infection (AI).
- ART is also recommended for HIV-infected individuals to prevent HIV transmission (AI).
- When initiating ART, it is important to educate patients regarding the benefits and considerations regarding ART, and to address strategies to optimize adherence. On a case-by-case basis, ART may be deferred because of clinical and/or psychosocial factors, but therapy should be initiated as soon as possible.

Considerations for Antiretroviral Use in Special Patient Populations: Acute and Recent (Early) HIV Infection (page I-1)

- Antiretroviral therapy (ART) is recommended for all individuals with HIV-1 infection (AI) including those with early HIV-1 infection.

HIV-Infected Adolescents and Young Adults (page I-8):

- ART is recommended for all HIV-infected individuals (AI) to reduce morbidity and mortality. Thus, ART is also recommended for ART-naïve adolescents. However, before initiation of therapy, adolescents' readiness and ability to adhere to therapy within their psychosocial context need to be carefully considered as partner of therapeutic decision making (AIII).

HIV-Infected Women (page I-20):

- Antiretroviral therapy (ART) is recommended for all HIV-infected women to improve their health and to reduce the risk of HIV transmission to HIV-uninfected sex partners (AI).

HIV/Hepatitis C Virus Coinfection (page J-6):

- Antiretroviral therapy (ART) may slow the progression of liver disease by preserving or restoring immune function and reducing HIV related immune activation and inflammation. For most HCV/HIV-coinfected patients, including those with cirrhosis, the benefits of ART outweigh concerns regarding drug-induced liver injury. Therefore, ART should be initiated in all HCV/HIV-coinfected patients, regardless of CD4 T lymphocyte (CD4) cell count (AI).

WHO:

4.3 When to start ART (page xxxi)

4.3.1 When to start ART in adults (>19 years old)

- ART should be initiated in all adults living with HIV, regardless of WHO clinical stage and at any CD4 cell count (strong recommendation, moderate-quality evidence).
- As a priority, ART should be initiated in all adults with severe or advanced HIV clinical disease (WHO clinical stage 3 or 4) and adults with a CD4 count ≤ 350 cells/mm³ (strong recommendation, moderate-quality evidence).

4.3.2 When to start ART in pregnant and breastfeeding women

- ART should be initiated in all pregnant and breastfeeding women living with HIV, regardless of WHO clinical stage and at any CD4 cell count and continued lifelong (strong recommendation, moderate-quality evidence).

4.3.3 When to start HIV antiretroviral therapy in adolescents (10–19 years of age)

- ART should be initiated in all adolescents living with HIV, regardless of WHO clinical stage and at any CD4 cell count (conditional recommendation, low-quality evidence).
- As a priority, ART should be initiated in all adolescents with severe or advanced HIV clinical disease (WHO clinical stage 3 or 4) and adolescents with a CD4 count ≤ 350 cells/mm³ (strong recommendation, moderate-quality evidence).

4.3.4 When to start HIV antiretroviral therapy in children younger than 10 years of age

- ART should be initiated in all children living with HIV, regardless of WHO clinical stage or at any CD4 cell count:
- Infants diagnosed in the first year of life (strong recommendation, moderate-quality evidence).
- Children living with HIV 1-year-old to less than 10 years old (conditional recommendation, low-quality evidence).
- As a priority, ART should be initiated in all children <2 years of age or children younger than 5 years of age with WHO clinical stage 3 or 4 or CD4 count ≤ 750 cells/mm³ or CD4 percentage <25% and children 5 years of age and older with WHO clinical stage 3 or 4 or CD4 count ≤ 350 cells/mm³ (strong recommendation, moderate-quality evidence).

4.3.5 Timing of HIV ANTIRETROVIRAL THERAPY for adults and children with TB

- ART should be started in all TB patients living with HIV regardless of CD4 count (strong recommendation, high-quality evidence).

International Advisory Panel on HIV Care Continuum Optimization (IAPAC):

Increasing HIV treatment coverage (page 3)

- The immediate offer of ART after HIV diagnosis, irrespective of CD4 count or clinical stage, is recommended. (AI)

Antiretroviral Drugs for Treatment and Prevention of HIV Infection in Adults 2016 Recommendations of the International Antiviral Society–USA Panel

Box 1. Recommendations for When to Start (page 193)

- Antiretroviral therapy (HIV ANTIRETROVIRAL THERAPY) is recommended for all viremic patients with established HIV infection, regardless of CD4 cell count (evidence rating AIa).
- Initiation of ART is recommended as soon as possible in the setting of acute HIV infection (evidence rating BIII).
- Planned discontinuation of early ART after a specific duration of treatment is not recommended outside a research setting (evidence rating AIa).
- Initiation of ART is recommended for individuals who have persistent undetectable viral load without ART but have declining CD4 cell counts (evidence rating BIII).

1a.4.3. Grade assigned to the quoted recommendation with definition of the grade:

Guidelines for the Use of Antiretroviral Agents in HIV-1-Infected Adults and Adolescents:

Basis for Recommendations

Recommendations in these guidelines are based upon scientific evidence and expert opinion. Each recommended statement includes a letter (A, B, or C) that represents the strength of the recommendation and a Roman numeral (I, II, or III) that represents the quality of the evidence that supports the recommendation (see Table 2).

Table 2. Rating Scheme for Recommendations

Strength of Recommendation	Quality of Evidence for Recommendation
A: Strong recommendation for the statement	I: One or more randomized trials with clinical outcomes and/or validated laboratory endpoints II: One or more well-designed, non-randomized trials or observational cohort studies with long-term clinical outcomes III: Expert opinion
B: Moderate recommendation for the statement	
C: Optional recommendation for the statement	

International Advisory Panel on HIV Care Continuum Optimization; IAPAC Guidelines for Optimizing the HIV Care Continuum for Adults and Adolescents.

Strong (A) = Almost all patients should receive the recommended course of action.

Moderate (B) = Most patients should receive the recommended course of action. However, other choices may be appropriate for some patients.

Optional (C) There may be consideration for this recommendation based on individual patient circumstances. Not recommended routinely.

Quality of the Body of Evidence and its Interpretation:

Excellent (I) = Randomized control trial (RCT) evidence without important limitations; overwhelming evidence from observational studies

High (II) = RCT evidence with important limitations; strong evidence from observational studies

Medium (III) = RCT evidence with critical limitations; observational study without important limitations

Low (IV) = Other evidence, including extrapolations from bench research, usual practice, expert opinion, consensus guidelines; observational study evidence with important or critical limitations

World Health Organization. Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection Recommendations for a public health approach - Second edition.:

The strength of a recommendation can be either strong or conditional. Process of guideline development This edition of the guidelines was revised in accordance with procedures established by the WHO Guidelines Review Committee. New clinical and operational recommendations in the guidelines are based on the GRADE (Grading of Recommendations, Assessment, Development and Evaluation) approach to reviewing evidence. Modelling, expert consultations and country case studies have all strongly informed the guidelines. The process has also identified key gaps in knowledge that will help to guide the future HIV research agenda. A strong recommendation is one for which there is confidence that the desirable effects of adherence to the recommendation clearly outweigh the undesirable effects.

A conditional recommendation is one for which the Guideline Development Group concludes that the desirable effects of adherence to the recommendation probably outweigh the undesirable effects or are closely balanced, but the Groups are not confident about these trade-offs in all situations. At implementation, monitoring and rigorous evaluation is needed to address these uncertainties, which are likely to provide new evidence that may change the calculation of the balance of trade-offs and to suggest how to overcome any implementation challenges.

Quality of evidence Definition

Table 1.1. GRADE quality of evidence

Quality of evidence	Definition
High	We are very confident that the true effect lies close to that of the estimate of the effect

Middle	We are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of effect, but there is a possibility that it is substantially different
Low	Our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect
Very low	We have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of the effect

Antiretroviral Drugs for Treatment and Prevention of HIV Infection in Adults, 2016 Recommendations of the International Antiviral Society–USA Panel:

Table 1. Strength of Recommendation and Quality of Evidence Rating Scale

Rating	Definition
Strength of recommendation	
A	Strong support for the recommendation
B	Moderate support for the recommendation
C	Limited support for the recommendation
Quality of evidence	
Ia	Evidence for ≥ 1 randomized clinical trials published in the peer-reviewed literature
Ib	Evidence for ≥ 1 randomized clinical trials presented in abstract form at peer-reviewed scientific meetings
IIa	Evidence from nonrandomized clinical trials or cohorts or case-control studies published in the peer-reviewed literature
IIb	Evidence from nonrandomized clinical trials or cohorts or case-control studies published in the peer-reviewed scientific meeting
III	Recommendation based on panel’s analysis of the accumulated available evidence

1a.4.4. Provide all other grades and associated definitions for recommendations in the grading system.

(Note: If separate grades for the strength of the evidence, report them in section 1a.7.)

All grade and definitions noted in 1a.4.3.

1a.4.5. Citation and URL for methodology for grading recommendations (if different from 1a.4.1):

Citations noted in 1a.4.1.

1a.4.6. If guideline is evidence-based (rather than expert opinion), are the details of the quantity, quality, and consistency of the body of evidence available (e.g., evidence tables)?

Yes → *complete section 1a.7*

No → *report on another systematic review of the evidence in sections 1a.6 and 1a.7; if another review does not exist, provide what is known from the guideline review of evidence in 1a.7*

1. Evidence, Performance Gap, Priority – Importance to Measure and Report

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

1a. Evidence to Support the Measure Focus – See attached Evidence Submission Form

[ART_evidence_NQF-636174955634964398-636177547774061167.docx](#)

1a.1 For Maintenance of Endorsement: Is there new evidence about the measure since the last update/submission?

Please update any changes in the evidence attachment in red. Do not remove any existing information. If there have been any changes to evidence, the Committee will consider the new evidence. If there is no new evidence, no updating of the evidence information is needed.

Yes

1b. Performance Gap

Demonstration of quality problems and opportunity for improvement, i.e., data demonstrating:

- considerable variation, or overall less-than-optimal performance, in the quality of care across providers; and/or
- Disparities in care across population groups.

1b.1. Briefly explain the rationale for this measure (e.g., how the measure will improve the quality of care, the benefits or improvements in quality envisioned by use of this measure)

IF a PRO-PM (e.g. HRQoL/functional status, symptom/burden, experience with care, health-related behaviors), provide evidence that the target population values the measured PRO and finds it meaningful. (Describe how and from whom their input was obtained.)

IF a COMPOSITE (e.g., combination of component measure scores, all-or-none, any-or-none), SKIP this question and provide rationale for composite in question 1c.3 on the composite tab.

Human immunodeficiency virus (HIV) is a communicable infection that leads to a progressive disease with a long asymptomatic period. Approximately 50,000 persons in the United States are newly infected with HIV each year. Without treatment, most persons develop acquired immunodeficiency syndrome (AIDS) within 10 years of HIV infection. HIV antiretroviral therapy delays this progression and increases the length of survival.

Current HIV treatment guidelines now recommend universal prescription of HIV antiretroviral therapy for sustained viral load suppression which in turn is directly related to reduction in disease progression and reduction in potential for transmission of HIV infection. Among persons in care, sustained viral load suppression represents the cumulative effect of prescribed therapy, ongoing monitoring, and patient adherence. The proposed measure will direct providers' attention and quality improvement efforts towards this important outcome.

In 2011, the HIV community saw the emergence of the HIV care continuum. This simple model outlines the sequential steps of medical care that people living with HIV go through from initial diagnosis to achieving the goal of viral suppression. The steps include diagnosis, linkage to care, retention in care, receipt of HIV antiretroviral therapy and viral suppression. This model has been incorporated into the National HIV/AIDS Strategy as it has focused all HIV prevention, care, and treatment efforts in the United States. As outlined in the model, all though there are five different steps, each step is dependent upon each other. For instance, you cannot become virally suppressed if you are not receiving HIV antiretroviral therapy or retained in medical care.

The most recent nationwide data from CDC dated 2014 estimates that although 86% of people living with HIV have been diagnosed, only 40% are engaged in care, 37% have been prescribed HIV antiretroviral therapy, and 30% have achieved viral suppression.

Right now, we are at a very special time and place. Many states and large metropolitan areas across the United States have developed plans to end the HIV epidemic in the communities. These jurisdictions have used the HIV care continuum and its steps as the framework by which they have developed their plans.

1b.2. Provide performance scores on the measure as specified (current and over time) at the specified level of analysis. *(This is required for maintenance of endorsement. Include mean, std dev, min, max, interquartile range, scores by decile. Describe the data source including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities include.) This information also will be used to address the sub-criterion on improvement (4b) under Usability and Use.*
Please see attachment "ART submission form" for formatted data.

1b.3. If no or limited performance data on the measure as specified is reported in 1b2, then provide a summary of data from the literature that indicates opportunity for improvement or overall less than optimal performance on the specific focus of measurement.

N/A

1b.4. Provide disparities data from the measure as specified (current and over time) by population group, e.g., by race/ethnicity, gender, age, insurance status, socioeconomic status, and/or disability. *(This is required for maintenance of endorsement. Describe the data source including number of measured entities; number of patients; dates of data; if a sample, characteristics of the entities included.) For measures that show high levels of performance, i.e., "topped out", disparities data may demonstrate an opportunity for improvement/gap in care for certain sub-populations. This information also will be used to address the sub-criterion on improvement (4b) under Usability and Use.*
Please see attachment "ART submission form" for formatted data.

1b.5. If no or limited data on disparities from the measure as specified is reported in 1b.4, then provide a summary of data from the literature that addresses disparities in care on the specific focus of measurement. Include citations. Not necessary if performance data provided in 1b.4

N/A

2. Reliability and Validity—Scientific Acceptability of Measure Properties

Extent to which the measure, as specified, produces consistent (reliable) and credible (valid) results about the quality of care when implemented. **Measures must be judged to meet the sub criteria for both reliability and validity to pass this criterion and be evaluated against the remaining criteria.**

2a.1. Specifications The measure is well defined and precisely specified so it can be implemented consistently within and across organizations and allows for comparability. eMeasures should be specified in the Health Quality Measures Format (HQMF) and the Quality Data Model (QDM).

De.5. Subject/Topic Area *(check all the areas that apply):*
Infectious Diseases (ID) : HIV/AIDS

De.6. Cross Cutting Areas *(check all the areas that apply):*
«crosscutting_area»

De.7. Target Population Category *(Check all the populations for which the measure is specified and tested if any):*
Populations at Risk

S.1. Measure-specific Web Page *(Provide a URL link to a web page specific for this measure that contains current detailed specifications including code lists, risk model details, and supplemental materials. Do not enter a URL linking to a home page or to general information.)*

There is no measure-specific web page for the electronic version of this measure.

S.2a. If this is an eMeasure, HQMF specifications must be attached. Attach the zipped output from the eMeasure authoring tool (MAT) - if the MAT was not used, contact staff. (Use the specification fields in this online form for the plain-language description of the specifications)

This is an eMeasure **Attachment:** [NQFXXX_PrescriptionOfAntiretroviralTherapy_Artifacts-636177547766417118.zip](#), [NQFXXX_PrescriptionOfAntiretroviralTherapy_MeasureSubmissionForm-636177547766573119.docx](#)

S.2b. Data Dictionary, Code Table, or Value Sets (and risk model codes and coefficients when applicable) must be attached. (Excel or csv file in the suggested format preferred - if not, contact staff)

Attachment Attachment: [HIVPAT_v4_6_Thu_Dec_15_20.34.08_CST_2016.xls](#)

S.3.1. For maintenance of endorsement: Are there changes to the specifications since the last updates/submission. If yes, update the specifications for S1-2 and S4-22 and explain reasons for the changes in S3.2.

No

S.3.2. For maintenance of endorsement, please briefly describe any important changes to the measure specifications since last measure update and explain the reasons.

S.4. Numerator Statement (Brief, narrative description of the measure focus or what is being measured about the target population, i.e., cases from the target population with the target process, condition, event, or outcome) DO NOT include the rationale for the measure.

IF an OUTCOME MEASURE, state the outcome being measured. Calculation of the risk-adjusted outcome should be described in the calculation algorithm (S.14).

Number of patients from the denominator prescribed HIV antiretroviral therapy during the measurement year.

S.5. Numerator Details (All information required to identify and calculate the cases from the target population with the target process, condition, event, or outcome such as definitions, time period for data collection, specific data collection items/responses, code/value sets – Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at S.2b)

IF an OUTCOME MEASURE, describe how the observed outcome is identified/counted. Calculation of the risk-adjusted outcome should be described in the calculation algorithm (S.14).

The antiretroviral therapy medication order is represented by the QDM element "Medication, Order: FDA Approved HIV Antiretroviral Therapy" using "HIV Antiretroviral Therapy RXNORM Value Set (2.16.840.1.113762.1.4.1032.1)." In order to be included in the numerator, the "Medication, Order: FDA Approved HIV Antiretroviral Therapy" element must start during the measurement period.

S.6. Denominator Statement (Brief, narrative description of the target population being measured)

Number of patients, regardless of age, with a diagnosis of HIV with at least one medical visit in the measurement year

S.7. Denominator Details (All information required to identify and calculate the target population/denominator such as definitions, time period for data collection, specific data collection items/responses, code/value sets – Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at S.2b.)

IF an OUTCOME MEASURE, describe how the target population is identified. Calculation of the risk-adjusted outcome should be described in the calculation algorithm (S.14).

The patient's HIV diagnosis is represented by the QDM element "Diagnosis: HIV" using "HIV Grouping Value Set (2.16.840.1.113883.3.464.1003.120.12.1003)".

The patient's medical visits are represented by the following QDM elements:

- "Diagnosis: HIV 1" using "HIV 1 Grouping Value Set (2.16.840.1.113883.3.464.1003.120.12.1004)"
- "Encounter, Performed: Face-to-Face Interaction" using "Face-to-Face Interaction Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1048)"
- "Encounter, Performed: Office Visit" using "Office Visit Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1001)"

- "Encounter, Performed: Outpatient Consultation" using "Outpatient Consultation Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1008)"
- "Encounter, Performed: Preventive Care - Established Office Visit, 0 to 17" using "Preventive Care - Established Office Visit, 0 to 17 Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1024)"
- "Encounter, Performed: Preventive Care Services - Established Office Visit, 18 and Up" using "Preventive Care Services - Established Office Visit, 18 and Up Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1025)"
- "Encounter, Performed: Preventive Care Services-Initial Office Visit, 18 and Up" using "Preventive Care Services-Initial Office Visit, 18 and Up Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1023)"
- "Encounter, Performed: Preventive Care- Initial Office Visit, 0 to 17" using "Preventive Care- Initial Office Visit, 0 to 17 Grouping Value Set (2.16.840.1.113883.3.464.1003.101.12.1022)"

S.8. Denominator Exclusions (Brief narrative description of exclusions from the target population)

There are no patient exclusions.

S.9. Denominator Exclusion Details (All information required to identify and calculate exclusions from the denominator such as definitions, time period for data collection, specific data collection items/responses, code/value sets – Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format at S.2b.)

There are no patient exclusions.

S.10. Stratification Information (Provide all information required to stratify the measure results, if necessary, including the stratification variables, definitions, specific data collection items/responses, code/value sets, and the risk-model covariates and coefficients for the clinically-adjusted version of the measure when appropriate – Note: lists of individual codes with descriptors that exceed 1 page should be provided in an Excel or csv file in required format with at S.2b.)

N/A

S.11. Risk Adjustment Type (Select type. Provide specifications for risk stratification in measure testing attachment)

No risk adjustment or risk stratification

If other:

S.12. Type of score:

Rate/proportion

If other:

S.13. Interpretation of Score (Classifies interpretation of score according to whether better quality is associated with a higher score, a lower score, a score falling within a defined interval, or a passing score)

Better quality = Higher score

S.14. Calculation Algorithm/Measure Logic (Diagram or describe the calculation of the measure score as an ordered sequence of steps including identifying the target population; exclusions; cases meeting the target process, condition, event, or outcome; time period for data, aggregating data; risk adjustment; etc.)

1. Identify the individuals who satisfy all specific criteria for inclusion in the denominator: 1.) diagnosed with HIV during the first 3 months of the measurement year or prior to the measurement year; and 2.) had at least one medical visit during the measurement year. The individuals who met these criteria are the denominator population.
2. Identify the individuals from the denominator population who meet the criterion for inclusion in the numerator: prescribed HIV antiretroviral therapy during the measurement year.
3. Calculate the percentage by dividing the numerator population by the denominator population and multiply by 100.

S.15. Sampling (If measure is based on a sample, provide instructions for obtaining the sample and guidance on minimum sample size.)

IF a PRO-PM, identify whether (and how) proxy responses are allowed.

Not applicable; not based on a sample.

S.16. Survey/Patient-reported data (If measure is based on a survey or instrument, provide instructions for data collection and guidance on minimum response rate.)

IF a PRO-PM, specify calculation of response rates to be reported with performance measure results.

This measure is not based on a survey or instrument.

S.17. Data Source (Check ONLY the sources for which the measure is SPECIFIED AND TESTED).

If other, please describe in S.18.

Electronic Health Record (Only)

S.18. Data Source or Collection Instrument (Identify the specific data source/data collection instrument (e.g. name of database, clinical registry, collection instrument, etc., and describe how data is collected.)

IF a PRO-PM, identify the specific PROM(s); and standard methods, modes, and languages of administration.

Not applicable.

S.19. Data Source or Collection Instrument (available at measure-specific Web page URL identified in S.1 OR in attached appendix at A.1)

No data collection instrument provided

S.20. Level of Analysis (Check ONLY the levels of analysis for which the measure is SPECIFIED AND TESTED)

Facility

S.21. Care Setting (Check ONLY the settings for which the measure is SPECIFIED AND TESTED)

Clinician Office/Clinic

If other:

S.22. COMPOSITE Performance Measure - Additional Specifications (Use this section as needed for aggregation and weighting rules, or calculation of individual performance measures if not individually endorsed.)

This is not a composite measure.

2. Validity – See attached Measure Testing Submission Form

[ART_testing-636177547781081212.docx](#), [NQFXXX_PrescriptionOfAntiretroviralTherapy_BonnieTestingAttachment-636177547781237213.zip](#)

2.1 For maintenance of endorsement

Reliability testing: If testing of reliability of the measure score was not presented in prior submission(s), has reliability testing of the measure score been conducted? If yes, please provide results in the Testing attachment. (Do not remove prior testing information – include date of new information in red.)

Yes

2.2 For maintenance of endorsement

Has additional empirical validity testing of the measure score been conducted? If yes, please provide results in the Testing attachment. (Do not remove prior testing information – include date of new information in red.)

Yes

2.3 For maintenance of endorsement

Risk adjustment: For outcome, resource use, cost, and some process measures, risk-adjustment that includes SDS factors is no longer prohibited during the SDS Trial Period (2015-2016). Please update sections 1.8, 2a2, 2b2, 2b4, and 2b6 in the Testing attachment and S.14 and S.15 in the online submission form in accordance with the requirements for the SDS Trial Period. NOTE: These sections must be updated even if SDS factors are not included in the risk-adjustment strategy. If yes, and your testing attachment does not have the additional questions for the SDS Trial please add these questions to your testing attachment:

What were the patient-level sociodemographic (SDS) variables that were available and analyzed in the data or sample used? For example, patient-reported data (e.g., income, education, language), proxy variables when SDS data are not collected from each patient (e.g. census tract), or patient community characteristics (e.g. percent vacant housing, crime rate).

Describe the conceptual/clinical and statistical methods and criteria used to select patient factors (clinical factors or sociodemographic factors) used in the statistical risk model or for stratification by risk (e.g., potential factors identified in the literature and/or expert panel; regression analysis; statistical significance of $p < 0.10$; correlation of x or higher; patient factors should be present at the start of care)

What were the statistical results of the analyses used to select risk factors?

Describe the analyses and interpretation resulting in the decision to select SDS factors (e.g. prevalence of the factor across measured entities, empirical association with the outcome, contribution of unique variation in the outcome, assessment of

between-unit effects and within-unit effects)
 No - This measure is not risk-adjusted

NATIONAL QUALITY FORUM—Measure Testing (subcriteria 2a2, 2b2-2b7)

Measure Number (if previously endorsed):

Measure Title: [Prescription of HIV Antiretroviral Therapy](#)

Date of Submission: [12/16/2016](#)

Type of Measure:

<input type="checkbox"/> Outcome (including PRO-PM)	<input type="checkbox"/> Composite – <i>STOP – use composite testing form</i>
<input checked="" type="checkbox"/> Intermediate Clinical Outcome	<input type="checkbox"/> Cost/resource
<input checked="" type="checkbox"/> Process	<input type="checkbox"/> Efficiency
<input type="checkbox"/> Structure	

Instructions

- Measures must be tested for all the data sources and levels of analyses that are specified. *If there is more than one set of data specifications or more than one level of analysis, contact NQF staff* about how to present all the testing information in one form.
- For all measures, sections 1, 2a2, 2b2, 2b3, and 2b5 must be completed.**
- For outcome and resource use measures**, section **2b4** also must be completed.
- If specified for **multiple data sources/sets of specifications** (e.g., claims and EHRs), section **2b6** also must be completed.
- Respond to **all** questions as instructed with answers immediately following the question. All information on testing to demonstrate meeting the subcriteria for reliability (2a2) and validity (2b2-2b6) must be in this form. An appendix for *supplemental* materials may be submitted, but there is no guarantee it will be reviewed.
- If you are unable to check a box, please highlight or shade the box for your response.
- Maximum of 20 pages (*including questions/instructions*; minimum font size 11 pt; do not change margins). **Contact NQF staff if more pages are needed.**
- Contact NQF staff regarding questions. Check for resources at [Submitting Standards webpage](#).
- For information on the most updated guidance on how to address sociodemographic variables and testing in this form refer to the release notes for version 6.6 of the Measure Testing Attachment.

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

2a2. Reliability testing ¹⁰ demonstrates the measure data elements are repeatable, producing the same results a high proportion of the time when assessed in the same population in the same time period and/or that the measure score is precise. For **PRO-PMs and composite performance measures**, reliability should be demonstrated for the computed performance score.

2b2. Validity testing ¹¹ demonstrates that the measure data elements are correct and/or the measure score correctly reflects the quality of care provided, adequately identifying differences in quality. For **PRO-PMs and composite performance measures**, validity should be demonstrated for the computed performance score.

2b3. Exclusions are supported by the clinical evidence; otherwise, they are supported by evidence of sufficient frequency of occurrence so that results are distorted without the exclusion; ¹²

AND

If patient preference (e.g., informed decisionmaking) is a basis for exclusion, there must be evidence that the exclusion impacts performance on the measure; in such cases, the measure must be specified so that the information about patient preference and the effect on the measure is transparent (e.g., numerator category computed separately, denominator exclusion category computed separately). ¹³

2b4. For outcome measures and other measures when indicated (e.g., resource use):

- **an evidence-based risk-adjustment strategy** (e.g., risk models, risk stratification) is specified; is based on patient factors (including clinical and sociodemographic factors) that influence the measured outcome and are present at start of care; ^{14,15} and has demonstrated adequate discrimination and calibration

OR

- rationale/data support no risk adjustment/ stratification.

2b5. Data analysis of computed measure scores demonstrates that methods for scoring and analysis of the specified measure allow for identification of statistically significant and practically/clinically meaningful ¹⁶ differences in performance;

OR

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

2b6. If multiple data sources/methods are specified, there is demonstration they produce comparable results.

2b7. For eMeasures, composites, and PRO-PMs (or other measures susceptible to missing data), analyses identify the extent and distribution of missing data (or nonresponse) and demonstrate that performance results are not biased due to systematic missing data (or differences between responders and nonresponders) and how the specified handling of missing data minimizes bias.

Notes

10. Reliability testing applies to both the data elements and computed measure score. Examples of reliability testing for data elements include, but are not limited to: inter-rater/abstractor or intra-rater/abstractor studies; internal consistency for multi-item scales; test-retest for survey items. Reliability testing of the measure score addresses precision of measurement (e.g., signal-to-noise).

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

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

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

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

15. With large enough sample sizes, small differences that are statistically significant may or may not be practically or clinically meaningful. The substantive question may be, for example, whether a statistically significant difference of one percentage point in the percentage of patients who received smoking cessation counseling (e.g., 74 percent v. 75 percent) is clinically meaningful; or whether a statistically significant difference of

\$25 in cost for an episode of care (e.g., \$5,000 v. \$5,025) is practically meaningful. Measures with overall less-than-optimal performance may not demonstrate much variability across providers.

1. DATA/SAMPLE USED FOR ALL TESTING OF THIS MEASURE

Often the same data are used for all aspects of measure testing. In an effort to eliminate duplication, the first five questions apply to all measure testing. If there are differences by aspect of testing, (e.g., reliability vs. validity) be sure to indicate the specific differences in question 1.7.

1.1. What type of data was used for testing? (Check all the sources of data identified in the measure specifications and data used for testing the measure. Testing must be provided for all the sources of data specified and intended for measure implementation. If different data sources are used for the numerator and denominator, indicate N [numerator] or D [denominator] after the checkbox.)

Measure Specified to Use Data From: (must be consistent with data sources entered in S.23)	Measure Tested with Data From:
<input type="checkbox"/> abstracted from paper record	<input type="checkbox"/> abstracted from paper record
<input type="checkbox"/> administrative claims	<input type="checkbox"/> administrative claims
<input type="checkbox"/> clinical database/registry	<input type="checkbox"/> clinical database/registry
<input type="checkbox"/> abstracted from electronic health record	<input type="checkbox"/> abstracted from electronic health record
<input checked="" type="checkbox"/> eMeasure (HQMF) implemented in EHRs	<input type="checkbox"/> eMeasure (HQMF) implemented in EHRs
<input checked="" type="checkbox"/> other: Synthetic Bonnie test patients	<input checked="" type="checkbox"/> other: Synthetic Bonnie test patients

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

This measure is a legacy electronic clinical quality measure (eCQM) – an NQF endorsed measure currently used in federal quality programs that has been respecified into eMeasure. Per NQF modified testing requirements for legacy eCQMs, the measure was tested in the Bonnie testing tool. Bonnie is designed to validate eCQM specifications (HQMF output and value sets) against the measure’s expected behavior for user-developed synthetic test patients.

The synthetic patient bundle used to test this measure was designed to simulate clinically relevant, realistic patient scenarios aligned with the target population for this measure. Full details on the Bonnie synthetic patient bundle used to test this measure are included in the Bonnie testing attachment.

For more information on Bonnie, please visit <https://bonnie.healthit.gov/>.

1.3. What are the dates of the data used in testing? The Bonnie test environment simulates the year 2012 as the measurement period.

1.4. What levels of analysis were tested? (testing must be provided for all the levels specified and intended for measure implementation, e.g., individual clinician, hospital, health plan)

Measure Specified to Measure Performance of: (must be consistent with levels entered in item S.26)	Measure Tested at Level of:
<input type="checkbox"/> individual clinician	<input type="checkbox"/> individual clinician

<input type="checkbox"/> group/practice	<input type="checkbox"/> group/practice
<input checked="" type="checkbox"/> hospital/facility/agency	<input checked="" type="checkbox"/> hospital/facility/agency
<input type="checkbox"/> health plan	<input type="checkbox"/> health plan
<input type="checkbox"/> other:	<input checked="" type="checkbox"/> other: Synthetic Bonnie test patients

1.5. How many and which measured entities were included in the testing and analysis (by level of analysis and data source)? (identify the number and descriptive characteristics of measured entities included in the analysis (e.g., size, location, type); if a sample was used, describe how entities were selected for inclusion in the sample)

Not applicable. The Bonnie synthetic patient bundle was used to test the measure.

1.6. How many and which patients were included in the testing and analysis (by level of analysis and data source)? (identify the number and descriptive characteristics of patients included in the analysis (e.g., age, sex, race, diagnosis); if a sample was used, describe how patients were selected for inclusion in the sample)

A test bundle of 34 patients was designed and built within the Bonnie testing tool to evaluate the measure logic. Information documented for each patient within the bundle include:

Patient name
Date of birth
Race
Ethnicity
Gender
Payer

Additional elements contained within the patient profiles as appropriate for testing against expected outcomes include:

Diagnosis
Medication orders
Encounters

The patient bundle's demographics were designed to mimic the HIV/AIDS population, specifically drawing from the patient characteristics collected via the Ryan White HIV/AIDS Program Services Report (RSR).

The breakdown of test bundle demographics for the 34 patients included (represented by number of patients/percentage of bundle): males 23/68%; females 11/32%; American Indian/Alaska Native 1/3%; Asian 1/3%; Black/African American 15/44%; Native Hawaiian/Pacific Islander 0/0%; White 9/26%; Hispanic/Latino 8/24%; younger than 13 1/3%; 13-17 years old 1/3%; 18-24 years old 2/6%; 25-34 years old 6/18%; 35-44 years old 6/18%; 45-54 years old 10/29%; 55-65 years old 6/18%; older than 65 2/6%.

Full details on the Bonnie synthetic patient bundle used to test this measure, including human-readable and QRDA Category 1 format documents for each synthetic patient record, are included in the Bonnie testing attachment.

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

The Bonnie patient test deck was used to satisfy all testing requirements for this measure. The testing results are further supported by testing data for the chart-abstracted version of this measure collected through the Health Resources and Services Administration HIV/AIDS Bureau's Ryan White HIV/AIDS Program Services Report.

1.8 What were the patient-level sociodemographic (SDS) variables that were available and analyzed in the data or sample used? For example, patient-reported data (e.g., income, education, language), proxy variables when SDS data are not collected from each patient (e.g. census tract), or patient community characteristics (e.g. percent vacant housing, crime rate).

Patient sociodemographic variables considered in the analysis of the chart-abstracted version of this measure were included in the eCQM specifications and modeled in the Bonnie patient bundle. These variables included age, race, ethnicity, gender and payer.

2a2. RELIABILITY TESTING

Note: *If accuracy/correctness (validity) of data elements was empirically tested, separate reliability testing of data elements is not required – in 2a2.1 check critical data elements; in 2a2.2 enter “see section 2b2 for validity testing of data elements”; and skip 2a2.3 and 2a2.4.*

2a2.1. What level of reliability testing was conducted? (may be one or both levels)

- Critical data elements used in the measure** (e.g., inter-abstractor reliability; data element reliability must address ALL critical data elements)
- Performance measure score** (e.g., signal-to-noise analysis)

2a2.2. For each level checked above, describe the method of reliability testing and what it tests (describe the steps—do not just name a method; what type of error does it test; what statistical analysis was used)

Currently, there is no performance data available to test the eCQM. However, the chart-abstracted version of this measure has been in use in national quality reporting programs since as early as 2010.

The most recent reliability analysis of the chart-abstracted measure was calculated according to the methods outlined in a technical report prepared by J.L. Adams for the National Committee for Quality Assurance titled “The Reliability of Provider Profiling: A Tutorial” (RAND Corporation, TR-653-NCQA, 2009). In this context, reliability represents the ability of a measure to confidently distinguish the performance of one physician from another. As discussed in the report: “Conceptually, it is the ratio of signal to noise. The signal in this case is the proportion of variability in measured performance that can be explained by real differences in performance. There are 3 main drivers of reliability; sample size, differences between physicians, and measurement error.”

According to this approach, reliability is estimated with a beta-binomial model. The beta-binomial model is appropriate for measuring the reliability of pass/fail measures such as those proposed here. Reliability scores vary from 0.0 to 1.0, with a score of zero indicating that all variation is attributable to measurement error (noise, or individual accountable entity variance) whereas a reliability of 1.0 implies that all variation is caused by real difference in performance across accountable entities.

2a2.3. For each level of testing checked above, what were the statistical results from reliability testing? (e.g., percent agreement and kappa for the critical data elements; distribution of reliability statistics from a signal-to-noise analysis)

Overall reliability scores (i.e., median of provider-level reliability [R_median], minimum [R_min], maximum [R_max]) by year, and the overall variance between sites, are summarized below.

Table 1. Overall reliability scores by year, 2010-2014

Year	% suppressed	Var_between	R_median	R_min	R_max
2010	68.4%	0.069	0.990	0.354	1.000
2011	71.1%	0.066	0.991	0.347	1.000
2012	74.3%	0.059	0.991	0.322	1.000

2013	77.5%	0.048	0.991	0.276	1.000
2014	77.6%	0.073	0.996	0.368	1.000

Reliability varied across providers by year. The proportion of providers with reliability greater than or equal to 0.9, 0.8, and 0.7 are shown below.

Table 2. Distribution of provider-level reliability scores by year, 2010-2014

Year	N	≥0.9 n (%)	≥0.8 n (%)	≥0.7 n (%)
2010	846	793 (93.7)	819 (96.8)	836 (98.8)
2011	811	752 (92.7)	788 (97.2)	792 (97.7)
2012	816	753 (92.3)	788 (96.6)	801 (98.2)
2013	823	753 (91.5)	794 (96.5)	806 (97.9)
2014	813	771 (94.8)	794 (97.7)	802 (98.7)

2a2.4 What is your interpretation of the results in terms of demonstrating reliability? (i.e., what do the results mean and what are the norms for the test conducted?)

There is no established cut-off for minimum reliability level. Values above 0.7 are considered sufficient to see differences between providers and the mean, and values above 0.9 are considered sufficient to see differences between pairs of providers (RAND Corporation, TR-653-NCQA, 2009).

Each year, more than 91% of providers had reliability scores of 0.9 or greater. Therefore, the reliability of viral suppression can be considered to be sufficient to identify real differences in performance across providers. As previously mentioned, sample size is another driver of reliability and likely contributed to the lowest reliability scores (e.g., in 2014 site 2081 had a reliability of 0.368, and reported 1 of 2 had been prescribed ART). However, median reliability was consistently 0.99 during 2010-2014, supporting the conclusion that the reliability of this measure can be considered very good.

2b2. VALIDITY TESTING

2b2.1. What level of validity testing was conducted? (may be one or both levels)

Critical data elements (data element validity must address ALL critical data elements)

Performance measure score

Empirical validity testing

Systematic assessment of face validity of performance measure score as an indicator of quality or resource use (i.e., is an accurate reflection of performance on quality or resource use and can distinguish good from poor performance)

2b2.2. For each level of testing checked above, describe the method of validity testing and what it tests (describe the steps—do not just name a method; what was tested, e.g., accuracy of data elements compared to authoritative source, relationship to another measure as expected; what statistical analysis was used)

The Bonnie testing environment was used to test the validity of the measure logic and data elements. For each Bonnie synthetic patient, an expected measure result was assigned to reflect the expected outcome of the measure given the specific patient scenario and associated data. The synthetic patients were run against the HQMF output loaded into Bonnie, which produces a measure outcome for each patient and evaluates it against the expected outcome. A patient is considered to pass Bonnie testing when the expected outcome matches the actual outcome, e.g. when a patient is expected to be in the numerator population and the computation of the

synthetic patient data against the eCQM logic places the patient in the numerator. In order to achieve a rigorous, clinically relevant test bundle, synthetic patients were designed following the below principles and test areas:

- Clinical relevance. References cited within the chart abstracted measure specification were used to design clinically relevant, realistic patient profiles for the measure’s target population. This approach ensured the eCQM logic maintained alignment with the clinical intent of the chart abstracted measure.
- 100% logic coverage: The resulting bundle of synthetic patients collectively includes all data elements and conditions logic that are specified within the measure logic, including at least one patient evaluating against each measure population pathway. Fully testing the measure logic increases test rigor and mitigates risk of unexpected outcomes.
- Edge case testing. Edge cases refer to those data elements that test the upper or lower boundary of measure logic conditions, e.g. a diagnosis starting on the latest qualifying date or a medication order for antiretroviral therapy starting on the first day or last day of the measurement period. Edge cases are designed to test each edge that exists within each measure population.
- Negative testing. Negative testing involves use of test cases do not evaluate positively against measure logic, but are otherwise clinically relevant and realistic, e.g. scenarios where an HIV diagnosis was not documented or a medication order for antiretroviral therapy starting on the first day or last day of the measurement period. Negative testing further validates measure logic by accurately evaluating patients against expected outcomes and simulating the effect of missing data on measure results.

In addition to Bonnie testing, the measure specifications were reviewed independently by three eCQM experts to confirm the logic was syntactically correct, using appropriate and current versions of the eCQM standards and terminologies, and consistent with the intent of the chart-abstracted measure.

2b2.3. What were the statistical results from validity testing? (e.g., correlation; t-test)

Bonnie testing results provide logic coverage and passing rates. The synthetic bundle reached 100% coverage, confirming each logic pathway was tested. The results also showed 100% passing rate, confirming all synthetic patients performed as expected.

Full details on Bonnie testing results are contained in the Bonnie testing attachment. The attachment includes a human-readable (HTML) summary document that lists each patient within the bundle and its passing status against expected measure outcomes. The attachment also includes a summary spreadsheet for the synthetic patient bundle which lists each patient, associated demographics, expected and actual measure population outcomes, and which portions of each measure population logic the patient meets expectations for.

2b2.4. What is your interpretation of the results in terms of demonstrating validity? (i.e., what do the results mean and what are the norms for the test conducted?)

The results of measure logic testing through use of Bonnie provided confidence in the measure logic accurately representing the clinical intent and alignment with the chart abstracted measure.

2b3. EXCLUSIONS ANALYSIS (FOR MEASURES WITH EXCLUSIONS --- gap in visits and medical visit frequency)

NA no exclusions — skip to section [2b4](#)

2b3.1. Describe the method of testing exclusions and what it tests (describe the steps—do not just name a method; what was tested, e.g., whether exclusions affect overall performance scores; what statistical analysis was used)

Not applicable.

2b3.2. What were the statistical results from testing exclusions? (include overall number and percentage of individuals excluded, frequency distribution of exclusions across measured entities, and impact on performance measure scores)

Not applicable.

2b3.3. What is your interpretation of the results in terms of demonstrating that exclusions are needed to prevent unfair distortion of performance results? (i.e., the value outweighs the burden of increased data collection and analysis. *Note: If patient preference is an exclusion, the measure must be specified so that the effect on the performance score is transparent, e.g., scores with and without exclusion*)

Not applicable.

2b4. RISK ADJUSTMENT/STRATIFICATION FOR OUTCOME OR RESOURCE USE MEASURES

If not an intermediate or health outcome, or PRO-PM, or resource use measure, skip to section 2b5.

2b4.1. What method of controlling for differences in case mix is used?

- No risk adjustment or stratification
- Statistical risk model with risk factors
- Stratification by risk categories
- Other,

2b4.1.1 If using a statistical risk model, provide detailed risk model specifications, including the risk model method, risk factors, coefficients, equations, codes with descriptors, and definitions.

Not applicable.

2b4.2. If an outcome or resource use component measure is not risk adjusted or stratified, provide rationale and analyses to demonstrate that controlling for differences in patient characteristics (case mix) is not needed to achieve fair comparisons across measured entities.

The Ryan White HIV/AIDS Program provides a comprehensive system of care that includes primary medical care and essential support services for people living with HIV who are uninsured or underinsured. The Program works with cities, states, and local community-based organizations to provide HIV care and treatment services to more than half a million people each year. The Program reaches approximately 52% of all people diagnosed with HIV in the United States.

As indicated in data presented earlier, the Ryan White HIV/AIDS Program is a public health, safety net program providing care to a high proportion of racial/ethnic minority, transgender, unstable housing, and low income people living with HIV. Many of people served by the Ryan White HIV/AIDS Program represent sociodemographics factors incorporate in risk adjusting models by many measures stewards.

As a result, the Ryan White HIV/AIDS Program does not adjust for risk in its performance measures. Rather, it is a fundamental aspect of the Ryan White HIV/AIDS Program to identify disparities and work to improve quality of care for subpopulations. Additionally, this measure is not used for pay-for-performance, bonuses, or penalties.

2b4.3. Describe the conceptual/clinical and statistical methods and criteria used to select patient factors (clinical factors or sociodemographic factors) used in the statistical risk model or for stratification by risk (e.g., potential factors identified in the literature and/or expert panel; regression analysis; statistical significance of $p < 0.10$; correlation of x or higher; patient factors should be present at the start of care)

Not applicable.

2b4.4a. What were the statistical results of the analyses used to select risk factors?

Not applicable.

2b4.4b. Describe the analyses and interpretation resulting in the decision to select SDS factors (e.g. prevalence of the factor across measured entities, empirical association with the outcome, contribution of unique variation in the outcome, assessment of between-unit effects and within-unit effects)

Not applicable.

2b4.5. Describe the method of testing/analysis used to develop and validate the adequacy of the statistical model or stratification approach (*describe the steps—do not just name a method; what statistical analysis was used*)

Provide the statistical results from testing the approach to controlling for differences in patient characteristics (case mix) below.

Not applicable.

If stratified, skip to [2b4.9](#)

2b4.6. Statistical Risk Model Discrimination Statistics (*e.g., c-statistic, R-squared*): Not applicable.

2b4.7. Statistical Risk Model Calibration Statistics (*e.g., Hosmer-Lemeshow statistic*): Not applicable.

2b4.8. Statistical Risk Model Calibration – Risk decile plots or calibration curves: Not applicable.

2b4.9. Results of Risk Stratification Analysis: Not applicable.

2b4.10. What is your interpretation of the results in terms of demonstrating adequacy of controlling for differences in patient characteristics (case mix)? (*i.e., what do the results mean and what are the norms for the test conducted*)

Not applicable.

2b4.11. Optional Additional Testing for Risk Adjustment (*not required, but would provide additional support of adequacy of risk model, e.g., testing of risk model in another data set; sensitivity analysis for missing data; other methods that were assessed*)

2b5. IDENTIFICATION OF STATISTICALLY SIGNIFICANT & MEANINGFUL DIFFERENCES IN PERFORMANCE

2b5.1. Describe the method for determining if statistically significant and clinically/practically meaningful differences in performance measure scores among the measured entities can be identified (*describe the steps—do not just name a method; what statistical analysis was used? Do not just repeat the information provided related to performance gap in 1b*)

The chart-abstracted version of this measure has been in use since 2010. To examine meaningful differences in performance, we examined the distribution of the proportion of patients with viral suppression across providers, by year. Performance scores were broken into the bottom 10% and top 90% providers to better characterize the

gaps that remain across providers. Moreover, performance scores were examined with respect the proportion of providers with least 80 percent of patients that were prescribed ART in a given year.

2b5.2. What were the statistical results from testing the ability to identify statistically significant and/or clinically/practically meaningful differences in performance measure scores across measured entities? (e.g., number and percentage of entities with scores that were statistically significantly different from mean or some benchmark, different from expected; how was meaningful difference defined)

Year	% patients with viral suppression across providers					providers with \geq 80% patients prescribed ART		
	Mean	SD	Median	10th %ile	90th %ile	N	n	%
2010	65.9%	27.5%	76.5%	17.8%	91.2%	846	353	41.7
2011	70.1%	26.4%	79.8%	26.1%	93.2%	811	402	49.6
2012	73.4%	25.4%	83.8%	31.7%	94.7%	816	471	57.7
2013	77.5%	24.1%	86.5%	42.9%	96.4%	823	532	64.6
2014	78.0%	28.0%	90.0%	29.6%	98.3%	813	565	69.5

2b5.3. What is your interpretation of the results in terms of demonstrating the ability to identify statistically significant and/or clinically/practically meaningful differences in performance across measured entities? (i.e., what do the results mean in terms of statistical and meaningful differences?)

The table above demonstrates meaningful variability across providers, allowing for the identification of meaningful differences across sites. Specifically, the measure is able to detect providers with better or worse than median performance scores. In 2014, the bottom 10% of providers had ART prescription rates of 29.6% or lower; the top 90% of providers had rates of 98.3% or higher. These differences demonstrate the continued value of the measure in identifying sites based on poor performance relative to the top performers.

Provider-level performance differences observed in the table above also underscore improvements in the proportion of patients prescribed ART. In 2014, of 813 providers, 565 (69.5%) had prescribed ART for at least 80% of patients. Additionally, on average by provider, nearly 80% (78%) of patients were prescribed ART; however, given the large population that the RWHAP serves, even the poorest performing sites (e.g., bottom 10%) represent a substantial number of patients.

2b6. COMPARABILITY OF PERFORMANCE SCORES WHEN MORE THAN ONE SET OF SPECIFICATIONS

If only one set of specifications, this section can be skipped.

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

2b6.1. Describe the method of testing conducted to compare performance scores for the same entities across the different data sources/specifications (describe the steps—do not just name a method; what statistical analysis was used)

Not applicable

2b6.2. What were the statistical results from testing comparability of performance scores for the same entities when using different data sources/specifications? (e.g., correlation, rank order)

Not applicable

2b6.3. What is your interpretation of the results in terms of the differences in performance measure scores for the same entities across the different data sources/specifications? (i.e., what do the results mean and what are the norms for the test conducted)

Not applicable

2b7. MISSING DATA ANALYSIS AND MINIMIZING BIAS

2b7.1. Describe the method of testing conducted to identify the extent and distribution of missing data (or nonresponse) and demonstrate that performance results are not biased due to systematic missing data (or differences between responders and nonresponders) and how the specified handling of missing data minimizes bias (describe the steps—do not just name a method; what statistical analysis was used)

The HQMF standard specifies that if data are unknown or missing, they shall fail the criterion. This constraint embodies the notion that absence of evidence is evidence of absence, i.e. data not present in a structured field from which the measure draws will not be considered for measure calculation. In certain cases, missing data may have no impact on the measure outcome for a given patient. For example, a data element used in a series of OR statements will not impact the measure outcome if another data element in the OR statement is present and meets all other defined constraints.

2b7.2. What is the overall frequency of missing data, the distribution of missing data across providers, and the results from testing related to missing data? (e.g., results of sensitivity analysis of the effect of various rules for missing data/nonresponse; if no empirical sensitivity analysis, identify the approaches for handling missing data that were considered and pros and cons of each)

The Bonnie synthetic patient bundle includes scenarios for missing data elements, which are a form of negative testing. All Bonnie synthetic patients with missing data performed according to the HQMF standard specification and as expected.

2b7.3. What is your interpretation of the results in terms of demonstrating that performance results are not biased due to systematic missing data (or differences between responders and nonresponders) and how the specified handling of missing data minimizes bias? (i.e., what do the results mean in terms of supporting the selected approach for missing data and what are the norms for the test conducted; if no empirical analysis, provide rationale for the selected approach for missing data)

Please see response for question 2b7.1 above.

3. Feasibility

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

3a. Byproduct of Care Processes

For clinical measures, the required data elements are routinely generated and used during care delivery (e.g., blood pressure, lab test, diagnosis, medication order).

3a.1. Data Elements Generated as Byproduct of Care Processes.

Generated or collected by and used by healthcare personnel during the provision of care (e.g., blood pressure, lab value, diagnosis, depression score)

If other:

3b. Electronic Sources

The required data elements are available in electronic health records or other electronic sources. If the required data are not in electronic health records or existing electronic sources, a credible, near-term path to electronic collection is specified.

3b.1. To what extent are the specified data elements available electronically in defined fields (i.e., data elements that are needed to compute the performance measure score are in defined, computer-readable fields) Update this field for **maintenance of endorsement**.

ALL data elements are in defined fields in electronic health records (EHRs)

3b.2. If ALL the data elements needed to compute the performance measure score are not from electronic sources, specify a credible, near-term path to electronic capture, OR provide a rationale for using other than electronic sources. For **maintenance of endorsement**, if this measure is not an eMeasure (eCQM), please describe any efforts to develop an eMeasure (eCQM).

For this measure, we are presenting an e-measure and paper measure.

3b.3. If this is an eMeasure, provide a summary of the feasibility assessment in an attached file or make available at a measure-specific URL. Please also complete and attach the NQF Feasibility Score Card.

Attachment: [NQFXXX_PrescriptionOfAntiretroviralTherapy_Feasibility_Scorecard_v1.0-636177547788569260.xlsx](#), [NQFXXX_PrescriptionOfAntiretroviralTherapy_MeasureTestingAttachment-636177547788725261.docx](#)

3c. Data Collection Strategy

Demonstration that the data collection strategy (e.g., source, timing, frequency, sampling, patient confidentiality, costs associated with fees/licensing of proprietary measures) can be implemented (e.g., already in operational use, or testing demonstrates that it is ready to put into operational use). For eMeasures, a feasibility assessment addresses the data elements and measure logic and demonstrates the eMeasure can be implemented or feasibility concerns can be adequately addressed.

3c.1. Required for maintenance of endorsement. Describe difficulties (as a result of testing and/or operational use of the measure) regarding data collection, availability of data, missing data, timing and frequency of data collection, sampling, patient confidentiality, time and cost of data collection, other feasibility/implementation issues.

IF a PRO-PM, consider implications for both individuals providing PRO data (patients, service recipients, respondents) and those whose performance is being measured.

Not applicable.

3c.2. Describe any fees, licensing, or other requirements to use any aspect of the measure as specified (e.g., value/code set, risk model, programming code, algorithm).

The measure specifications contain limited proprietary codes for convenience. Users of CPT(R) should obtain all necessary licenses from the owners of these code sets.

The use of SNOMED Clinical Terms(R) requires a Unified Medical Language System (UMLS) license. These licenses are freely available, from the National Library of Medicine.

4. Usability and Use

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

4a. Accountability and Transparency

Performance results are used in at least one accountability application within three years after initial endorsement and are publicly reported within six years after initial endorsement (or the data on performance results are available). If not in use at the time of initial endorsement, then a credible plan for implementation within the specified timeframes is provided.

4.1. Current and Planned Use

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

Specific Plan for Use	Current Use (for current use provide URL)
	<p>Public Reporting</p> <p>Ryan White HIV/AIDS Program https://hab.hrsa.gov/clinical-quality-management/performance-measure-portfolio</p> <p>Public Health/Disease Surveillance National HIV/AIDS Strategy https://www.aids.gov/federal-resources/national-hiv-aids-strategy/nhas-update.pdf</p> <p>Payment Program PQRS https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/PQRS/index.html?redirect=/pqri</p> <p>Quality Improvement (external benchmarking to organizations) Ryan White HIV/AIDS Program https://hab.hrsa.gov/clinical-quality-management/performance-measure-portfolio</p> <p>Quality Improvement (Internal to the specific organization) yan White HIV/AIDS Program https://hab.hrsa.gov/clinical-quality-management/performance-measure-portfolio</p>

4a.1. For each CURRENT use, checked above (update for maintenance of endorsement), provide:

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

Ryan White HIV/AIDS Program

Sponsor: Federal government

Geographic area: Nationwide

Accountable entities: Approximately 600 Ryan White HIV/AIDS Program grant recipients and their providers

Patients: Approximately 316,000 patients

Physician Quality Report System and Value Based Modifier

Sponsor: Federal government

Geographic area: Nationwide

Accountable entities: Physicians and practitioners

Patients: Unknown

Merit-Based Incentive Payment System

Sponsor: Federal government

Geographic area: Nationwide

Accountable entities: Physicians, Physician Assistant, Nurse Practitioner, and Clinical Nurse Specialist

Patients: Unknown

National HIV/AIDS Strategy

Sponsor: Federal government

Geographic area: Nationwide

Accountable entities: Federal agencies and service providers

Patients: All people living with HIV in the United States

4a.2. If not currently publicly reported OR used in at least one other accountability application (e.g., payment program, certification, licensing) what are the reasons? (e.g., Do policies or actions of the developer/steward or accountable entities restrict access to performance results or impede implementation?)

N/A

4a.3. If not currently publicly reported OR used in at least one other accountability application, provide a credible plan for implementation within the expected timeframes -- any accountability application within 3 years and publicly reported within 6 years of initial endorsement. (Credible plan includes the specific program, purpose, intended audience, and timeline for implementing the measure within the specified timeframes. A plan for accountability applications addresses mechanisms for data aggregation and reporting.)

N/A

Improvement

Progress toward achieving the goal of high-quality, efficient healthcare for individuals or populations is demonstrated. If not in use for performance improvement at the time of initial endorsement, then a credible rationale describes how the performance results could be used to further the goal of high-quality, efficient healthcare for individuals or populations.

4b. Refer to data provided in 1b but do not repeat here. Discuss any progress on improvement (trends in performance results, number and percentage of people receiving high-quality healthcare; Geographic area and number and percentage of accountable entities and patients included.)

If no improvement was demonstrated, what are the reasons? If not in use for performance improvement at the time of initial endorsement, provide a credible rationale that describes how the performance results could be used to further the goal of high-quality, efficient healthcare for individuals or populations.

Prescription of HIV antiretroviral therapy has been improving in the United States since the first release of publically available data. The Ryan White HIV/AIDS Program served more than 300,000 unduplicated patients annually between 2010-2014 across 2,000+ grant recipients and subrecipients. The Ryan White HIV/AIDS Program has experienced a 10 + point increase in viral suppression from 65.9% in 2010 to 78.0% in 2014. Prescription of HIV antiretroviral therapy has increased across all demographic groups and subpopulations.

4c. Unintended Consequences

The benefits of the performance measure in facilitating progress toward achieving high-quality, efficient healthcare for individuals or populations outweigh evidence of unintended negative consequences to individuals or populations (if such evidence exists).

4c.1. Please explain any unexpected findings (positive or negative) during implementation of this measure including unintended impacts on patients.

The adoption and use of this measure has continued to spread since the initial development of this measure. This measure has been adopted by Centers for Medicare and Medicaid measurement programs, Department of Health and Human Service Secretary as a one of the core HIV indicators, countless outpatient/ambulatory care settings, and health departments. National learning collaborates have used this measure to focus the improvement efforts of grant recipients and subrecipients.

Additionally, prescription of HIV antiretroviral therapy is one of five stages of the HIV care continuum. This measure has become the standard when measuring prescription of HIV antiretroviral therapy.

4c.2. Please explain any unexpected benefits from implementation of this measure.

N/A

4d1.1. Describe how performance results, data, and assistance with interpretation have been provided to those being measured or other users during development or implementation.

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

Starting in 2015, Health Resources and Services Administration began releasing December 1st – World AIDS Day – an annual data report (Ryan White HIV/AIDS Program Annual Client-Level Data Report) that contains data similar to those presenting in the report. Building upon the success of the state profiles (<http://hab.hrsa.gov/stateprofiles/>), Health Resources and Services

Administration worked diligently to release the annual data report in the same year it was collected (collected in April and released in December of the same year). The report is publically available on the Health Resources and Services Administration website (<http://hab.hrsa.gov/data/data-reports>) and is released via an accompanying webinar (recorded and archived). A supplemental report exploring data for the eligible metropolitan areas and transitional grant areas and youth/young adults has been released as well as slides sets for fact sheets by program and population, special populations (<http://hab.hrsa.gov/publications/hivaids-bureau-fact-sheets>), and infographics (contained in fact sheets). Additionally, grant recipient level reports are prepared and disseminated to all Ryan White HIV/AIDS Program grant recipients.

4d1.2. Describe the process(es) involved, including when/how often results were provided, what data were provided, what educational/explanatory efforts were made, etc.

Starting in 2015, Health Resources and Services Administration began releasing December 1st – World AIDS Day – an annual data report (Ryan White HIV/AIDS Program Annual Client-Level Data Report) that contains data similar to those presenting in the report. Building upon the success of the state profiles (<http://hab.hrsa.gov/stateprofiles/>), Health Resources and Services Administration worked diligently to release the annual data report in the same year it was collected (collected in April and released in December of the same year). The report is publically available on the Health Resources and Services Administration website (<http://hab.hrsa.gov/data/data-reports>) and is released via an accompanying webinar (recorded and archived). A supplemental report exploring data for the eligible metropolitan areas and transitional grant areas and youth/young adults has been released as well as slides sets for fact sheets by program and population, special populations (<http://hab.hrsa.gov/publications/hivaids-bureau-fact-sheets>), and infographics (contained in fact sheets). Additionally, grant recipient level reports are prepared and disseminated to all Ryan White HIV/AIDS Program grant recipients.

4d2.1. Summarize the feedback on measure performance and implementation from the measured entities and others described in 4d.1.

Describe how feedback was obtained.

Antidotal feedback has been received from Ryan White HIV/AIDS Program grant recipients and subrecipients regarding the feasibility and usefulness of the data presented in the Ryan White HIV/AIDS Program Annual Client-Level Data Report. Significant feedback has been provided about the timeliness and expansions of the data release. Grant recipient report using the data for benchmarking their program, setting goals/targets, and gaining a fuller understanding of all aspects of the Ryan White HIV/AIDS Program (i.e. other regions of the country). Grant recipients and subrecipients have also requested additional analyses. Health Resources and Services Administration responded with supplemental reports (Ryan White HIV/AIDS Program Supplemental Client-Level Data Report, Eligible Metropolitan Areas and Transitional Grant Areas; special population reports); slide decks for the overall client population and special populations; grant recipient reports; and infographics – all of which will be updated and released annually. Health Resources and Services Administration plans to release additional analyses and special reports this year based on feedback from Ryan White HIV/AIDS Program grant recipients and subrecipients.

4d2.2. Summarize the feedback obtained from those being measured.

See 4d2.2

4d2.3. Summarize the feedback obtained from other users

Ryan White HIV/AIDS Program national partners (national organizations that represent grant recipients, subrecipients, and patients) has provided antidotal feedback regarding the timeliness, feasibility, and usability of the release of the Ryan White HIV/AIDS Program Annual Client-Level Data Report, supplemental reports, slide decks, fact sheets, and infographics. The national partners encourage the continued release of the data in all its formats.

4d.3. Describe how the feedback described in 4d.2 has been considered when developing or revising the measure specifications or implementation, including whether the measure was modified and why or why not.

During the initial development of the measure, formal feedback was gathered. The measures were modified during the development phase and have not been modified since. A concerted effort was made to develop a measure that would likely stand the test of time from a scientific, clinical, and patient perspective. On an annual basis, the measure is review for clinical relevance, change in scientific acceptability, and consistency with guidelines. This measure has not been modified as a result of the annual reviews. Additionally, this measure is used by a number of measurement programs and strategies. Each of those programs require a separate annual review. No modifications have been made for those programs.

5. Comparison to Related or Competing Measures

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

5. Relation to Other NQF-endorsed Measures

Are there related measures (conceptually, either same measure focus or target population) or competing measures (conceptually both the same measure focus and same target population)? If yes, list the NQF # and title of all related and/or competing measures.

Yes

5.1a. List of related or competing measures (selected from NQF-endorsed measures)

5.1b. If related or competing measures are not NQF endorsed please indicate measure title and steward.

0405 HIV/AIDS: Pneumocystis Jiroveci Pneumonia (PCP) Prophylaxis

0409 HIV/AIDS: Sexually Transmitted Disease Screening for Chlamydia, Gonorrhea, and Syphilis

2079 HIV Medical Visit Frequency

2080 Gap in HIV Medical Visits

2082 HIV Viral Suppression

3210 HIV Viral Suppression

3010 HIV Medical Visit Frequency

5a. Harmonization of Related Measures

The measure specifications are harmonized with related measures;

OR

The differences in specifications are justified

5a.1. If this measure conceptually addresses EITHER the same measure focus OR the same target population as NQF-endorsed measure(s):

Are the measure specifications harmonized to the extent possible?

Yes

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

This measure does not have a competing measure.

5b. Competing Measures

The measure is superior to competing measures (e.g., is a more valid or efficient way to measure);

OR

Multiple measures are justified.

5b.1. If this measure conceptually addresses both the same measure focus and the same target population as NQF-endorsed measure(s):

Describe why this measure is superior to competing measures (e.g., a more valid or efficient way to measure quality); OR provide a rationale for the additive value of endorsing an additional measure. (Provide analyses when possible.)

Appendix

A.1 Supplemental materials may be provided in an appendix. All supplemental materials (such as data collection instrument or methodology reports) should be organized in one file with a table of contents or bookmarks. If material pertains to a specific

submission form number, that should be indicated. Requested information should be provided in the submission form and required attachments. There is no guarantee that supplemental materials will be reviewed.

Attachment:

Contact Information

Co.1 Measure Steward (Intellectual Property Owner): Health Resources and Services Administration - HIV/AIDS Bureau

Co.2 Point of Contact: Marlene, Matosky, mmatosky@hrsa.gov, 301-443-0798-

Co.3 Measure Developer if different from Measure Steward: Health Resources and Services Administration - HIV/AIDS Bureau

Co.4 Point of Contact: Marlene, Matosky, mmatosky@hrsa.gov, 301-443-0798-

Additional Information

Ad.1 Workgroup/Expert Panel involved in measure development

Provide a list of sponsoring organizations and workgroup/panel members' names and organizations. Describe the members' role in measure development.

Employees of the following governmental and non-governmental organizations/agencies participated in the development of this measure and assisted in assessing face validity:

- HHS Office of HIV/AIDS and Infectious Disease Policy
- Centers for Disease Control
- Center for Medicaid and Medicare
- Health Resources and Services Administration
- Indian Health Service
- National Institutes of Health
- Substances Abuse and Mental Health Services Administration
- U.S. Department of Veterans Affairs
- HIV Medical Association
- Kaiser Permanente
- National Associate of State and Territorial AIDS Directors
- Urban Coalition for HIV/AIDS Prevention Services
- National Minority AIDS Council
- Iowa Department of Health
- Washington D.C. Department of Health
- Maryland Department of Health
- University of Alabama
- University of San Francisco
- Johns Hopkins University

Measure Developer/Steward Updates and Ongoing Maintenance

Ad.2 Year the measure was first released: 2011

Ad.3 Month and Year of most recent revision: 05, 2016

Ad.4 What is your frequency for review/update of this measure? Annual

Ad.5 When is the next scheduled review/update for this measure? 05, 2016

Ad.6 Copyright statement: None

Ad.7 Disclaimers: None

Ad.8 Additional Information/Comments: It is our intention that this measure will be used in quality improvement in addition to public reporting. As it is involved in quality improvement, it is not our intent that the performance goal will be 100%. When we do set the performance goal, we will take into consideration appropriate reasons why the patient may not be able to meet the numerator criterion.