MEDICARE-MEDICAID
CAPITATED FINANCIAL ALIGNMENT MODEL
REPORTING REQUIREMENTS:
NEW YORK FIDA-IDD-SPECIFIC REPORTING REQUIREMENTS

Issued February 26, 2021
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New York FIDA-IDD-Specific Reporting Requirements Appendix

Introduction

The measures in this appendix are required reporting for the plan participating in the New York Fully Integrated Duals Advantage for Individuals with Intellectual and Developmental Disabilities (FIDA-IDD) Demonstration. CMS and the State of New York reserve the right to update the measures in this appendix for subsequent demonstration years. These state-specific measures directly supplement the Medicare-Medicaid Capitated Financial Alignment Model Core Reporting Requirements, which can be found at the following web address:


The FIDA-IDD Plan should refer to the core document for additional details regarding definitions, reporting phases and timelines, and sampling methodology, except as otherwise specified in this document.

The core and state-specific measures supplement existing Part C and Part D Reporting Requirements, as well as measures that the FIDA-IDD Plan reports via other vehicles or venues, such as HEDIS® and HOS. CMS and the State will also track key utilization measures, which are not included in this document, using encounter and claims data. The quantitative measures are part of broader oversight, monitoring, and performance improvement processes that include several other components and data sources not described in this document.

The FIDA-IDD Plan should contact the FIDA-IDD HelpDesk at FIDA-IDDHelpDesk@norc.org with any questions about the New York FIDA-IDD state-specific appendix or the data submission process.

Definitions

Calendar Quarter: All quarterly measures are reported on calendar quarters. The four calendar quarters of each calendar year will be as follows: January 1 to March 31, April 1 to June 30, July 1 to September 30, and October 1 to December 31.

Calendar Year: All annual measures are reported on a calendar year basis. For example, Calendar Year (CY) 2021 represents January 1, 2021 through December 31, 2021.

Community-based Long Term Services and Supports (LTSS): A range of medical, habilitation, rehabilitation, home care, or social services a person needs over months or years in order to improve or maintain function or health which are provided in the person’s home or community-based setting. These home and community-based services are designed to meet an individual’s needs as an alternative to long-term care in a nursing facility or intermediate care facility for individuals with intellectual disabilities

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1 HEDIS® is a registered trademark of the National Committee for Quality Assurance (NCQA).
(ICF-IID) and to enable a person to live as independently as possible. Examples include assistance with bathing, dressing, and other basic activities of daily life and self-care, as well as support for everyday tasks such as laundry, shopping, and transportation.

**Facility-based Long-Term Services and Supports (LTSS):** Facility-based LTSS are a range of medical, social, habilitation, or rehabilitation services a person needs over months or years in order to improve or maintain function or health which are provided in a long-term care facility, such as a nursing facility or intermediate care facility, ICF-IID (not including assisted living residences).

**Implementation Period:** The initial months of the demonstration during which the FIDA-IDD Plan reported to CMS and the State on a more intensive reporting schedule. The Implementation Period started on the first effective enrollment date and continued for six months (April 1, 2016 – September 30, 2016).

**Primary Care Provider (PCP):** Primary care physicians licensed by the State of New York and board certified in family practice, internal medicine, general practice, obstetrics/gynecology, or geriatrics, State licensed physician assistants, or a physician extender who is a registered nurse practitioner or advanced practice nurse or advanced practice nurse group practice within an acceptable specialty as required under State regulation.

**Interdisciplinary Team (IDT):** The team of individuals that will provide person-centered care management to Participants. Each Participant will have an IDT.

**Variation from the Core Reporting Requirements Document**

**Core 9.2**

The following section provides additional guidance about identifying individuals enrolled in the FIDA-IDD Plan as “nursing home certifiable,” or meeting the nursing facility level of care (NF LOC), for the purposes of reporting Core 9.2.

Core 9.2 focuses on “nursing home certifiable” members, defined as “members living in the community, but requiring an institutional level of care” (see the Core Reporting Requirements for more information). The FIDA-IDD Plan should include in reporting for “nursing home certifiable” all Participants meeting the intermediate care facility level of care (ICF LOC). The FIDA-IDD Plan must confirm that such Participants are living in the community and not in long-term nursing facility stays or other institutional settings.

**Quality Withhold Measures**

CMS and the State of New York established a set of quality withhold measures, and the FIDA-IDD Plan is required to meet established thresholds. Throughout this document, state-specific quality withhold measures are marked with the following symbol for Demonstration Year 1: (i) and the following symbol for Demonstration Years 2 through 7: (ii). Note that additional state-specific quality withhold measures are reported separately through HEDIS®.² For more information about the state-specific quality withhold measures, refer to the Quality Withhold Technical Notes (DY 1): New York

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² HEDIS® is a registered trademark of NCQA.
Reporting on Comprehensive Service Planning Assessments (CSPAs) and Life Plans (LPs) Completed Prior to First Effective Enrollment Date

The FIDA-IDD Plan may complete CSPAs prior to individuals’ effective date of enrollment, provided that the FIDA-IDD Plan meets the requirements as articulated in the National MMP Enrollment and Disenrollment Guidance. The following section provides information on how the FIDA-IDD Plan should report completion of early CSPAs.

For purposes of reporting data on assessments (Core 2.1 and Core 2.2), the FIDA-IDD Plan should report any CSPAs completed prior to the first effective enrollment date as if they were completed on the first effective enrollment date. For example, if a Participant’s first effective enrollment date was June 1 and the CSPA for that Participant was completed on May 25, the FIDA-IDD Plan should report the CSPA as if it were completed on June 1.

The FIDA-IDD Plan must comply with the IDT Policy regarding completion of LPs within 90 days of enrollment. In the event that an LP is also finalized prior to the first effective enrollment date, the FIDA-IDD Plan should report completion of the LP (for measures Core 3.2 and IDD1.2) as if it were completed on the first effective enrollment date. For example, if a Participant’s first effective enrollment date was June 1 and the LP for that Participant was completed on May 27, the FIDA-IDD Plan should report the LP as if it were completed on June 1.

Guidance on CSPAs and LPs for Participants with a Break in Coverage

CSPAs

If the FIDA-IDD Plan already completed a CSPA for a Participant who was previously enrolled, the FIDA-IDD Plan is not necessarily required to conduct a new CSPA if the Participant rejoins the FIDA-IDD Plan within one year of his/her most recent CSPA. Instead, the FIDA-IDD Plan can:

1. Perform any risk stratification, claims data review, or other analyses as required by the three-way contract and IDT Policy to detect any changes in the Participant’s condition since the CSPA was conducted; and
2. Ask the Participant (or his/her authorized representative) if there has been a change in the Participant’s health status or needs since the CSPA was conducted.

The FIDA-IDD Plan must document any risk stratification, claims data review, or other analyses that are performed to detect any changes in the Participant’s condition. The FIDA-IDD Plan must also document its outreach attempts and the discussion(s) with the Participant (or his/her authorized representative) to determine if there was a change in the Participant’s health status or needs.

If a change is identified, the FIDA-IDD Plan must conduct a new CSPA within the timeframe prescribed by the three-way contract and IDT Policy. If there are no changes, the FIDA-IDD Plan is not required to conduct a new CSPA unless requested by the Participant (or his/her authorized representative). Please note, if the FIDA-IDD Plan prefers to conduct CSPAs on all re-enrollees regardless of status, it may continue to do so.

Once the FIDA-IDD Plan has conducted a new CSPA as needed or confirmed that the prior CSPA is still accurate, the FIDA-IDD Plan can mark the CSPA as complete for the Participant’s current enrollment. The FIDA-IDD Plan would then report that completion according to the specifications for Core 2.1 and Core 2.2. When reporting these core measures, the FIDA-IDD Plan should count the number of enrollment days from the Participant’s most recent enrollment effective date and should report the CSPA based on the date the prior CSPA was either confirmed to be accurate or a new CSPA was completed. Additionally, in certain circumstances a new CSPA that has been completed for a Participant upon reenrollment may also be reported in Core 2.3.

If the FIDA-IDD Plan is unable to reach a re-enrolled Participant to determine if there was a change in health status, then the FIDA-IDD Plan may report that Participant as unable to be reached so long as the FIDA-IDD Plan made the requisite number of outreach attempts. If a re-enrolled Participant refuses to discuss his/her health status with the FIDA-IDD Plan, then the FIDA-IDD Plan may report that Participant as unwilling to participate in the CSPA.

If the FIDA-IDD Plan did not complete a CSPA for the re-enrolled Participant during his/her prior enrollment period, or if it has been more than one year since the Participant’s CSPA was completed, the FIDA-IDD Plan is required to conduct a CSPA for the Participant within the timeframe prescribed by the three-way contract and IDT Policy. The FIDA-IDD Plan must make the requisite number of attempts to reach the Participant (at minimum) after his/her most recent enrollment effective date, even if the FIDA-IDD Plan reported that the Participant was unable to be reached during his/her prior enrollment. Similarly, Participants who refused the CSPA during his/her prior enrollment must be asked again to participate (i.e., the FIDA-IDD Plan may not carry over a refusal from one enrollment period to the next).

LPs

If the FIDA-IDD Plan conducts a new CSPA for the re-enrolled Participant, the IDT must revise the LP accordingly within the timeframe prescribed by the three-way contract and IDT Policy. Once the LP is revised, the FIDA-IDD Plan may mark the LP as complete for
the Participant’s current enrollment. If the FIDA-IDD Plan determines that the prior CSPA is still accurate and therefore the IDT does not need to update the previously completed LP, the FIDA-IDD Plan may mark the LP as complete for the current enrollment at the same time that the CSPA is marked complete. The FIDA-IDD Plan would then follow the Core 3.2 and IDD1.2 measure specifications for reporting the completion. Please note, for purposes of reporting, the LP for the re-enrolled Participant should be classified as an initial LP.

If the IDT did not complete an LP for the re-enrolled Participant during his/her prior enrollment period, or if it has been more than one year since the Participant’s LP was completed, the IDT is required to complete an LP for the Participant within the timeframe prescribed by the three-way contract and IDT Policy. The IDT must also follow the above guidance regarding reaching out to Participants who previously refused to participate or were not reached.

Comprehensive Reassessments (CRs) and LP updates

The FIDA-IDD Plan must follow the three-way contract requirements and the IDT Policy regarding the completion of CRs at least annually based on the completion date of the previous CSPA, and the IDT should update the LP as necessary within 30 days of the CR. If the FIDA-IDD Plan determined that a CSPA from a Participant’s prior enrollment was accurate and marked that CSPA as complete for the Participant’s current enrollment, the FIDA-IDD Plan should count from the date that the CSPA was completed in the prior enrollment period to determine the due date for the CR and LP update. For example, when reporting Core 2.3, the FIDA-IDD Plan should count 365 days from the date when the CSPA was actually completed, even if that date was during the Participant’s prior enrollment period.

Reporting on Passively Enrolled and Opt-In Enrolled Participants

When reporting all New York FIDA-IDD state-specific measures, the FIDA-IDD Plan should include all Participants who meet the criteria for inclusion in the measure regardless of whether the Participant was enrolled through passive enrollment or opt-in enrollment. Medicaid-only Participants should not be included.

Reporting on Disenrolled and Retro-disenrolled Participants

Unless otherwise indicated in the Reporting Requirements, the FIDA-IDD Plan should report on all Participants enrolled in the demonstration who meet the definition of the data elements, regardless of whether that Participant was subsequently disenrolled from the FIDA-IDD Plan. Measure-specific guidance on how to report on disenrolled Participants is provided under the Notes section of each state-specific measure.

Due to retro-disenrollment of Participants, there may be instances where there is a lag between a Participant’s effective disenrollment date and the date on which the FIDA-IDD Plan is informed about that disenrollment. This time lag might create occasional data inaccuracies if the FIDA-IDD Plan includes Participants in reports who had in fact disenrolled before the start of the reporting period. If the FIDA-IDD Plan is aware at the time of reporting that a Participant has been retro-disenrolled with a disenrollment effective date prior to the reporting period (and, therefore, was not enrolled during the
reporting period in question), then the FIDA-IDD Plan may exclude that Participant from reporting. Please note that the FIDA-IDD Plan is not required to re-submit corrected data should it be informed of a retro-disenrollment subsequent to a reporting deadline. The FIDA-IDD Plan should act upon its best and most current knowledge at the time of reporting regarding each Participant’s enrollment status.

Value Sets

The measure specifications in this document refer to code value sets that must be used to determine and report measure data element values. A value set is the complete set of codes used to identify a service or condition included in a measure. The New York FIDA-IDD-Specific Value Sets Workbook includes all value sets and codes needed to report certain measures included in the New York FIDA-IDD-Specific Reporting Requirements and is intended to be used in conjunction with the measure specifications outlined in this document. The New York FIDA-IDD-Specific Value Sets Workbook can be found on the CMS website at the following address: https://www.cms.gov/Medicare-Medicaid-Coordination/Medicare-and-Medicaid-Coordination/Medicare-Medicaid-Coordination-Office/FinancialAlignmentInitiative/MMPIInformationandGuidance/MMPRreportingRequirements.html.

New York FIDA-IDD’s Implementation, Ongoing, and Continuous Reporting Periods

<table>
<thead>
<tr>
<th>Phase</th>
<th>Dates</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demonstration Year 1</td>
<td></td>
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<tr>
<td>Continuous Reporting</td>
<td>Implementation Period</td>
<td>4-1-16 through 9-30-16</td>
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<tr>
<td>Ongoing Period</td>
<td>4-1-16 through 12-31-17</td>
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<tr>
<td>Demonstration Year 2</td>
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<td>Continuous Reporting</td>
<td>Ongoing Period</td>
<td>1-1-18 through 12-31-18</td>
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<td>Demonstration Year 3</td>
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<td>Continuous Reporting</td>
<td>Ongoing Period</td>
<td>1-1-19 through 12-31-19</td>
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<tr>
<td>Phase</td>
<td>Dates</td>
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<tr>
<td>---------------</td>
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<td>Demonstration Year 4</td>
<td>Continuous Reporting</td>
<td>Ongoing Period</td>
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<td>Demonstration Year 5</td>
<td>Continuous Reporting</td>
<td>Ongoing Period</td>
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<td>Demonstration Year 6</td>
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<td>Ongoing Period</td>
</tr>
<tr>
<td>Demonstration Year 7</td>
<td>Continuous Reporting</td>
<td>Ongoing Period</td>
</tr>
</tbody>
</table>

**Data Submission**

The FIDA-IDD Plan will submit state-specific measure data through the web-based Financial Alignment Initiative Data Collection System (FAI DCS), unless otherwise specified in the measure description. All data submissions must be submitted to this site by 5:00 p.m. ET on the applicable due date. This site can be accessed at the following web address: [https://Financial-Alignment-Initiative.NORC.org](https://Financial-Alignment-Initiative.NORC.org).

(Note: Prior to the first use of the system, the FIDA-IDD Plan will receive an email notification with the username and password that it has been assigned. This information will be used to log in to the FAI DCS and complete the data submission).

The FIDA-IDD Plan will submit core measure data in accordance with the Core Reporting Requirements. Submission requirements vary by measure, but most core measures are reported through the Health Plan Management System (HPMS).

Please note, late submissions may result in compliance action from CMS.

**Resubmission of Data**

The FIDA-IDD Plan must comply with the following steps to resubmit data after an established due date:

1. Email the FIDA-IDD HelpDesk ([FIDA-IDDHelpDesk@norc.org](mailto:FIDA-IDDHelpDesk@norc.org)) to request resubmission.
   a. Specify in the email which measure(s) need resubmission;
b. Specify for which reporting period(s) the resubmission is needed; and

c. Provide a brief explanation for why the data need to be resubmitted.

2. After review of the request, the FIDA-IDD HelpDesk will notify the FIDA-IDD Plan once the FAI DCS and/or HPMS has been re-opened.

3. Resubmit data through the applicable reporting system.

4. Notify the FIDA-IDD HelpDesk again after resubmission has been completed.

Please note, requests for resubmission after an established due date may result in compliance action from CMS.
Section FIDA-IDD I. Care Coordination

IDD1.1 Participants with Life Plans (LPs) updated within 30 days of a Comprehensive Reassessment (CR).

### IMPLEMENTATION

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
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<td>FIDA-IDD1. Care Coordination</td>
<td>Monthly</td>
<td>Contract</td>
<td>Current Month Ex: 1/1-1/31</td>
<td>By the end of the third month following the last day of the reporting period</td>
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### ONGOING

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<tr>
<th>Reporting Section</th>
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<td>FIDA-IDD1. Care Coordination</td>
<td>Quarterly</td>
<td>Contract</td>
<td>Current Calendar Quarter Ex: 1/1-3/31 4/1-6/30 7/1-9/30 10/1-12/31</td>
<td>By the end of the third month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Comprehensive Reassessments (CRs) completed during the reporting period</td>
<td>Total number of CRs completed during the reporting period for Participants who were continuously enrolled for 30 days after the completion of the CR, with no gaps in enrollment.</td>
<td>Field Type: Numeric</td>
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<tr>
<td>Element Letter</td>
<td>Element Name</td>
<td>Definition</td>
<td>Allowable Values</td>
</tr>
<tr>
<td>---------------</td>
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<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------------------------------------</td>
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<tr>
<td>B.</td>
<td>Total number of CRs for which the Participant was documented as unwilling to participate in the revised LP process within 30 days after the completion of the CR.</td>
<td>Of the total reported in A, the number of CRs for which the Participant was documented as unwilling to participate in the revised LP process within 30 days after the completion of the CR.</td>
<td>Field Type: Numeric</td>
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<tr>
<td></td>
<td></td>
<td>Note: Is a subset of A.</td>
<td></td>
</tr>
<tr>
<td>C.</td>
<td>Total number of CRs for which the Participant was unable to be reached, following no fewer than three documented outreach attempts, to complete the revised LP within 30 days after the completion of the CR.</td>
<td>Of the total reported in A, the number of CRs for which the Participant was unable to be reached to revise the LP, following no fewer than three documented outreach attempts, to complete the revised LP within 30 days after the completion of the CR.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Note: Is a subset of A.</td>
<td></td>
</tr>
<tr>
<td>D.</td>
<td>Total number of CRs for which a revised LP was completed within 30 days after the completion of the CR.</td>
<td>Of the total reported in A, the number of CRs for which a revised LP was completed within 30 days after the completion of the CR.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Note: Is a subset of A.</td>
<td></td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.

- The FIDA-IDD Plan should validate that data elements B, C, and D are less than or equal to data element A.
D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored. CMS and the state will evaluate the percentage of CRs completed for which:

- Participants were documented as unwilling to participate in the revised LP process within 30 days after the completion of the CR.
  - Percentage = \( \frac{B}{A} \times 100 \)
- Participants were unable to be reached to have a revised LP completed within 30 days of the completion of the CR.
  - Percentage = \( \frac{C}{A} \times 100 \)
- A revised LP was completed within 30 days after the completion of the CR.
  - Percentage = \( \frac{D}{A} \times 100 \)
- Participants were willing to participate, could be reached and for whom a revised LP was completed within 30 days after the completion of the CR.
  - Percentage = \( \frac{D}{A - B - C} \times 100 \)

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

**Data Element A**

- The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are slated for disenrollment as of the end of the reporting period (i.e., include all Participants whose 30th day of coverage following the completion of a CR falls within the reporting period, even if that day is his/her last effective day of coverage).

**Data Element A Exclusion**

- For data element A, exclude CRs where the Participant has a trigger event that occurs within 30 days after completing the CR. These CRs are excluded because the trigger event results in the need for the Participant to undergo another CR. Any subsequent CR should be reported in this measure in the reporting period in which it occurs.

**Data Element B**

- For data element B, the FIDA-IDD Plan should report the number of CRs for which Participants were documented as unwilling to participate in the revised LP process if a Participant (or his or her authorized representative):
  - Affirmatively declines to participate in the LP process, affirmatively declines care management activities overall, or refuses any contact with the FIDA-IDD Plan. The Participant may communicate the declination or refusal by phone, mail, fax, or in person. The declination or refusal must be documented by the FIDA-IDD Plan.
  - Expresses willingness to complete the LP process but asks for it to be conducted after the indicated timeframe (despite being offered a reasonable opportunity to complete the LP process within that timeframe). Discussions with the Participant must be documented by the FIDA-IDD Plan.
• Schedules an appointment to complete the LP process, but cancels or is a no-show and then is subsequently non-responsive to additional outreach attempts by the FIDA-IDD Plan. All attempts to contact the Participant must be documented by the FIDA-IDD Plan.

• Initially agrees to complete the LP process, but then declines to participate in the LP process. The declination must be documented by the FIDA-IDD Plan.

• If a Participant was not reached after three outreach attempts, but then subsequently is reached and refuses to complete the LP process within the specified timeframe, the CR for that Participant should be classified in data element B.

Data Element C

• For data element C, the FIDA-IDD Plan should report the number of CRs for which the FIDA-IDD Plan was unable to reach the Participant to revise the LP after three documented outreach attempts to contact the Participant.

• For Participants reported in data element C, the three documented outreach attempts to contact the Participant should be specific to updating the LP after completing the CR. The FIDA-IDD Plan should refer to the NY FIDA-IDD three-way contract, IDT Policy, or state guidance for any specific requirements pertaining to the method of outreach to Participants.

• The FIDA-IDD Plan must document each attempt to reach the Participant, including the method of the attempt (e.g., phone, mail, or email), as CMS and the state may validate this number.

• There may be instances when the FIDA-IDD Plan has a high degree of confidence that a Participant’s contact information is correct, yet that Participant is not responsive to the FIDA-IDD Plan’s outreach efforts. So long as the FIDA-IDD Plan follows the guidance regarding outreach attempts, these Participants may be included in the count for this data element.

• If the FIDA-IDD Plan was previously able to reach a Participant for the purpose of completing a CR, at least three new and distinct outreach attempts for the purpose of completing the LP must be made and documented.

Data Element D

• If a CR is completed and the FIDA-IDD Plan reviews the LP within 30 days of the completion of that CR and finds that no revisions to the LP are necessary, the FIDA-IDD Plan may report this CR in data element D.

• The FIDA-IDD Plan should only report revised LPs where the Participant or the Participant’s authorized representative was involved in the revision of the LP.

• If a Participant initially refused to participate in the LP process or could not be reached after three outreach attempts, but then subsequently completes an LP within the indicated timeframe, the CR for that Participant should be classified in data element D.
General Guidance

- For purposes of reporting, 30 days from the date of the CR completion will be equivalent to one full calendar month.
- The FIDA-IDD Plan should refer to the IDT Policy and the FIDA-IDD three-way contract for specific requirements pertaining to LPs and CRs.
- The CR must be completed within the reporting period, but the LP may not be completed in the same reporting period. For example, if a CR is completed less than 30 days before the end of the quarterly reporting period (e.g., March 15), look up to 30 days past the end of the reporting period to identify whether an LP was completed.
- There may be certain circumstances that make it impossible or inappropriate to complete an LP within the specified timeframe.
  - For example, a Participant may become medically unable to respond and have no authorized representative to do so on their behalf, or a Participant may be experiencing an acute medical or behavioral health crisis that requires immediate attention and outweighs the need for an LP. However, the FIDA-IDD Plan should not include such CRs in the counts for data elements B or C.
- If an LP was started but not completed within 30 days of a completed CR, then the LP should not be considered completed and, therefore, would not be counted in data elements B, C, or D. However, this Participant would be included in data element A if a CR was completed within the reporting period.
- CRs reported in data elements B, C, and D must also be reported in data element A, since these data elements are subsets of data element A. Additionally, subset data elements should be mutually exclusive (e.g., a Participant reported in data element B or C should not also be reported in data element D).

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.
### IMPLEMENTATION

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD1. Care Coordination</td>
<td>Monthly</td>
<td>Contract</td>
<td>Current Month Ex: 1/1-1/31</td>
<td>By the end of the month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

### ONGOING

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Periods</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD1. Care Coordination</td>
<td>Quarterly</td>
<td>Contract</td>
<td>Current Calendar Quarter Ex: 1/1-3/31 4/1-6/30 7/1-9/30 10/1-12/31</td>
<td>By the end of the second month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants with an initial LP completed.</td>
<td>Total number of Participants with an initial LP completed during the reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of Participants with at least one documented discussion of care goals in the initial LP.</td>
<td>Of the total reported in A, the number of Participants with at least one documented discussion of care goals in the initial LP.</td>
<td>Field Type: Numeric Note: Is a subset of A.</td>
</tr>
<tr>
<td>C.</td>
<td>Total number of existing LPs revised.</td>
<td>Total number of existing LPs revised during the reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>Element Letter</td>
<td>Element Name</td>
<td>Definition</td>
<td>Allowable Values</td>
</tr>
<tr>
<td>----------------</td>
<td>--------------</td>
<td>------------</td>
<td>------------------</td>
</tr>
<tr>
<td>D.</td>
<td>Total number of revised LPs with at least one documented discussion of new or existing care goals.</td>
<td>Of the total reported in C, the number of revised LPs with at least one documented discussion of new or existing care goals.</td>
<td>Field Type: Numeric Note: Is a subset of C.</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.

- The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.
- The FIDA-IDD Plan should validate that data element D is less than or equal to data element C.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored. CMS and the state will evaluate the percentage of:

- Participants with an initial LP completed during the reporting period who had at least one documented discussion of care goals in the initial LP.
  - Percentage = (B / A) * 100
- Existing LPs revised during the reporting period that had at least one documented discussion of new or existing care goals.
  - Percentage = (D / C) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

**Data Element A**

- The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of the end of the reporting period (i.e., include all Participants regardless of whether they are currently enrolled or disenrolled as of the last day of the reporting period).
- Data element A should include all Participants whose LP was completed for the first time during the reporting period (i.e., the Participant did not previously have an LP completed prior to the start of the reporting period). There can be no more than one initial LP completed per Participant.
- Only LPs that included participation from the Participant (or his/her authorized representative) in the completion of the LP should be reported.
Data Element B

- The FIDA-IDD Plan should only include Participants in data element B when the discussion of care goals with the Participant (or his/her authorized representative) is clearly documented in the Participant’s initial LP.

Data Element C

- The FIDA-IDD Plan should include all LPs that meet the criteria outlined in data element C, regardless of whether the Participants are disenrolled as of the end of the reporting period (i.e., include all LPs regardless of whether the Participants are currently enrolled or disenrolled as of the last day of the reporting period).
- Data element C should include all existing LPs that were revised during the reporting period. The FIDA-IDD Plan should refer to the IDT Policy and the FIDA-IDD three-way contract for specific requirements pertaining to updating the LP.
- Only LPs that included participation from the Participant (or his/her authorized representative) in the revision to the LP should be reported.
- If a Participant’s LP is revised multiple times during the same reporting period, each revision should be reported in data element C.
  - For example, if a Participant’s LP is revised twice during the same reporting period, two LPs should be counted in data element C.

Data Element D

- The FIDA-IDD Plan should only include LPs in data element D when a new or previously documented care goal is discussed with the Participant (or his/her authorized representative) and is clearly documented in the Participant’s revised LP.
- If the initial LP clearly documented the discussion of care goals, but those existing care goals were not revised or discussed, or new care goals are not discussed and documented during the revision of the LP, then that LP should not be reported in data element D.

General Guidance

- If a Participant has an initial LP completed during the reporting period, and has their LP revised during the same reporting period, then the Participant should be reported in data element A and the Participant’s revised LP should be reported in data element C.

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.
 IDD1.3  Participants with first follow-up visit within 30 days of hospital discharge.

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD1. Care Coordination</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year</td>
<td>By the end of the fourth month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of acute inpatient hospital discharges.</td>
<td>Total number of acute inpatient hospital discharges that occurred during the reporting period for Participants who were continuously enrolled from the date of the inpatient hospital discharge through 30 days after the inpatient hospital discharge, with no gaps in enrollment.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of acute inpatient hospital discharges that resulted in an ambulatory care follow-up visit within 30 days of discharge from the inpatient hospital stay.</td>
<td>Of the total reported in A, the number of acute inpatient hospital discharges that resulted in an ambulatory care follow-up visit within 30 days of discharge from the inpatient hospital stay.</td>
<td>Field Type: Numeric</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.
C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
   • The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored. CMS and the state will:
   • Evaluate the percentage of acute inpatient hospital discharges that resulted in an ambulatory care follow-up visit within 30 days of the discharge from the inpatient hospital stay.
     o Percentage = (B / A) * 100
   • Use enrollment data to evaluate the total number of acute inpatient hospital discharges per 10,000 member months during the reporting period.
     o Rate = (A / Total Member Months) * 10,000

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

Data Element A
   • The FIDA-IDD Plan should include all acute inpatient hospital discharges for Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of the end of the reporting period.
   • The denominator for this measure is based on acute inpatient hospital discharges, not Participants.
   • To identify all acute inpatient hospital discharges during the reporting period:
     o Identify all acute and nonacute inpatient stays (Inpatient Stay value set).
     o Exclude nonacute inpatient stays (Nonacute Inpatient Stay value set).
     o Identify the discharge date for the stay. The date of discharge must be within the reporting period.
     o Report on all inpatient stays identified with discharges within the reporting period, including denied and pended claims.
   Additionally, the FIDA-IDD Plan should use UB Type of Bill codes 11x, 12x, 41x, and 84x or any acute inpatient facility code to identify discharges from an inpatient hospital stay.
   • If the discharge is followed by readmission or direct transfer to an acute inpatient care setting within the 30-day follow-up period, count only the last discharge for reporting in data element A. To identify readmissions and direct transfers to an acute inpatient care setting:
     o Identify all acute and nonacute inpatient stays (Inpatient Stay value set).
     o Exclude nonacute inpatient stays (Nonacute Inpatient Stay value set).
     o Identify the admission date for the stay.
Data Element A Exclusions

- Exclude discharges for Participants who use hospice services or elect to use a hospice benefit at any time between the hospital discharge date and 30 days following the hospital discharge. These Participants may be identified using various methods, which may include but are not limited to enrollment data, medical record, claims/encounter data (Hospice Encounter value set; Hospice Intervention value set), or supplemental data.

- Exclude discharges due to death, using the Discharges due to Death value set.

- Exclude from data element A any discharges followed by readmission or direct transfer to a nonacute inpatient care setting within the 30-day follow-up period. To identify readmissions and direct transfers to a nonacute inpatient care setting:
  o Identify all acute and nonacute inpatient stays (Inpatient Stay value set).
  o Confirm the stay was for nonacute care based on the presence of a nonacute code (Nonacute Inpatient Stay value set) on the claim.
  o Identify the admission date for the stay.

  These discharges are excluded from the measure because rehospitalization or direct transfer may prevent an outpatient follow-up visit from taking place.

- For example, the following direct transfers/readmissions should be excluded from this measure:
  o An inpatient discharge on June 1, followed by an admission to another inpatient setting on June 1 (a direct transfer).
  o An inpatient discharge on June 1, followed by a readmission to a hospital on June 15 (readmission within 30 days).

Data Element B

- The date of discharge must occur within the reporting period, but the follow-up visit may not be in the same reporting period.
  o For example, if a discharge occurs during the last month of the reporting period, look to the first month of the following reporting period to identify the follow-up visit.

- A follow-up visit is defined as an ambulatory care follow-up visit to assess the Participant’s health following a hospitalization. Codes to identify follow-up visits are provided in the Ambulatory Visits value set, Other Ambulatory Visits value set, and Telephone Visits value set.

- The FIDA-IDD Plan should report ambulatory care follow-up visits based on all visits identified, including denied and pended claims, and including encounter data as necessary in cases where follow-up care is included as part of a bundled payment covering the services delivered during the inpatient stay. The FIDA-IDD Plan should use all information available, including encounter data supplied by providers, to ensure complete and accurate reporting.
F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.

IDD1.4 Participants with a Comprehensive Service Planning Assessment completed within 30 days of enrollment.

<table>
<thead>
<tr>
<th>CONTINUOUS REPORTING</th>
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<tbody>
<tr>
<td><strong>Reporting Section</strong></td>
</tr>
<tr>
<td>FIDA-IDD1. Care Coordination</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants whose 30th day of enrollment occurred within the reporting period.</td>
<td>Total number of Participants whose 30th day of enrollment occurred within the reporting period.</td>
<td>Field type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of Participants who were documented as unwilling to participate in the Comprehensive Service Planning Assessment (CSPA) within 30 days of enrollment.</td>
<td>Of the total reported in A, the number of Participants who were documented as unwilling to participate in the CSPA within 30 days of enrollment.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Note: Is a subset of A.</td>
<td></td>
</tr>
<tr>
<td>Element Letter</td>
<td>Element Name</td>
<td>Definition</td>
<td>Allowable Values</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>C.</td>
<td>Total number of Participants the FIDA-IDD Plan was unable to reach, following three documented outreach attempts, within 30 days of enrollment.</td>
<td>Of the total reported in A, the number of Participants the FIDA-IDD Plan was unable to reach, following three documented outreach attempts, within 30 days of enrollment.</td>
<td>Field type: Numeric Note: Is a subset of A.</td>
</tr>
<tr>
<td>D.</td>
<td>Total number of Participants with a CSPA completed within 30 days of enrollment.</td>
<td>Of the total reported in A, the number of Participants with a CSPA completed within 30 days of enrollment.</td>
<td>Field type: Numeric Note: Is a subset of A.</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.
- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
- The FIDA-IDD Plan should validate that data elements B, C, and D are less than or equal to data element A.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored. CMS and the state will evaluate the percentage of Participants who:
- Were documented as unwilling to participate in the CSPA within 30 days of enrollment.
  - Percentage = (B / A) * 100
- Were unable to be reached, following three documented outreach attempts, to complete the CSPA within 30 days of enrollment.
  - Percentage = (C / A) * 100
- Had a CSPA completed within 30 days of enrollment.
  - Percentage = (D / A) * 100
- Were willing to participate and who could be reached who had a CSPA completed within 30 days of enrollment.
  - Percentage = (D / (A – B – C)) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.
Data Element A

- The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of the end of the reporting period (i.e., include all Participants regardless of whether they are currently enrolled or disenrolled as of the last day of the reporting period).
- The 30th day of enrollment should be based on each Participant’s effective date of enrollment. For the purposes of reporting this measure, 30 days of enrollment will be equivalent to one full calendar month.

Data Element B

- For data element B, the FIDA-IDD Plan should report the number of Participants who were documented as unwilling to participate in the CSPA if a Participant (or his or her authorized representative):
  - Affirmatively declines to participate in the CSPA, affirmatively declines care management activities overall, or refuses any contact with the FIDA-IDD Plan. The Participant may communicate the declination or refusal by phone, mail, fax, or in person. The declination or refusal must be documented by the FIDA-IDD Plan.
  - Expresses willingness to complete the CSPA but asks for it to be conducted after 30 days (despite being offered a reasonable opportunity to complete the CSPA within 30 days). Discussions with the Participant must be documented by the FIDA-IDD Plan.
  - Schedules an appointment to complete the CSPA, but cancels or is a no-show and then is subsequently non-responsive to additional outreach attempts by the FIDA-IDD Plan. All attempts to contact the Participant must be documented by the FIDA-IDD Plan.
  - Initially agrees to complete the CSPA, but then declines to answer a sufficient number of questions in the CSPA, as determined by the FIDA-IDD Plan. The declination must be documented by the FIDA-IDD Plan.
- If a Participant was not reached after three outreach attempts, but then subsequently is reached and refuses the CSPA within 30 days of enrollment, the Participant should be classified in data element B.

Data Element C

- For data element C, the FIDA-IDD Plan should report the number of Participants the FIDA-IDD Plan was unable to reach after three attempts to contact the Participant. The FIDA-IDD Plan should refer to the FIDA-IDD IDT Policy for any specific requirements pertaining to the method of outreach to Participants.
- The FIDA-IDD Plan must document each attempt to reach the Participant, including the method of the attempt (e.g., phone, mail, or email), as CMS and the state may validate this number. If less than three outreach attempts are made to the Participant within 30 days of enrollment, the Participant should not be included in data element C.
There may be instances when the FIDA-IDD Plan has a high degree of confidence that a Participant’s contact information is correct, yet that Participant is not responsive to the FIDA-IDD Plan’s outreach efforts. So long as the FIDA-IDD Plan follows the guidance regarding outreach attempts, these Participants may be included in the count for data element C.

**Data Element D**

- If a Participant’s CSPA is in progress, but is not completed within 30 days of enrollment, then the CSPA should not be considered completed, and therefore, would not be counted in data element D. However, this Participant would be included in data element A.
- If a Participant initially refused the CSPA or could not be reached after three outreach attempts, but then subsequently completes the CSPA within 30 days of enrollment, the Participant should be classified in data element D.

**General Guidance**

- The FIDA-IDD Plan should refer to the FIDA-IDD IDT Policy for specific requirements pertaining to the CSPA.
- Participants reported in data elements B, C, and D must also be reported in data element A, since these data elements are subsets of data element A. Additionally, data elements B, C, and D should be mutually exclusive (e.g., a Participant reported in element B or C should not also be reported in element D).
- There may be certain circumstances that make it impossible or inappropriate to complete a CSPA within the required timeframe.
  - For example, a Participant may be medically unable to respond and have no authorized representative to do so on their behalf, or a Participant may be experiencing an acute medical or behavioral health crisis that requires immediate attention and outweighs the need for a CSPA. However, the FIDA-IDD Plan should not include such Participants in the counts for data elements B or C.

**F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.**

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: [https://Financial-Alignment-Initiative.NORC.org](https://Financial-Alignment-Initiative.NORC.org).
Section FIDA-IDD II. Long Term Care Quality

IDD2.1 Long Term Care Overall Balance.¹

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD2. Long Term Care Quality</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year</td>
<td>By the end of the sixth month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants continuously enrolled in the FIDA-IDD Plan for 6 months.</td>
<td>Total number of Participants continuously enrolled in the FIDA-IDD Plan for 6 months during the reporting period, with no gaps in enrollment.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of Participants who did not reside in an Intermediate Care Facility (ICF-IID).</td>
<td>Of the total reported in A, the number of Participants who did not reside in an ICF-IID at the earliest point of their enrollment during the reporting period.</td>
<td>Field Type: Numeric Note: Is a subset of A.</td>
</tr>
<tr>
<td>C.</td>
<td>Total number of Participants who did not reside in an ICF-IID during the reporting period.</td>
<td>Of the total reported in B, the number of Participants who did not reside in an ICF-IID during the reporting period.</td>
<td>Field Type: Numeric Note: Is a subset of B.</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

IDD-26
• A higher number of Participants who did not reside in an ICF-IID during the reporting period (data element C) is better.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
  • The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.
  • The FIDA-IDD Plan should validate that data element C is less than or equal to data element B.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.
  • For Participants who did not reside in an ICF-IID at the earliest point of their enrollment during the reporting period, CMS and the state will evaluate the percentage of Participants who did not reside in an ICF-IID during the reporting period.
    o Percentage = (C / B) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

Definitions
  • ICF-IID services are those services provided by a residential facility certified by the Office of People with Developmental Disabilities (OPWDD) as an ICF-IID and providing comprehensive and individualized health care and habilitation services to individuals with IDD to promote their functional status and independence. ICF-IID is available only for individuals in need of, and receiving, active treatment (AT) services.
  • AT refers to aggressive, consistent implementation of a program of specialized and generic training, treatment, and health services.

Data Element A
  • The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of the end of the reporting period (i.e., include all Participants regardless of whether they are currently enrolled or disenrolled as of the last day of the reporting period).

Data Element B
  • For data element B, the FIDA-IDD Plan should include Participants who did not reside in an ICF-IID at the earliest point of their enrollment during the current reporting period:
    o For Participants enrolled as of January 1 of the reporting period, use the Participants’ status (i.e., did not reside in an ICF-IID) as of January 1 of current reporting period.

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For Participants enrolled after January 1 of the reporting period, use the Participants’ status (i.e., did not reside in an ICF-IID) on the first day of enrollment during the reporting period.

- For example, if a Participant enrolls on April 1 and does not reside in an ICF-IID at the time of enrollment, the Participant would be reported in data element B.

Data Element C

- To establish a Participant’s ICF-IID stay for data element C, the FIDA-IDD Plan should evaluate the entire reporting period in which the Participant was enrolled to determine if the Participant did not reside in an ICF-IID.

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.

IDD2.2 Community Reintegration.

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD2. Long Term Care Quality</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year, beginning in CY2</td>
<td>By the end of the second month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants who resided in an Intermediate Care Facility (ICF-IID) during the previous reporting period.</td>
<td>Total number of Participants who resided in an ICF-IID during the previous reporting period and who were continuously enrolled during the previous and current reporting periods.</td>
<td>Field Type: Numeric</td>
</tr>
</tbody>
</table>
B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.
  - CMS and the state will perform an outlier analysis.
  - As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
  - The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.
  - CMS and the state will evaluate the percentage of Participants who resided in an ICF-IID during the previous reporting period who were continuously enrolled during the previous and current reporting periods who were discharged to a community-based setting during the previous or current reporting period who did not return to the ICF-IID during the current reporting period.
    - Percentage = (B / A) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.
  **Definitions**
  - **ICF-IID services** are those services provided by a residential facility certified by OPWDD as an ICF-IID and providing comprehensive and individualized health care and habilitation services to individuals with IDD to promote their functional status and independence. ICF-IID is available only for individuals in need of, and receiving, active treatment (AT) services.
• AT refers to aggressive, consistent implementation of a program of specialized and generic training, treatment, and health services.
• A community-based setting is defined as a private home, apartment, Individual Residential Alternative, or Family Care Home.

Data Element A

• Continuous enrollment is defined as no more than one gap in enrollment of up to 45 days during the reporting period (i.e., January through December). To determine continuous enrollment for a Participant for whom enrollment is verified monthly, the Participant may not have more than a 1-month gap in coverage (i.e., a Participant whose coverage lapses for 2 months [60 days] is not considered continuously enrolled).
• For the purposes of this measure, the “previous reporting period” is defined as the previous calendar year. The “current reporting period” is defined as the current calendar year. For example, for data submitted on February 28, 2022, the previous reporting period is January 1, 2020 – December 31, 2020, and the current reporting period is January 1, 2021 – December 31, 2021.

Data Element B

• The discharge to a community-based setting could have occurred during either the previous reporting period or the current reporting period.
• Codes to identify a discharge to a community-based setting are provided in the Discharges to the Community value set.

General Guidance

• This measure is reported starting with the FIDA-IDD Plan’s second year of operation (i.e., Calendar Year 2). Calendar Year 2017 was Calendar Year 2 for the FIDA-IDD Plan.

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

• The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.
Intermediate Care Facilities for Individuals with Intellectual Disabilities (ICF-IID) Diversion.ii

CONTINUOUS REPORTING

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD2. Long Term Care Quality</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year, beginning CY2</td>
<td>By the end of the second month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Total number of Participants who were continuously enrolled in the FIDA-IDD Plan for at least 5 out of the last 6 months during the previous reporting period and continuously enrolled in the FIDA-IDD Plan for at least 11 out of 12 months during the current reporting period.</td>
<td>Total number of Participants who were continuously enrolled in the FIDA-IDD Plan for at least 5 out of the last 6 months during the previous reporting period and continuously enrolled in the FIDA-IDD Plan for at least 11 out of 12 months during the current reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B</td>
<td>The total number of Participants who did not reside in an ICF-IID during the previous reporting period.</td>
<td>Of the total reported in A, the number of Participants who did not reside in an ICF-IID during the previous reporting period.</td>
<td>Field Type: Numeric Note: Is a subset of A.</td>
</tr>
<tr>
<td>C</td>
<td>Total number of Participants who did not reside in an ICF-IID during the current reporting period.</td>
<td>Of the total reported in B, the number of Participants who did not reside in an ICF-IID during the current reporting period.</td>
<td>Field Type: Numeric Note: Is a subset of B.</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.
• The quality withhold benchmark for DY 2 through 7 is timely and accurate reporting according to the measure specifications. For more information, refer to the Quality Withhold Technical Notes (DY 2-7): New York FIDA-IDD-Specific Measures.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.

• The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.
• The FIDA-IDD Plan should validate that data element C is less than or equal to data element B.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.

• For Participants who did not reside in an ICF-IID during the previous reporting period, CMS and the state will evaluate the percentage of Participants who did not reside in an ICF-IID during the current reporting period.
  o Percentage = (C / B) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

**Definitions**

• ICF-IID services are those services provided by a residential facility certified by OPWDD as an ICF-IID and providing comprehensive and individualized health care and habilitation services to individuals with IDD to promote their functional status and independence. ICF-IID is available only for individuals in need of, and receiving, active treatment (AT) services.

• AT refers to aggressive, consistent implementation of a program of specialized and generic training, treatment, and health services.

**Data Element A**

• Continuous enrollment is defined as no more than one gap in enrollment of up to 45 days during each reporting period (i.e., January through December). To determine continuous enrollment for a Participant for whom enrollment is verified monthly, the Participant may not have more than a 1-month gap in coverage (i.e., a Participant whose coverage lapses for 2 months [60 days] is not considered continuously enrolled).

• The Participant must be enrolled as of the last day of both the previous and current reporting periods to be included in this measure.

• For the purposes of this measure, the “previous reporting period” is defined as the previous calendar year. The “current reporting period” is defined as the current calendar year. For example, for data submitted on February 28, 2022, the previous reporting period is January 1, 2020 – December 31, 2020, and the current reporting period is January 1, 2021 – December 31, 2021.
Data Element A Exclusions

- Exclude Participants who are transitioned to hospice services in either the current or previous reporting periods when reporting this measure. These Participants may be identified using various methods, which may include but are not limited to enrollment data, medical record, claims/encounter data (Hospice Encounter value set; Hospice Intervention value set), or supplemental data.
- Exclude Participants who expired in either the current or previous reporting period when reporting this measure using the Discharges due to Death value set.

General Guidance

- This measure is reported starting with the FIDA-IDD Plan’s second year of operation (i.e., Calendar Year 2). Calendar Year 2017 was Calendar Year 2 for the FIDA-IDD Plan.

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.
Section FIDA-IDD III. Enrollee Protections

IDD3.1 The number of critical incident and abuse reports for Participants receiving LTSS.

<table>
<thead>
<tr>
<th>IMPLEMENTATION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reporting</strong></td>
</tr>
<tr>
<td><strong>Section</strong></td>
</tr>
<tr>
<td>FIDA-IDD3. Enrollee Protections</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ONGOING</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reporting</strong></td>
</tr>
<tr>
<td><strong>Section</strong></td>
</tr>
<tr>
<td>FIDA-IDD3. Enrollee Protections</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants receiving LTSS.</td>
<td>Total number of Participants receiving LTSS during the reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of critical incident and abuse reports.</td>
<td>Of the total reported in A, the number of critical incident and abuse reports during the reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
- N/A.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored. CMS and the state will evaluate the:
- Number of critical incident and abuse reports per 1,000 Participants receiving LTSS during the current reporting period.
  - Rate = (B / A) * 1,000
- Average number of critical incident and abuse reports for Participants receiving LTSS during the prior four reporting periods (i.e., rolling year).
  - Average number = Sum of B for prior four reporting periods / 4
- Weighted average number of critical incident and abuse reports per 1,000 Participants receiving LTSS during the prior four reporting periods.
  - Rate = (Sum of B for prior four reporting periods / Sum of A for prior four reporting periods) * 1,000

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

Definitions
- **Critical incident** refers to any actual or alleged event or situation that creates a significant risk of substantial or serious harm to the physical or mental health, safety or well-being of a Participant including, neglect, financial exploitation, and mandated Reporting Requirements called for under the three-way contract, and under New York State and Federal requirements for reporting on incidents and residents of Intermediate Care Facilities for Individuals with Intellectual Disabilities (ICF-IID) or Nursing Facilities.
- **Abuse** refers to any of the following:
  - Willful use of offensive, abusive, or demeaning language by a caretaker that causes mental anguish;
  - Knowing, reckless, or intentional acts or failures to act which cause injury or death to an individual or which place that individual at risk of injury or death;
  - Rape or sexual assault;
  - Corporal punishment or striking of an individual;
  - Unauthorized use or the use of excessive force in the placement of bodily restraints on an individual; and
  - Use of bodily or chemical restraints on an individual which is not in compliance with federal or state laws and administrative regulations.

Data Element A
- The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of
the end of the reporting period (i.e., include all Participants regardless of whether they are currently enrolled or disenrolled as of the last day of the reporting period).

- Data element A should include all Participants enrolled in the FIDA-IDD Plan who received LTSS for any amount of time during the reporting period.

**Data Element B**

- For data element B, the FIDA-IDD Plan should include all new critical incident and abuse cases that are reported during the reporting period, regardless of whether the case status is open or closed as of the last day of the reporting period.
- Critical incident and abuse reports could be reported by the FIDA-IDD Plan or any provider and are not limited to only those providers defined as LTSS providers.
- It is possible for Participants to have more than one critical incident and/or abuse report during the reporting period. All critical incident and abuse reports during the reporting period should be counted.

**F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.**

- The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.
Section FIDA-IDD IV. Utilization

IDD4.1 Participants self-directing their services through employer authority or budget authority.

### CONTINUOUS REPORTING

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD4. Utilization</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year</td>
<td>By the end of the second month following the last day of the reporting period</td>
</tr>
</tbody>
</table>

A. Data Element Definitions – details for each data element reported to CMS and the state, including examples, calculation methods, and how various data elements are associated.

<table>
<thead>
<tr>
<th>Element Letter</th>
<th>Element Name</th>
<th>Definition</th>
<th>Allowable Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Total number of Participants enrolled in the FIDA-IDD Plan.</td>
<td>Total number of Participants continuously enrolled in the FIDA-IDD Plan during the reporting period.</td>
<td>Field Type: Numeric</td>
</tr>
<tr>
<td>B.</td>
<td>Total number of Participants who were self-directing their services through employer authority or budget authority.</td>
<td>Of the total reported in A, the number of Participants who were self-directing their services through employer authority or budget authority during the reporting period.</td>
<td>Field Type: Numeric Note: Is a subset of A.</td>
</tr>
</tbody>
</table>

B. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- CMS and the state will perform an outlier analysis.
- As data are received from the FIDA-IDD Plan over time, CMS and the state will apply threshold checks.

C. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.
• The FIDA-IDD Plan should validate that data element B is less than or equal to data element A.

D. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.

• CMS and the state will evaluate the percentage of Participants continuously enrolled in the FIDA-IDD Plan who were self-directing their services through employer authority or budget authority during the reporting period.
  o Percentage = (B / A) * 100

E. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

Definition

• Self-direction is defined as the ability for a Participant and his/her Representative to direct his/her own services through the Self-Direction in the Section 1915(c) OPWDD Comprehensive Waiver or the consumer-directed personal assistance option.

Data Element A

• The FIDA-IDD Plan should include all Participants who meet the criteria outlined in data element A, regardless of whether they are disenrolled as of the end of the reporting period (i.e., include all Participants regardless of whether they are currently enrolled or disenrolled as of the last day of the reporting period).

• Continuous enrollment is defined as no more than one gap in enrollment of up to 45 days during the reporting period (i.e., January through December). To determine continuous enrollment for a Participant for whom enrollment is verified monthly, the Participant may not have more than a 1-month gap in coverage (i.e., a Participant whose coverage lapses for 2 months [60 days] is not considered continuously enrolled).

F. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.

• The FIDA-IDD Plan will submit data collected for this measure in the above specified format through a secure data collection site established by CMS. This site can be accessed at the following web address: https://Financial-Alignment-Initiative.NORC.org.

IDD4.2 Participants who received an eye exam in the last year. – Retired
IDD4.3 Annual Dental Visit.

Please note: No FIDA-IDD Plan reporting is required for this measure; however, the FIDA-IDD Plan must assist NYSDOH with data collection and analysis as needed. NYSDOH will calculate the measure using claims and encounter data.

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD4. Utilization</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year</td>
<td>N/A</td>
</tr>
</tbody>
</table>

A. QA Checks/Thresholds – procedures used by CMS and the state to establish benchmarks in order to identify outliers or data that are potentially erroneous.

- The quality withhold benchmark is 79% for DY 3 and 82% for DY 4 through 7. For more information, refer to the Quality Withhold Technical Notes (DY 2-7): New York FIDA-IDD-Specific Measures.

B. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.

- CMS and the state will evaluate the percent of Participants continuously enrolled in the FIDA-IDD Plan during the measurement year (with no more than a one-month enrollment gap) who had at least one dental procedure code during the measurement year.
Section FIDA-IDD V. Participant-Level File

The New York Office of Quality and Patient Safety (OQPS) will be evaluating measures using the Medicaid Encounter Data System (MEDS), the OPWDD Approved Assessment Tool (OAA), and Participant-level data. The State will conduct ongoing research on Participants’ personal experiences in FIDA-IDD and on potential relationship of these experiences with Participants’ health outcomes.

The following table provides instructions on the submission of Participant-level data on a subset of measures the FIDA-IDD Plan is reporting at the plan-level and a number of IDT-related performance measures. This table does not provide any additional measures; it only provides guidance on the subset of measures for which CMS/NYSDOH require Participant-level data. Please see the source of each measure for the technical specifications for each. In particular:

- Columns 17-51 provide instructions on reporting Participant-level data on HEDIS measures.
- Columns 52-61 provide instructions on reporting some of the NY FIDA-IDD-Specific measures defined elsewhere within this appendix and some of the MMP Core Measures defined in the Core Reporting Requirements: https://www.cms.gov/Medicare-Medicaid-Coordination/Medicare-and-Medicaid-Coordination/Medicare-Medicaid-Coordination-Office/FinancialAlignmentInitiative/MMPIInformationandGuidance/MMPRreportingRequirements.html.
- Columns 65-360 provide instructions on reporting Participant-level data on IDT activities. As noted for each element, some of the IDT requirements are specified within this appendix while others are specified within the final IDT Policy. CMS and NYSDOH may establish thresholds and conduct outlier analysis based on this data.

Questions from the FIDA-IDD Plan regarding the Participant-Level File or the data submission process should be directed to NYSDOH and OPWDD.

### IDD5.1 Participant-Level File

<table>
<thead>
<tr>
<th>Reporting Section</th>
<th>Reporting Frequency</th>
<th>Level</th>
<th>Reporting Period</th>
<th>Due Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIDA-IDD Participant-Level Measures</td>
<td>Annually</td>
<td>Contract</td>
<td>Calendar Year</td>
<td>By the end of the sixth month following the last day of the reporting period</td>
</tr>
</tbody>
</table>
A. File Format Definitions – details for each data element reported to CMS and the state.

<table>
<thead>
<tr>
<th>Column Placement</th>
<th>Name</th>
<th>Direction</th>
<th>Allowed Values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Instructions for Participant Identification</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Column 1-8</td>
<td>MMIS ID</td>
<td>Enter the Plan’s numeric eight-digit ID.</td>
<td>#######</td>
</tr>
<tr>
<td>Column 9–16</td>
<td>Medicaid CIN</td>
<td>A Participant’s client identification number. The field should be continuous without any spaces or hyphens. The field is alpha-numeric and should be treated as a text field. The CIN entered in this field should be for the CIN for the measurement period. For example, CINs for 2015 should be used.</td>
<td>Dual eligible individuals only</td>
</tr>
<tr>
<td><strong>Instructions for Participant-level data on HEDIS Measures</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Column 17</td>
<td>Denominator for Antidepressant Medication Management (AMM)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Antidepressant Medication Management measures, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td></td>
<td>Numerator for Antidepressant Medication Management – Effective Acute Phase Treatment (AMM)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Antidepressant Medication Management – Effective Acute Phase Treatment measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td></td>
<td>Numerator for Antidepressant Medication Management–Effective Continuation Phase Treatment (AMM)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Antidepressant Medication Management – Effective Continuation Phase Treatment measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 20-21</td>
<td>Denominator for Pharmacotherapy Management of COPD Exacerbation (PCE)</td>
<td>Enter the number of times the Participant is in the denominator of the Pharmacotherapy Management of COPD PCE measure, ‘0’ if the Participant is not in the denominator.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Column 22-23</td>
<td>Numerator for Pharmacotherapy Management of COPD Exacerbation (PCE): Systemic Corticosteroid</td>
<td>Enter the number of times the Participant is in the numerator of the Pharmacotherapy Management of COPD PCE Corticosteroid measure, ‘0’ if the Participant is not in the numerator.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column 24-25</td>
<td>Numerator for Pharmacotherapy Management of COPD Exacerbation (PCE): Bronchodilator</td>
<td>Enter the number of times the Participant is in the numerator of the Pharmacotherapy Management of COPD PCE Bronchodilator measure, ‘0’ if the Participant is not in the numerator.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column 26-27</td>
<td>Denominator for Follow-Up After Hospitalization for Mental Illness (FUH)</td>
<td>Enter the number of times the Participant appears in the denominator of the Follow-Up After Hospitalization for Mental Illness; ‘0’ if the Participant is not in the denominator.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column 28-29</td>
<td>Numerator 1 for Follow-Up After Hospitalization for Mental Illness, 7 days after discharge (FUH)</td>
<td>Enter the number of times the Participant appears in numerator 1 of the Follow-Up After Hospitalization for Mental Illness, 7 days after discharge. ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column 30-31</td>
<td>Numerator 2 for Follow-Up After Hospitalization for Mental Illness, 30 days after discharge (FUH)</td>
<td>Enter the number of times the Participant appears in numerator 2 of the Follow-Up After Hospitalization for Mental Illness, 30 days after discharge. ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>00-98</td>
</tr>
<tr>
<td>Column 32</td>
<td>Denominator for Osteoporosis Management in Women who had a Fracture (OMW)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Osteoporosis Management OMW measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes, 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>Column 33</td>
<td>Numerator for Osteoporosis Management in Women who had a Fracture (OMW)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Osteoporosis Management OMW measure, ‘0’ if the Participant is not in the numerator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 34</td>
<td>Denominator for the Care for Older Adults (COA)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Care for Older Adults measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 35</td>
<td>Numerator for Care for Older Adults – Medication Review (COA)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Care for Older Adults Medication Review measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 36</td>
<td>Numerator for Care for Older Adults – Functional Status Assessment (COA)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Care for Older Adults Functional Status Assessment, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 37</td>
<td>Numerator for Care for Older Adults – Pain Screening (COA)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Care for Older Adults Pain Screening, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 38</td>
<td>Denominator for the Use of High-Risk Medications in the Elderly (DAE)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Use of High-Risk Medications DAE measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 39</td>
<td>Numerator for the Use of High-Risk Medications in the Elderly (DAE)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Use of High-Risk Medications DAE measure, ‘0’ if the Participant is not in the numerator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 40</td>
<td>Denominator for Persistence of Beta-Blocker Treatment After a Heart Attack (PBH)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Persistence of Beta-Blocker PBH measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------</td>
</tr>
</tbody>
</table>
| Column 41        | Numerator for Persistence of Beta-Blocker Treatment After a Heart Attack (PBH) | Enter a ‘1’ if the Participant is in the numerator of the Persistence of Beta-Blocker PBH measure, ‘0’ if the Participant is not in the numerator.                                                             | 1 = Yes  
0 = No |
| Column 42        | Denominator for Potentially Harmful Drug-Disease Interactions in the Elderly (DDE) | Enter a ‘1’ if the Participant is in the denominator of the Potentially Harmful Drug-Disease DDE measure, ‘0’ if the Participant is not in the denominator.                                                   | 1 = Yes  
0 = No |
| Column 43        | Numerator for Potentially Harmful Drug-Disease Interactions in the Elderly (DDE) Rate 1 | Enter a ‘1’ if the Participant is in the numerator of the Potentially Harmful Drug-Disease DDE measure Rate 1, ‘0’ if the Participant is not in the numerator.                                      | 1 = Yes  
0 = No |
| Column 44        | Numerator for Potentially Harmful Drug-Disease Interactions in the Elderly (DDE) Rate 2 | Enter a ‘1’ if the Participant is in the numerator of the Potentially Harmful Drug-Disease DDE measure Rate 2, ‘0’ if the Participant is not in the numerator.                                      | 1 = Yes  
0 = No |
| Column 45        | Numerator for Potentially Harmful Drug-Disease Interactions in the Elderly (DDE) Rate 3 | Enter a ‘1’ if the Participant is in the numerator of the Potentially Harmful Drug-Disease DDE measure Rate 3, ‘0’ if the Participant is not in the numerator.                                      | 1 = Yes  
0 = No |
| Column 46        | Denominator for Controlling High Blood Pressure (CBP)                 | Enter a ‘1’ if the Participant is in the denominator of the Controlling High Blood Pressure measure, ‘0’ if the Participant is not in the denominator.                                                       | 1 = Yes  
0 = No |
<table>
<thead>
<tr>
<th>Column Placement</th>
<th>Name</th>
<th>Direction</th>
<th>Allowed Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Column 47</td>
<td>Numerator for Controlling High Blood Pressure (CBP)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Controlling High Blood Pressure measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 48</td>
<td>Denominator for Spirometry Testing (SPR)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Use of Spirometry Testing in the Assessment and Dx of COPD measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 49</td>
<td>Numerator for Spirometry Testing (SPR)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Use of Spirometry Testing in the Assessment and Dx of COPD measure. ‘0’ if the Participant is not in the numerator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 50</td>
<td>Denominator for Colorectal Cancer Screening (COL)</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the Colorectal Cancer Screening measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 51</td>
<td>Numerator for Colorectal Cancer Screening (COL)</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the Colorectal Cancer Screening measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
</tbody>
</table>

**Instructions for Participant Level Data on NY FIDA-IDD-Specific Measures**

<table>
<thead>
<tr>
<th>Column Placement</th>
<th>Name</th>
<th>Direction</th>
<th>Allowed Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Column 52</td>
<td>Denominator for IDD2.1 Long Term Care Overall Balance</td>
<td>Enter a ‘1’ if the Participant is in the denominator of IDD2.1 Long Term Care Overall Balance measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 53</td>
<td>Numerator for IDD2.1 Long Term Care Overall Balance</td>
<td>Enter a ‘1’ if the Participant is in the numerator of IDD2.1 Long Term Care Overall Balance measure, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
</tr>
<tr>
<td>------------------</td>
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<td>----------------</td>
</tr>
<tr>
<td>Column 54</td>
<td>Denominator for IDD4.1 Participants self-directing their services through employer authority or budget authority</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the IDD4.1 Participants self-directing their services through employer authority or budget authority, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 55</td>
<td>Numerator for IDD4.1 Participants self-directing their services through employer authority or budget authority</td>
<td>Enter a ‘1’ if the Participant is in the numerator of IDD4.1 Participants self-directing their services through employer authority or budget authority, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 56</td>
<td>Denominator for IDD2.2 Community Reintegration</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the IDD2.2 Community Reintegration measure above, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 57</td>
<td>Numerator for IDD2.2 Community Reintegration</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the IDD2.2 Community Reintegration measure above, ‘0’ if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 58</td>
<td>Denominator for IDD2.3 ICF-IID Diversion</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the IDD2.3 ICF-IID Diversion measure above, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 59</td>
<td>Numerator for IDD2.3 ICF-IID Diversion</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the IDD2.3 ICF-IID Diversion measure above, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
</tr>
<tr>
<td>------------------</td>
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<td>----------------</td>
</tr>
<tr>
<td><strong>Instructions for Participant Level Data on MMP Specific Core Reporting Requirements</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Column 60</td>
<td>Denominator for MMP Core Measure 9.2 Nursing Facility Diversion</td>
<td>Enter a ‘1’ if the Participant is in the denominator of the MMP Core Measure 9.2 Nursing Facility Diversion measure, ‘0’ if the Participant is not in the denominator.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 61</td>
<td>Numerator for MMP Core Measure 9.2 Nursing Facility Diversion</td>
<td>Enter a ‘1’ if the Participant is in the numerator of the MMP Core Measure 9.2 Nursing Facility Diversion measure, '0' if the Participant is not in the numerator or the information is missing.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 62-64</td>
<td>Retain file layout</td>
<td>Enter 000, to maintain the same layout to the file after column 64 as last year.</td>
<td>000</td>
</tr>
<tr>
<td><strong>Instructions for Participant Level Data on IDT Requirements</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Column 65</td>
<td>Did the Participant refuse to have his/her OAA completed at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant refused to have his/her OAA completed at least once in the reporting year, otherwise enter ‘0’.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Columns 66-161 (up to 12 dates for this measure)</td>
<td>Date(s) on which the Participant had an LP completed or updated</td>
<td>Enter the date(s) on which the Participant had their LP completed or updated as specified in the final IDT Policy. Leave the cell blank if the Participant did not have an LP completed/updated or the information is missing.</td>
<td>MMDDYYYY</td>
</tr>
<tr>
<td>Column 162</td>
<td>Did the Participant refuse to have his/her LP completed at least once in the reporting year?</td>
<td>Enter a ‘1’ if this Participant refused to have his/her LP completed at least once in the reporting year as specified in the final IDT Policy, otherwise enter ‘0’.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
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<td>----------------</td>
</tr>
<tr>
<td>Columns 163-258</td>
<td>Date(s) on which the Participant had an IDT meeting within reporting year</td>
<td>Enter the date(s) on which the Participant had IDT meetings as specified in the final IDT Policy. Leave the cell blank if the Participant did not have any IDT meetings or the information is missing.</td>
<td>MMDDYYYY</td>
</tr>
<tr>
<td>Column 259</td>
<td>Did the Participant refuse to have an IDT meeting at least once in the reporting year?</td>
<td>Enter a ‘1’ if this Participant refused to have an IDT meeting as specified in the final IDT Policy at least once in the reporting year, otherwise enter ‘0’.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Columns 260-355</td>
<td>Date(s) on which the Participant was discharged from a hospital inpatient or ICF–IID to the Community</td>
<td>Enter the date(s) on which the Participant was discharged from a hospital inpatient or ICF–IID to the community as specified in the final IDT Policy. Leave the cell blank if the Participant was not discharged from a hospital inpatient or ICF–IID to the community or the information is missing.</td>
<td>MMDDYYYY</td>
</tr>
<tr>
<td>Column 356</td>
<td>Did the Participant have Behavioral Health Specialist as a member of the IDT at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant had a Behavioral Health Specialist as a member of the IDT as specified in the final IDT Policy, ‘0’ if the Participant did not have a Behavioral Health Specialist as a member of the IDT.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 357</td>
<td>Did the Participant have an RN assessor as a member of the IDT at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant had the RN assessor as a member of the IDT as specified in the final IDT Policy, ‘0’ if the Participant did not have the RN Assessor as a member of the IDT.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column Placement</td>
<td>Name</td>
<td>Direction</td>
<td>Allowed Values</td>
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</tr>
<tr>
<td>Column 358</td>
<td>Did the Participant have a Participant Designee on the IDT at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant had a Participant Designee as a member of the IDT as specified in the final IDT Policy, ‘0’ if the Participant did not have a Participant Designee as a member of the IDT.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 359</td>
<td>Did the Participant have a DD provider or a designee with clinical experience from the DD provider on the IDT at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant had a DD provider or a designee with clinical experience from the DD provider as a member of the IDT as specified in the final IDT Policy, ‘0’ if the Participant did not have a DD provider or a designee with clinical experience from the DD provider as a member of the IDT.</td>
<td>1 = Yes 0 = No</td>
</tr>
<tr>
<td>Column 360</td>
<td>Did the Participant have an ICF–IID representative on the IDT at least once in the reporting year?</td>
<td>Enter a ‘1’ if the Participant had an ICF-IID representative as a member of the IDT as specified in the final IDT Policy, ‘0’ if the Participant did not have an ICF-IID representative as a member of the IDT.</td>
<td>1 = Yes 0 = No</td>
</tr>
</tbody>
</table>

B. Edits and Validation Checks – validation checks that should be performed by the FIDA-IDD Plan prior to data submission.

- The FIDA-IDD Plan should ensure that all data values are recorded in the prescribed format (see column “Allowed Values”).

C. Analysis – how CMS and the state will evaluate reported data, as well as how other data sources may be monitored.

- NYSDOH will use the Participant-level data to conduct ongoing research on Participants’ personal experiences in the FIDA-IDD program and on potential relationship of these experiences with Participants’ health outcomes.

D. Notes – additional clarifications to a reporting section. This section incorporates previously answered frequently asked questions.

Overall Format

- Prepare a fixed width text file in the following format.
  - Include one row for every Participant who was enrolled in the FIDA-IDD Plan and who meets criteria for one or more of the specified FIDA-IDD measures for the measurement year.
Numeric values should be right-justified and blank filled to the left of the value; text fields should be left-justified and blank filled to the right of the value.

The file should be named ParticipantFIDAIDD.txt.

**General Guidance**

- The sum of the field should equal the numerator or denominator for the corresponding measure entered in the NYS submission tools for the plan-level reporting. Measures that are not applicable to the Participant should be zero-filled.

- The FIDA-IDD Plan should use the continuous enrollment specifications applicable to each element as specified in HEDIS, elsewhere in this appendix, and/or the MMP Core Reporting Requirements.

**E. Data Submission – how the FIDA-IDD Plan will submit data collected to CMS and the state.**

- The FIDA-IDD Plan will submit the file in the above specified format to the OQPS MLTC Evaluation mailbox at the NYSDOH via the Health Commerce System (HCS) Secure File Transfer 2.0.