Updated
State Medicaid Health Information Technology Plan (SMHP)
November 3, 2017
Revised March 2018

State of Mississippi

Division of Medicaid
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   November 3, 2017

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1 Executive Summary

The Mississippi Division of Medicaid (DOM) submits this update to the State Medicaid Health Information Technology Plan (SMHP), in accordance with implementation activities authorized by the American Recovery and Reinvestment Act of 2009 (ARRA) (Pub. L. 111-5). This SMHP update provides a description of the strategic planning process that DOM has undertaken, and continues to undertake to participate in the provider incentive program; the business and operational plan for payment of the incentives; and an HIT Roadmap presenting the direction that DOM plans to take to achieve the HIT vision described in this document.

In 2017, DOM completed a new Environmental Scan of the State of Mississippi to evaluate provider adoption of CEHRT and HIT/HIE. Data from the initial 2010 scan and the recent 2017 scan provide a comprehensive understanding of the HIT landscape within the State of Mississippi. The current HIT landscape, is discussed in Section 3 – Current HIT Landscape Assessment – The “As-Is” Environment. The results of the 2017 Environmental Scan serve as a “line in the sand” revamp for revisions to the As-Is Environment, To-Be Landscape, and the HIT Roadmap of this IAPDU.

With a thorough understanding of the current EHR and HIT/HIE landscape, DOM’s planning effort for this update focused on the vision of DOM’s HIT for the next five years, with emphasis on the next two years (2018 and 2019). DOM has specific goals to upgrade and transform the existing Medicaid Management Information System (MMIS) with strategically implemented modular components over the next several years as a part of a new Modularized Medicaid Enterprise System (MMES). As part of the overall DOM HIT program, including interoperability with the new MES, DOM will utilize clinical data to improve care coordination and the quality of care of Medicaid beneficiaries in the State of Mississippi via four goals: 1) Achieve greater interoperability with Medicaid providers and provider clinical systems (EHRs, other clinical systems) to aggregate provider-based Medicaid clinical data and store/utilize this data in the existing DOM Clinical Data Repository; 2) Utilize the aggregated provider-based Medicaid clinical data in the DOM Clinical Data Repository for Agency goals and programs including clinical data analytics and clinical data population health; 3) Offer tools and interfaces to providers so that providers may access and utilize the aggregated clinical data in the DOM Clinical Data Repository, including such tools as a Medicaid clinical data provider portal and real-time, bi-directional clinical data interfaces to support the sharing and updating of Medicaid clinical data interoperability within provider EHRs and provider EHR workflows; and 4) promote adoption of CEHRT for DOM providers with the goal of using CEHRT and HIT/HIE to promote coordinated health care for DOM beneficiaries, better health care outcomes, and improvements in care quality. The effort to promote electronic exchange of clinical data, will be enhanced by the improvement of access to broadband technology for the citizens of Mississippi. Discussion of DOM’s future vision of HIT and HIE can be found in this document at Section 4 – To-Be Landscape.

As one of the key elements to this SMHP, DOM underwent a comprehensive technical, business and operational planning endeavor to be ready to pay Mississippi Medicaid Provider Incentive Payments (MPIP) as quickly as possible. This commitment resulted in Mississippi being one of the first states in the nation to make incentive payments to its providers. DOM has implemented rigorous administration and oversight of the MPIP, including A/I/U post payment audits, and continues to promote the adoption of CEHRT for its providers. As part of its promotion efforts, DOM implemented a communication plan to inform providers of the availability of the incentives and will continue to conduct provider outreach and
education. The discussion of the MPIP and its processes is found in this document in Section 5 – Provider Incentive Program Blueprint.

Using DOM’s strategy as defined by the To-Be Landscape, DOM defined the Mississippi HIT Roadmap for achievement of its future vision. The HIT Roadmap articulates the major milestones and activities that DOM will achieve as it moves from its current environment (As-Is) to its future vision (To-Be). Discussion of DOM’s HIT Roadmap is found in this document in Section 6 – HIT Roadmap.

In addition to the submission of an updated SMHP last year, DOM submitted an updated IAPD to CMS in November 2016 (with a revised submission in February 2017), requesting implementation funding for federal fiscal years (FFY) 2017 and 2018. The updated SMHP and IAPD were each approved February 22, 2017. As requested by CMS, a 2017 SMHP Addendum was submitted in February 2017 regarding the 2015-2017 Modifications Rule.

DOM is pleased to submit this updated SMHP dated November 3, 2017, as documentation of its continued activities to comprehensively plan and implement the future vision of DOM as a partner to its providers and stakeholders in the adoption of CEHRT and the promotion of HIE. An updated IAPD will be submitted in conjunction with this SMHP update to request proposed implementation funding through FFY 2019.
2 Introduction and Overview

DOM submits an updated SMHP annually to provide CMS with a summary of the activities that DOM has completed and expects to undertake in the future to successfully implement its HIT promotion program. For ease of use, an acronym table is attached hereto as Appendix A and a glossary of terms is attached hereto as Appendix B. And, to facilitate CMS review, tables are now provided in Appendix M that cross-reference sections of the SMHP to the CMS Guidelines of April 27, 2010.

In order to submit this FFY 2018 SMHP update, DOM has completed a rigorous planning process designed to consider and incorporate all of the requirements for implementation of its HIT promotion program. These requirements include payment of the incentives for A/I/U and MU of CEHRT for Mississippi Medicaid providers.

The results of DOM’s meticulous planning process are incorporated into this SMHP update, including all of the elements required by CMS. This document includes a description of the following elements required by CMS:

- The current and future vision for the MMIS;
- A re-assessment of the current HIT environment in the State of Mississippi through a 2017 Environmental Scan;
- The State of Mississippi’s HIT To-Be landscape, taking into account perspectives learned from the 2017 Environmental Scan;
- The State of Mississippi’s HIT Roadmap and plan;
- A description of how the SMHP was designed and developed;
- The MPIP payment system and how the MMIS has been considered in developing the HIT Roadmap;
- Infrastructure enhancements that will support the overall goals of DOM;
- Data sharing components of the HIT Roadmap;
- Promotion of secure data exchange in accordance with the Health Insurance Portability and Accountability Act (HIPAA);
- A description of how DOM will promote the adoption and use of data technical standards;
- The process for improvements in health outcomes, clinical quality, or efficiency resulting from the adoption of CEHRT by DOM Medicaid providers, including the methods by which DOM will measure success;
- The method by which DOM will support the integration of clinical and administrative data;
- The method by which DOM will adopt national data standards for health and data exchange and open standards for technical solutions as they become available;
- A list of specific actions completed to implement the MPIP; and
- A Blueprint of the MPIP.
Section 5 – Provider Incentive Program Blueprint, of this SMHP update, details the following processes used by DOM for oversight and administration of the MPIP, as required by CMS:

- The oversight of the MPIP that is conducted to ensure that providers meet all program requirements are met, including:
  - Compliance based upon their participation year;
  - Enrollment eligibility criteria;
  - Patient volume requirements;
  - EH incentive payment calculations remain consistent with CMS rules;
  - A/I/U and MU requirements are met prior to payment;
  - Monitoring and validation information; and
  - A process for combating fraud and abuse;

- Assurance that no amounts higher than 100 percent of Federal Financial Participation (FFP) will be claimed by DOM for reimbursement of expenditures for payments to providers;

- Assurance that no amounts higher than 90 percent FFP will be claimed by DOM for administrative expenses in administering the MPIP;

- Assurance that payments made to the approved providers are paid directly (or to an employer of facility to which the provider has assigned payments) without any reduction or rebate, and that incentive payment reassignments to an entity promoting the adoption of CEHRT as validated by DOM are voluntary for the provider involved;

- Assurance that providers receive only one incentive payment per program year;

- The Mississippi State Level Registry (MS SLR) attestation process, including specific identifiers used by DOM to coordinate with CMS on incentive payments;

- Assurance that only appropriate funding sources are used to make MPIP payments, including the methodology for verification;

- Assurance that MPIP payments are made for no more than a total of six years;

- Assurance that no provider begins receiving payments after Program Year 2016 and incentive payments cease after Program Year 2021;

- Assurance that an EH does not receive payments after fiscal year 2016 unless the hospital has received an incentive payment in the prior fiscal year;

- Executing timely and accurate payment of incentives;

- Recoupment/adjustment of incentive payments incorrectly disbursed; and

- The MPIP appeals process.
As DOM continues to refine this plan and provide updates to CMS, DOM will conduct operational and business planning to provide the following information:

- A description of the process to capture clinical quality data from each provider and a description of the methodology in place to verify this information; and
- The method by which DOM intends to address the needs of underserved and vulnerable populations, including information related to children, individuals with chronic conditions, Title IV-E foster care children, individuals in long term care settings, and the aged, blind, and disabled.

In addition to developing elements for the SMHP update, DOM has also been coordinating with the statewide HIE to promote the use of CEHRT to providers throughout the State of Mississippi as well as educate providers on the MPIP.

Based upon recommendations from a previously concluded CMS site visit, DOM will consider renaming the Mississippi State Level Registry to the Mississippi EHR Attestation System in future years. DOM plans to keep CMS informed of anticipated changes to activities, scope, or objectives. DOM will provide annual updates and as-needed updates to CMS as its plan evolves over the remaining program years.
3 Current HIT Landscape Assessment – The “As-Is” Environment

In the fall of 2010 DOM completed the first Environmental Scan of the State of Mississippi to ascertain the level of readiness of its providers. DOM also considered its current data-sharing partners and evaluated the level of readiness to expand its current data sharing capacity.

As requested by CMS, DOM completed a new Statewide Environmental Scan in 2017. The objective of the 2017 Environmental Scan was to assess the current status of the EHR program as well as HIT and HIE adoption within the State, then use that information to develop plans for the completion of the EHR program through its remaining years. The results of the completed 2017 Scan have been used to significantly update several sections of this SMHP and its related IAPDU.

3.1 The 2017 Statewide Environmental Scan

This section describes the 2017 Environmental Scan of the State of Mississippi’s Medicaid providers and the level of EHR adoption and Medicaid incentive payments. The subsections provide the assessment documents, the tools used, the analysis applied, and the outcomes of the 2017 Environmental Scan, and a historical context of any relevant HIT/HIE issues. These sections serve as a source of data for the development of the To-Be Landscape and completion of the HIT Roadmap and the IAPD.

DOM has conducted several comprehensive assessments of the current and planned levels of HIT adoption by Medicaid providers. These assessments began in June 2010, and include assessments up to September, 2012, as well as the full, statewide 2017 Environmental Scan. For the purposes of this document, HIT refers to health information technology (IT) that a provider might use, including practice management, health management records, EHRs, Laboratory Information Systems (LIS), ePrescribing Systems, electronic billing, and other clinical systems. The mechanisms utilized to collect this data included interviews, surveys, and focus groups. The entities interviewed or surveyed included all types and sizes of providers in a cross section of urban and rural settings, as well as providers in Federally Qualified Health Centers (FQHCs), Rural Health Clinics (RHCs), Agencies such as the State Department of Public Health and the United States Veterans Administration (VA), and Tribal settings. Data includes information gathered specifically for the SMHP, as well as information gathered for the Statewide HIE Strategic and Operational Plan (SOP), and other HIT related initiatives. As reflected in the information contained in Appendix H, DOM concludes that the incentive program has been a strong motivational factor for the adoption of CEHRT.

3.1.1 Background and Methodology for the 2017 Environmental Scan

The 2017 Environmental Scan was conducted in four phases that included planning, data collection, data analysis and reporting results. The planning process culminated in an Environmental Scan methodology that included data collected from a web survey, targeted interviews, and a focus group.
The web survey results were limited to a qualitative analysis due to a lower than anticipated response rate. The data collected indicates that, as expected, HIT/HIE has expanded significantly in Mississippi since the previous Environmental Scan of 2010.

The targeted interviews were conducted in-person and telephonically, and included provider types across Mississippi. Specifically, interviews were conducted with DOM, the Mississippi State Department of Health (MSDH), the Mississippi Health Information Network (MS-HIN), the Department of Veterans Affairs (VA), hospitals, physicians, professional healthcare associations such as the Mississippi Hospital Association, and community health centers. The results of the interviews reflected the various sentiments and experiences of each different stakeholder, but all were asked a specific set of questions and common themes and key information emerged from the participants.

A focus group was held to supplement the information from the targeted interviews. Discussion facilitated during the focus group related to strategic plans for EHR, technology status, HIE membership status and familiarity with the DOM Provider Access portal.

3.1.2 2017 Environmental Scan: Provider Web Survey

A web-based survey was determined to be the most efficient means of reaching the broadest audience possible. Therefore, although there was no guarantee of widespread participation, the web survey provided an appropriate start to the data collection effort. The survey was used to collect quantitative and qualitative data about adoption and use of electronic health information technology and the level of clinical data exchange throughout the state.

The following activities were completed to inform and develop the web survey approach:

- Analysis of past efforts and lessons learned
- Identification of the key questions and learning objectives for the web survey
- Identification of survey audience and outreach methods

Past survey efforts were leveraged to develop a starting point with the intention of focusing on the new information to be obtained regarding providers’ adoption and use of HIT and clinical data exchanges. A comprehensive list of providers throughout the state from varying locations and specialties was developed for survey distribution. The survey was ultimately distributed to over 1,650 providers using an official DOM e-mail address.

After development of a comprehensive set of survey questions and response selections, the survey was distributed to the target audience. As these types of surveys have historically received lower response rates, the survey was also posted on the DOM EHR Incentive Program website to promote the existence and importance of the survey to the provider community. Over the course of the five-week survey, each provider received at least two follow-up e-mails to drive additional responses. A final reminder was sent out one day before the survey closed to generate a last-minute push for increased responses. Understanding that the survey required 10-15 minutes for completion, the 5-week period was provided to allow for a reasonable and flexible window of time for survey completion.
The web survey gathered information from stakeholders and consumers in the Mississippi healthcare industry along strategic, operational and technical lines. The main areas of focus for technical and operational data gathered in the web survey included:

- Organization background
- HIT/EHR adoption
- Meaningful Use/Medicaid EHR Incentive Program
- Envision Web Portal knowledge and participation
- Provider Access portal knowledge and participation
- Electronic clinical data exchange
- Mississippi Health Information Network (MS-HIN) knowledge and participation
- Clinical data exchange trading partners.

The targeted audience for the web survey included a mix of stakeholders from the following types of organizations:

- Community Health Centers and Rural Health Clinics
- Hospital Association(s)
- Hospitals
- Long-term care providers
- Independent Laboratories / Reference Laboratories
- Independent Radiology Providers
- Pharmacies
- Physicians and Physician Practices
- Other healthcare providers.

### 3.1.3 2017 Environmental Scan: Web Survey Results

The results of the web survey were limited due to a low response rate of 69 respondents. However, the results revealed important findings and trends that indicate a significant expansion of electronic data exchange and technology in health care since the 2010 Environmental Scan.

The data collected from the survey was parsed and analyzed to determine key findings and trends in HIT/EHR adoption and clinical data exchange throughout the state. A statistical and qualitative analysis of the data received from responses was used to complement the focus group and targeted interview findings. The following pages highlight the key findings from the web survey in graphical and narrative form by topic.

**Organizational Background**

Over 60% of respondents represented primary care and family doctor practices, while 10% were dental, 7% behavioral health, and the remaining 23% were a mix of ophthalmology, pharmacy, radiology, and other organizations. The insurance utilization by patients included 36% Medicaid/CHIP, 23% Medicare, 30% private insurance, and 11% cash or uninsured.
Figure 1: Web Survey Results - Respondents by Classification

Level of HIT/EHR Adoption
EHR adoption tops the list of HIT with a usage rate of 94% among survey respondents, followed by adoption of electronic prescribing at 86%, adoption of practice management software at 75%, and adoption of clinical quality measure tools at 62%. Of those who have adopted an EHR, 56% plan to upgrade their system within the next 6 months.

Figure 2: Web Survey Results - Reported Usage of HIT

Respondents indicated that the main reasons preventing the usage of additional EHR functionalities include such functionality would slow clinical staff down (39%), high costs of
additional features (34%), lack of EHR feature knowledge (31%), and staffing and documentation issues (29%). Meanwhile, 26% indicated they have no concerns and use all their EHR features.

![Figure 3: Web Survey Results - Reasons Preventing use of more EHR Functionalities](image)

The web survey also found that telemedicine is used by or planned to be used by 27% of providers primarily for purposes of consultation with other physicians or hospitals, providing care, or viewing patient information at home.

**Meaningful Use / EHR Incentive Program Participation**

53% of respondents indicated that there are no barriers to participating in the EHR Incentive Program, while 36% found MU requirements to be confusing or burdensome. Only 3% of respondents were not familiar with the EHR Incentive Program. Of those survey participants that are participating in the EHR Incentive Program, 95% have achieved Modified Stage 2. According to information from the EHR incentive program, all participants from 2015-2017 have reached Modified Stage 2 status. There is no longer a Stage 1/Stage 2 classification.
Envision Web Portal
The Envision web portal, Mississippi’s MMIS portal for providers, is known by 77% of survey respondents, and of those, 52% use the portal frequently, 33% use the portal sometimes, while only 15% rarely use the portal. The survey found that the most beneficial Envision features, ranked highest to lowest, are to (1) check beneficiary eligibility, (2) review claim status, and (3) review patient claim information.
Provider Access Web Portal
The Medicaid provider clinical web portal, called Provider Access, allows providers to view Medicaid patient data online and was known by 47% of survey respondents. Of those familiar with Provider Access, 30% indicated they use the portal as part of their daily workflow.

Clinical Data Exchange
The most common method of clinical data exchange for survey respondents is via fax at 76%, followed by Direct Secure Messaging at 29%, and EHR systems are used by 22% of respondents for the exchanging of clinical data.

![Methods of Clinical Data Exchange](image)

Figure 6: Web Survey Results - Methods of Clinical Data Exchange

The following figure highlights provider usage of various electronic data exchange services (blue) as well as the services that providers would like to use in the future (orange). The top services in use today include order and lab results delivery, patient portals, public health data submission and direct messaging. Services with the highest percentage of provider interest in the future include active care coordination, MU analysis and reporting, historical lists, and discharge summaries.
The survey found that the most prevalent challenges faced by providers exchanging clinical data electronically are that referral partners’ technology can’t support exchanges (41%), software costs are too high (29%), and organizational concerns with sharing data (16%).

Mississippi HIE: Mississippi Health Information Network (MS-HIN)
HIE participation among respondents was 38%, or 24 providers. Of the 24, 12 respondents indicated membership with MS-HIN. Of the 12 who were members of MS-HIN, six indicated they use the lookup service less than monthly, while only two respondents use it daily. The MS-
HIN services most commonly used by respondents include the Mississippi Immunization Registry, Direct Secure Messaging, CCD/C-CDA, Community Health Record, and ADTs.

![Figure 9: Web Survey Results - Data Types Contributed to MS-HIN](image)

### Clinical Data Exchange Trading Partners
The survey found that the majority of Mississippi providers have clinical referrals in the central region (66%), followed by the northeast as the next highest region (36%). When asked who providers exchange or plan to exchange health information with, respondents indicated other physicians (75%), hospitals (62%), pharmacies (62%), laboratories and X-Ray facilities (58%), and governing agencies (26%).
3.1.4 2017 Environmental Scan: Targeted Interviews

The use of targeted interviews was the second data collection activity of the 2017 Environmental Scan. The targeted interviews were held with specific stakeholders to discuss positive and negative experiences with EHR and HIT adoption and use, how the EHR incentive program has impacted clinical practice, what barriers may exist to meaningful use, both to meet statutory MU measures as well as having a generally positive impact on clinical decision making and care coordination. The interviews were planned to identify specific activities that the state can do to foster greater HIT adoption and meaningful use.

To prepare for targeted interviews, the team focused on two main items:

- Analysis of the results of the web-based survey
- Identification of key information brokers

The qualitative information from the Web Survey was used to prepare for interviews and identify trends. These findings were paired with the findings from the previous Environmental Scan to identify historical trends as well.

The second preparation step for performing the targeted interviews was to identify key information brokers within the targeted audience. Through a comprehensive stakeholder analysis performed at the outset of the project, key stakeholders were identified who may offer valuable insight and interview questions were tailored specific to each stakeholder. Additional outreach was then conducted to the identified stakeholders to fill in information gaps.

The intent of all targeted interviews was to use an approved interview guide to ensure a consistent approach to obtaining desired information, but also to facilitate a fruitful conversation with the interviewees to gain additional useful information that will be used to
update the SMHP. An interview guide was developed and tailored for unique stakeholders and was also personalized based on the background research conducted prior to all interviews.

The targeted interview process gathered information from various stakeholders along strategic, operational, and technical subject matter. This information gathering method provided the most robust, candid and detailed data regarding HIT/HIE in Mississippi. Examples of the strategic and operational information that were gathered and assessed include, but are not limited to:

- Vision, mission and values
- Strategic plans for EHRs
- Privacy and security plans / concerns
- Oversight and enforcement concerns
- Accountability and transparency

Examples of technical information gathered and assessed include, but are not limited to:

- HIT / HIE / EHR adoption plans
- Utilization of DOM tools such as the Provider Access portal
- HIE membership and utilization
- HIT / HIE / EHR growth plans
- Interoperability

A modular interview guide was developed and used for each interview. The use of a modular guide allowed for quick adaptation to the specific interview audience. This guide was used to make sure the team obtained responses for identified information gaps. However, the targeted interviews were not a strict question and answer session, but instead an opportunity for interactive conversation that allowed more information to be unveiled that was not discovered through other methods of information gathering.

The preliminary list of targeted audiences included:

- Community Health Centers and Rural Health Clinics
- MSDH
- Health Plans and Payers
- Hospitals
- Long-term care providers
- Physicians and Physician Practices
- Trade Associations
- MS-HIN
- Other healthcare providers

In total, 41 stakeholders were interviewed, representing 31 organizations. The largest stakeholder group was hospitals and providers with 21 different organizations. The remaining stakeholder group breakdown is as follows: 6 Trade Associations; 1 Health Plan; 2 State Agencies; 1 Health Information Exchange Organization.
3.1.5 2017 Environmental Scan: Targeted Interview Results

The results of the targeted interviews reflected the varying sentiments and experiences each stakeholder group had with HIT in Mississippi.

Ongoing analysis of results from interviews was performed to confirm desired information was being discovered. This enabled the team to ensure all key information was being gathered and allowed for modification in tactics to ensure the Environmental Scan will meet its objectives.

Key findings include:

- Majority of health care providers have implemented and are effectively using an EHR
- Majority of electronic clinical data sharing is done through shared EHR vendors
- Common obstacles for exchanging data electronically include:
  - Technical infrastructure challenges (systems do not talk to each other)
  - Difficulties associated with HIT adoption (e.g., training and education of end users; interoperability tools not embedded within workflow)
  - Lack of timeliness of data exchange and accessing exchanged data
- Participation in the Medicaid EHR Incentive Program is high
- Multiple HIEs exist and membership in these multiple HIEs is high, although consistent use of HIEs for electronic clinical data exchange is low
- The most commonly used interoperability applications and/or HIEs include Care Everywhere, CommonWell, Relay Health, and MS-HIN

Results of the targeted interviews were analyzed and interpreted by subject matter experts and organized in a table by themes, examples, and common concerns heard from each stakeholder group.
3.1.6 2017 Environmental Scan: Focus Group

A focus group was used to supplement the targeted interviews and generate conversation among stakeholders to share ideas and future strategic plans of HIT adoption. The focus group further explored the data gathered during the web survey and targeted interviews which concluded the data gathering phase of the Environmental Scan.

The following activities were completed to inform and develop the focus group approach:

- Analysis of past efforts and lessons learned
- Identification of the key questions of the Environmental Scan
- Identification of target audience and outreach methods

The 2010 Environmental Scan was examined to determine what new information could be beneficial for the State and what information could be gained from the focus group. After the previous Environmental Scan was analyzed, gaps in information were identified, and the web survey results were considered, resulting in key learning objectives and questions to guide the focus group discussion.

The focus group methodology was comprised of five categories:

1. Targeted Participants
2. Structure and Design
3. Recruitment and Preparation
4. Focus Group Session
5. Data Analysis

Outreach and communications were conducted to solicit participation once the target audience was identified and the focus group questions were finalized. The following methods and tools were used for the focus group process:

- Communication methods: email (initial, follow-up, and confirmation)
- Documents: recruitment emails, focus group facilitator script, focus group ground rules, consent form, and short-form survey
- Locations: DOM office location
- Focus group sessions: One
- Data analysis: Transcription and recording

Target participants represented a cross-section of the market, such as providers and health care management leaders.

Six major topics were discussed in the focus group based on the results and preliminary analysis of the web survey and interview data. Open-ended questions that begin with “what”, “how”, and “why” were used to draw detailed conversations and answers. The team included three types of questions in the focus group:

1. Engagement: introducing the topic and making participants comfortable with the topic
2. Exploration: asking questions that will produce in-depth discussions
3. Exit: asking for all other opinions or ideas that were not discussed
The focus group was facilitated with a lead moderator and assistant moderator. The role of the moderator was to facilitate conversation and ensure all participants felt comfortable contributing honest viewpoints on each topic. The role of the assistant moderator was to manage the recorder and take notes.

Prior to the start of the focus group, participants were asked to review and sign a consent form acknowledging that the discussion would be recorded and that their identity would remain anonymous in any analysis or report. A facilitation script and ground rules were used to guide the focus group. This promoted professionalism and standardization in the session.

The web survey primarily provided quantitative data, while the targeted interviews provided a significant and productive set of qualitative data. These two data gathering methods were the basis for the 2017 Environmental Scan. However, the focus group was used to complement the targeted interviews by exploring the relationship between stakeholders and clarifying themes and concerns that were raised through analysis of the interview and web survey data.

### 3.1.7 2017 Environmental Scan: Focus Group Results

The data from the focus group in addition to the targeted interviews shed light on the perceived benefits and concerns of electronic data exchange and electronic health records. The stakeholders that participated in the focus group interviews were in favor of HIEs and EHRs, but also had important concerns and barriers. Workflow disruption, technical capabilities of disparate systems, cost, and prioritization were frequently discussed regarding HIT. The scope of HIE functionalities within each health system varied but clear trends were presented especially with smaller rural health systems versus larger urban health systems.

### 3.1.8 2017 Environmental Scan: Comparison with 2010 Scan

The team reviewed Section 3 – “Current HIT Landscape Assessment – The “As-Is” Environment of DOM’s most recent CMS approved SMHP. The SMHP As-Is Environment contains a series of findings from the 2010 Environmental Scan. The following table lists the conclusions from the 2010 Environmental Scan (Section 3.1.1.3) and updates the findings based on the 2017 Environmental Scan results.

**Table 3-1: 2010 versus 2017 Environmental Scan Results**

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<tr>
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<th>2010</th>
<th>2017</th>
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<tr>
<td><strong>Hospitals are becoming increasingly aware of the benefits of EHR technology and its positive impact on the quality of care for their patients.</strong></td>
<td>Mississippi paid hospital EHR incentive payments across a three-year span. All hospitals that participated have received full payments.</td>
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<td><strong>The exchange of electronic data between hospitals and</strong></td>
<td>Since 2010, Mississippi’s hospitals and networked providers have steadily moved to an integrated EHR model across their facilities.</td>
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Their providers is necessary for improvement of patient care and controlling costs.

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<td>The providers who have an integrated network can now seamlessly exchange clinical data with their integrated providers. Information from the targeted interviews confirms that hospitals are continuing to join and grow integrated networks to align hospitals and providers across the care continuum. The interview information also indicates that hospitals and providers are continuing to invest in improving and developing their integrated information systems to improve all performance metrics. Many indicated they were seeking population health improvement, and increasing their evidence-based practice to improve quality. Exchange of data across non-networked providers continues to expand, but continues to pose interoperability challenges. Many of the interviewees stated that they were not able to electronically exchange data with non-networked providers. Many indicated that fax is still a main method of data exchange. Examples of the challenges cited include:</td>
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<td>1) Different EHRs are not on the same platform or set of standards. Although, some hospitals and providers are indicating commitment to sharing through participation in HIEs such as CommonWell, Carequality or MSHIN; the use of a statewide or regional HIE is not a common solution considered.</td>
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<td>2) Organizations have direct messaging capabilities, however, they have encountered multiple issues of not being able to find a provider’s information and/or accurate information.</td>
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<td>3) Physicians will often not use a portal that is outside of their clinical workflow and EHR workspace. Therefore, HIE portals that require separate logins have limited use.</td>
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<td>4) Organizations have cited many competing priorities in regards to advancing their health information technology. Clinical data exchange efforts must be a priority in order for organizations to effectively invest in the technical setup and provider training and outreach.</td>
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<td>All hospitals recognize the inevitability of moving to an EMR/EHR system with the capability of exchanging clinical health care data beyond the integrated service delivery network.</td>
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<td>All hospitals have moved to an EHR system to meet Meaningful Use. Hospitals recognize the inevitability of moving towards data analytics and population health capabilities to meet MACRA and MIPS. To have full capability for data analytics and population health tools, hospitals recognize the need for more interoperability across the state.</td>
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| **The success of participation in exchanges relies on vendor ability to achieve certification.** | The 2010 finding was unclear whether it was applicable to the EHR vendor or the HIE vendor. Therefore, for 2017, the response includes a description of the current certification landscape and requirements for both EHR and HIE vendors.  

EHR vendors are required to be certified by the ONC. However, success of participation relies on cost, organizational priority, end user training and provider outreach in addition to having a certified EHR vendor.  

To participate in the Medicare and Medicaid EHR Incentive Program, EHR software must follow established standards and other criteria for structured data and be certified by CMS and ONC as a Certified Electronic Health Record Technology (CEHRT). Today, all providers must be using a CEHRT to the 2014 Edition Final Rule, or if available, the 2015 Edition Final Rule, or a combination of the two. Throughout this Environmental Scan, EHR certification was not a major concern or barrier to exchange participation.  

HIE vendors do not have a formal federal mandatory accreditation or certification requirement. However, there are currently multiple different organizations, some national and some state, that assess certain functions, including technology and security. For example, some of the certifications relate to secure messaging functions, some to query-based exchange. Minnesota, Pennsylvania, Vermont and Kansas have some mandatory requirements, and some are voluntary, such as Direct Trust and Healtheaway. However, there is not one comprehensive testing and certification service for HIE. |
| **The NwHIN (HealtheWay CONNECT) and the State HIE will provide the mechanisms to facilitate the secure exchange of patient data regardless of the location of the patient and his/her health records.** | The NwHIN has transitioned from ONC to an independent initiative, the eHealth Exchange, supported by the Sequoia Project (formerly HealtheWay) in 2012. The Sequoia Project manages the eHealth Exchange and Carequality interoperability initiatives. Three organizations that were interviewed are a member of the eHealth Exchange. The majority of organizations rely on the interoperability application of their EHR vendor to participate in electronic clinical data exchange. The most cited bi-directional interoperability platforms were Care Everywhere (Epic) and Relay Health (McKesson).  

HIEs (e.g., the Mississippi Coastal Health Information Exchange (MSCHIE)), RHIOs, and system-wide record sharing will continue to increase in parallel with a statewide HIE effort. The establishment of standards is  

After Hurricane Katrina in 2008, five health systems (Singing River Health System, Memorial of Gulfport, Hancock Medical, Biloxi Regional and Coastal Family Health Center clinics) partnered to create the Mississippi Coastal HIE. MSCHIE was the pilot project for an HIE in Mississippi. Subsequently, in 2009, HITECH passed, and thereafter, state legislation passed forming MS-HIN and its governance structure. MSCHIE expanded and became part of MS-HIN. The current statewide HIE in Mississippi is MS-HIN. MS-HIN |
critical to interoperability and alignment with the existing exchanges.

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<td>currently has 41 connected facilities. 19 more are being on-boarded with approximately 12 that will be live in the summer of 2017. There has been a 72% increase from 2015 in connected hospitals. MS-HIN offers a community health record, electronic referrals, Direct Secure Messaging, medication history, clinical results delivery, MU reporting, bi-directional immunization gateway, bi-directional CCD/C-CDA gateway, HISP services, orders and results services, and event notification services. MS-HIN remains a gateway for unaffiliated organizations and parties in Mississippi to exchange clinical data electronically. The establishment of agreements on rules of engagement for information sharing and how those rules can be changed is important for healthcare organizations to have trust in the platform and become an active member. CommonWell has created a vendor neutral platform with common standards and policies. Carequality, an initiative managed by the Sequoia Project, has developed similar standards including a common “rules of the road,” technical specifications and a participant director to enable cross network exchange. The eHealth Exchange functions as a HIE and is based on federal standards with which all members agree. Lastly, the Direct Project, which was launched in 2010, enables a secure, standards-based way to electronically send health information and has been a consistently cited method of electronic data exchange among Mississippi providers. Standardized data continues to be important to facilitate aggregation for population health and data analytics in the future.</td>
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Providers have a strong interest in improving their patients’ quality of care.

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<td>Since 2010, physician reimbursement has been updated to include incentives for quality of care. First there was MU in 2010 followed by the Medicare Access and CHIP Reauthorization Act (MACRA) in 2015. This legislation created a new payment methodology for quality of care. Part of MACRA is the Merit Based Incentive System (MIPS), and it combines three existing quality initiatives MU, PQRS and VBPM into a single program. Although MACRA and MIPS do not currently apply to Medicaid providers, this shift to value-based payment methodology will likely impact how CMS and states consider payment mechanisms for Medicaid. This is shown in the shift to ACOs, PCMH and other bundled payment models. In conclusion, with MU only continuing to apply to Medicaid providers, and the implementation of MACRA/MIPS quality requirements, providers continue to have a strong interest in improving their patients’ quality of care. Additionally, interviewees noted that the health system a physician is affiliated with, and therefore, the technical capabilities the physician has access to, determines their investment in electronic clinical data exchange. The</td>
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<td>physician’s age cohort was also determined to be a factor in their use of electronic clinical data exchange.</td>
<td>Health systems have moved to one system for Ambulatory and Inpatient. Smaller providers are also joining larger health systems and accessing the same EHR platform. Therefore, exchanging data from provider offices to hospitals within the same health system has become routine. The most common types of data shared include laboratory results, problem list, patient demographics, allergies, disease management data, and medication data. The top health data exchange partners continue to be other physicians, hospitals, pharmacies, laboratories and X-Ray facilities.</td>
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<td>Providers are focused on first exchanging data with hospitals and pharmacies.</td>
<td>Practices with fewer than ten practitioners are more likely to meet the 30 percent Medicaid requirement.</td>
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<td>Practices with fewer than ten practitioners are more likely to meet the 30 percent Medicaid requirement.</td>
<td>Practices with fewer than ten practitioners remain more likely to meet the 30 percent Medicaid population served requirement.</td>
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<td>Providers show a significant interest in the Health Information Technology for Economic and Clinical Health (HITECH) incentive program.</td>
<td>June 30, 2017 was the deadline for eligible providers to submit for EHR Program Year 2016 and be eligible to receive the full five years of incentive payments. Ninety-two percent of Medicaid providers in Mississippi who registered in the Medicaid EHR incentive program are participating in the program.</td>
</tr>
<tr>
<td>The large majority of respondents indicated they intend to apply for the stimulus payments in 2011. Most respondents intend to upgrade or replace their systems.</td>
<td>The majority of respondents indicated they have applied for HITECH stimulus payments since 2011. Some respondents implemented their first EHR systems, others have transitioned to a new vendor, while others upgraded their system because of the EHR incentive program.</td>
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<tr>
<td>Providers need community outreach programs to understand the incentive program details regarding eligibility.</td>
<td>2016 was the midpoint of the EHR Incentive program, and is the last year providers can begin to participate. From the level of continued participation through MU (92%), providers understand their eligibility for the EHR incentive program.</td>
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<td>Providers need community outreach programs to understand the requirements of MU and Clinical Quality Measures (CQM) for the Medicaid EHR incentive program.</td>
<td>The majority of providers understand the requirements of MU and Clinical Quality Measures (CQM) for the Medicaid EHR Incentive Program. However, as more value-based purchasing options are introduced for Mississippi providers, outreach efforts should continue as a main priority.</td>
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3.1.9 2017 Environmental Scan: Additional Findings

Apart from the updated findings relevant to the conclusions drawn from the 2010 Environmental Scan, additional findings were discovered as a result of changes in HIT and clinical data exchange capabilities across the state. Based on the survey, targeted interviews, and focus group, new findings from the 2017 Environmental Scan include:

- MS-HIN is viewed as one of several HIEs supporting clinical data exchange for hospitals and providers in Mississippi;
- Although Mississippi ranks low on broadband availability (34% of the population without broadband access ranks Mississippi 50th among all states per the Federal Communications Commission), broadband is not a concern nor a limiting factor among the provider community;
- Many organizations offer their own online patient portal;
- Workflow integration has driven value of interoperable EHR platforms that are able to perform all or nearly all of providers HIT needs in one system;
- Clinical data exchange and HIT adoption is generational; those less familiar with technology are more likely to be opposed to new initiatives until penalties become too costly;
- Providers are adopting telemedicine as the technology becomes increasingly more prevalent;
- The EHR Incentive Program has reached maturity so no new registrations will occur after June 2017.

3.2 MMIS Capabilities Assessment

Mississippi’s current MMIS is a three-tiered application architecture composed of:

1. A client work station (user interface tier);
2. An application server (business logic tier); and
3. A mainframe backend (data tier).

The business logic and data tier are housed in a secure data center facility in Pennsylvania with MMIS’ vendor Conduent. The user interface tier workstations are located in DOM facilities in the State of Mississippi. The workstations run a PowerBuilder runtime client and the presentation layer of the Envision system on the Windows Vista Professional operating system. The workstation application handles primary edit logic prior to sending the data on to the business logic tier for further processing.

The business logic tier provides: 1) middleware connectivity to the mainframe environment; 2) clustering, load-balancing, failover, and two-phase commit control over the database transactions; and 3) additional business logic processing via PowerBuilder and Java objects. The mainframe-based data tier uses IBM Customer Information Control System for transaction processing and DB2 for relational database management.

The major components of the current MMIS include:

- Data Entry
• Claims
• Managed Care Enrollment, Capitation Payment and Reporting
• Financial
• Reference
• Management and Administrative Reporting
• Third Party Liability
• Provider
• Surveillance and Utilization Review
• Beneficiary
• Medicare Buy-In
• Automated Voice Response System
• Provider Lookup
• Bulletin Board System
• WINASAP – Provider claims submission software
• Web Portal
• Electronic Data Interchange (EDI) Processing
• Computer Systems Request

The Decision Support System (DSS) and Data Warehouse (DW) components include:

• Data Warehousing
• Management and Administrative Reporting
• Surveillance and Utilization Review (J-SURS)
• Data Management Tools

Lastly, pharmacy claims processing include:

• Point of sale terminals
• Pharmacy Benefit Management (PBM)

The State of Mississippi completed final contract negotiations to procure a Service-Oriented Arrchitecture MMIS, including pharmacy claims processing, a DSS / DW solution and Fiscal Agent services. The procurement effort emphasized vendor achievement and alignment of Medicaid Information Technology Architecture (MITA) principles and goals as key outcomes of the process. As a result of subsequent CMS directives for modular solutions, CMS contract approval is contingent upon modular implementation where feasible. As changes to the awarded vendor solution become necessary, DOM will work with the vendor to define solutions that will also achieve the CMS requirements for modularity.
Additionally, DOM is procuring services to perform a revised State Self-Assessment (SS-A) using the new Medicaid Information Technology Architecture (MITA) 3.0 guidelines. DOM will consider the appropriate solution to address opportunities identified for MITA level advancement during each subsequent re-procurement effort. Funding for the new MITA SS-A is not included in the corresponding HIT IAPDU, but will be requested in a separate MES IAPD.

### 3.3 Feasibility of Incentive Payment Methodology

The State of Mississippi studied two possible solutions for administering the MPIP – one involving in-house development of a provider incentive payment system; and a second option involving a standalone Web-based hosted solution developed by Conduent. DOM elected to use the Conduent solution, which involved minimal changes to the current MMIS.

The Conduent solution was designed and implemented in conjunction with Conduent’s work on the California replacement MMIS. Since 2011 it has been implemented in multiple states as a Software as a Service (SaaS) solution. Conduent’s solution offers DOM a Web-based State registration, attestation, and tracking system to support provider incentive payments for the A/I/U and/or MU of CEHRT. This Web-based system was designed to provide a State Level Registry (SLR) to document, track, and attest to the provider’s use of EHRs in support of A/I/U and MU requirements. This SLR works in conjunction with and communicates with the CMS Registration & Attestation System in accordance with the published CMS interface specifications.

The Conduent solution provides two separate Web portals: one for the provider access and one for State staff to access.

The provider portal is a single location where providers can securely log in to complete their A/I/U and MU attestation information, including uploading any additional required documentation for acceptance and review by the State. The provider portal allows each eligible provider to complete registration and to review and edit their demographic information. However, data received from the CMS Registration & Attestation System must be edited through the Medicare/Medicaid EHR Incentive Program Registration Website.

As a part of the MS SLR attestation process, providers enter the following information into the provider portal:

- Medicaid patient volume percentage numerator and denominator to achieve eligibility. This will also be analyzed for non-hospital based eligibility;
- Required A/I/U data (or MU data and percentage information, including CQMs); and
- Supporting documentation.

The MS SLR automatically verifies provider data, such as license validation and exclusion checks, and indicates a preliminary approval or denial in accordance with current CMS and DOM requirements. Providers are able to track the status of their application and payments online, and view any messages from DOM.
The State access portal provides a location where DOM-approved users can securely log in to access provider attestation information and work queues. The work queues for DOM users are role-based so that the provider registration and attestation information can be routed to the correct user and/or department for approval, action, or denial. The State access portal provides a mechanism by which the State can track incentive payments and communicate with providers through a messaging system. In this way, the State can communicate “directly” with the providers on matters of approval or denial, or to request additional information.

Approved State users utilize the State access portal to:

- Review and approve provider attestation information and supporting materials;
- Calculate and initiate a provider payment cycle using an automated interface to the MMIS;
- Manage the audit, recoupment and adjustment, and appeals processes; and
- Review provider quality metrics.

The following is an alternatives analysis that DOM used to compare the Conduent proposed stand-alone solution with an effort to develop an in-house system to provide functionality needed for issuing provider incentive payments.

The in-house system was investigated and process flows were developed to show the required changes in workflow to accommodate provider payments. The outcome of that process is represented in the figures shown below. The first set of figures represents the proposed new process flow for EPs and the second set of figures represents the proposed new process flow for EHS.
Figure 12: Internal Process Flow - Professional Eligibility
The following table shows the comparison that DOM made between the internal solution and the Conduent solution. As noted above, the Conduent option was used as an adjunct to the current MMIS, requiring only minimal changes to the current MMIS. This option had several...
advantages that are discussed below. Critical factors in DOM’s decision-making process were critical timeline, availability of qualified staff, and investment in infrastructure.

**Table 3-2: Internal Solution versus. Conduent Solution**

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<thead>
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<th>Considerations</th>
<th>Internal Solution/SaaS Solution</th>
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| The State has indicated a desire to participate in Group 1 testing for the provider incentive payments with CMS. | **Internal Solution**: The changes necessary for participating in Group 1 testing will not be available in time.  
**SaaS Solution**: Vendor commits to meeting the required timelines. |
| The State desires a solution that poses the least risk of schedule delay.       | **Internal Solution**: The State does not have the required resources necessary to successfully develop and implement the solution.  
**SaaS Solution**: The vendor is devoting significant resources to creating a solution for multiple states. |
| The State desires a solution that requires the least amount of limited State resources. | **Internal Solution**: The required State resources will be significant under this scenario (support, maintenance, development, program, help desk, project management, and vendor oversight). The State would struggle to recruit sufficient qualified resources in a timely manner.  
**SaaS Solution**: The State’s required commitment of resources is significantly decreased by focusing its limited resources on the oversight of the proposed solution. |
| The State desires a solution that meets all Mississippi-specific requirements.   | **Internal Solution**: An internal solution will be able to meet any Mississippi-specific requirements.  
**SaaS Solution**: The Conduent solution may not meet all Mississippi-specific requirements. Small changes, such as additional fields are included in the cost, but substantial modifications may be expensive or time consuming. |
| The State desires a solution that conforms to all CMS requirements.             | **Internal Solution**: An internal solution would require additional manual processes for attestation and verification, but will be able to meet all CMS requirements fully.  
**SaaS Solution**: The Conduent proposal includes a Web-based system to support the MU requirements, incentive payments, and other ARRA HITECH Act requirements. This solution provides a more automated solution for the attestation and verification processes, therefore requiring fewer DOM resources for the oversight of the attestation and verification processes. |
### Considerations

| The State desires a solution that is flexible, easily modifiable, and maintainable. | **Internal Solution**: Building applications that are flexible, easy to modify, and maintain is a challenge. The State may struggle to create an internal solution to meet these objectives while altering a legacy MMIS at the same time.  
**SaaS Solution**: The vendor states that the application will be configurable for state specific requirements, but not enough information is known to verify flexibility. |

| The State desires a solution that provides as much automation as possible for audit functions. | **Internal Solution**: An internal solution may be able to automate audit functions fully; but design, development, and implementation would take a significant amount of time beyond the timeline for Group 1 or Group 2.  
**SaaS Solution**: The Conduent proposed solution provides automation of audit functions. The full extent of those automation capabilities is unknown at this point. |

Based on a review of the alternatives, the State chose to pursue the Conduent SaaS solution. The State believed that the SaaS offered the lowest risk and a lower cost alternative, long-term, than developing a new internal solution. The State worked closely with Conduent to finalize the requirements for the State of Mississippi in the commercial off the shelf (COTS) offering using configuration sessions and user group calls. Since implementation, the Conduent application has successfully accepted provider attestations for A/I/U and MU and DOM continues to work on shaping functionality within the Conduent solution to meet the needs of the MPIP and future regulatory changes to the MU program.

#### 3.4 Medicaid Electronic Health Record System and e-Prescribing System (MEHRS) Transition to the DOM Medicaid Clinical Infrastructure (MCI) subproject

##### 3.4.1 Background on the MEHRS System

With the use of funds from a Transformation grant, a provider Stabilization grant, and the MMIS enhanced funding match, the State of Mississippi implemented a system known as Medicaid Electronic Health Record System and e-Prescribing (MEHRS/eScript). The MEHRS/eScript system was launched in June 2010 and was available to all Mississippi Medicaid providers at no charge.

DOM requested and received funding for MEHRS design, development, and implementation, as well as ongoing support, via an IAPD that was approved by CMS in January 2009. CMS approved a four-year contract term with two two-year renewals with the understanding that the renewals would require further approval.

DOM contracted with the vendor Shared Health to provide a MEHRS/eScript for Mississippi Medicaid providers in 2009. Shared Health subsequently rolled out MEHRS/eScript with over
3,200 Medicaid providers and practice staff users registering for the system, enabling electronic health records with clinical data for over 600,000 Medicaid beneficiaries in MEHRS/eScript.

Shared Health was contracted to upgrade the deployed version of MEHRS/eScript (Version 7) to an Office of the National Coordinator for Healthcare Information Technology (ONC)-certified version, named MEHRS/eScript Version 8. MEHRS/eScript Version 8 was due for delivery to DOM in late 2011, as mutually and contractually agreed by both DOM and Shared Health; however, it was not delivered.

In early 2012, DOM was informed that Shared Health would not be delivering Version 8 of MEHRS/eScript, would not be delivering any ONC-certified version of MEHRS/eScript, and that Shared Health was stopping all development work on the MEHRS/eScript product and platform.

As DOM had providers who were relying on the MEHRS/eScript system for meeting the criteria of Stage 1 Meaningful Use, DOM and Shared Health entered into an agreement to migrate/upgrade the MEHRS/eScript system to a commercially available solution, through new (subcontracted) vendors, Orion Health and Mede/Analytics.

Orion Health began the Operations Phase of the MEHRS/eScript project on July 1, 2013 and continued working on the operations of the project through March 2014. As stipulated in Orion’s’ contract, Orion successfully implemented a certified EHR product to DOM. However, after the implementation of the upgraded ATCB MEHRS/eScript Electronic Health Record and integrated ePrescribing components to DOM, it was determined that many Medicaid providers had adopted commercially available EHR/ePrescribing solutions to comply with Meaningful Use (MU) requirements. With the deadline looming for the required ONC 2014 certification, and with the diminished need and requirements for the MEHRS EHR/ePrescribing components of the MEHRS solution, DOM made the decision to initiate a strategic realignment of the project as of June 30, 2014.

Core components of the MEHRS/eScript solution were retained and upgraded to support the DOM clinical data interoperability strategy (as defined in the SMHP as the ‘To-Be’ infrastructure and also defined in the HIT IAPD). These Medicaid-specific clinical data components include the Medicaid Clinical Data Repository (CDR), Medicaid Master Patient Index (MPI), and Provider Access (a provider web portal for Medicaid Providers), and these three components, along with the addition of clinical data Analytics formed the basis for the DOM Medicaid Clinical Infrastructure (MCI).

### 3.4.2 DOM Medicaid Clinical Data Infrastructure (MCI)

The existing DOM MCI, provided by MedeAnalytics, is composed of a Medicaid Clinical Data Repository (CDR), a Medicaid Master Patient Index (MPI), a Medicaid provider portal (Provider Access), and Medicaid Clinical Data Analytics. A description of each is below:

- DOM Medicaid Master Patient Index (MPI): The DOM MPI is a SaaS-based, modular component that is coupled with the DOM MCI. The DOM MCI allows for beneficiary identification via the complex, unattended probabilistic matching algorithm developed specifically for DOM and DOM’s use-cases. There are over 2.6 million historical
identities in the DOM MPI, with approximately 800,000 active beneficiaries, and a match rate of approximately 92%.

- DOM Medicaid Clinical Data Repository (CDR): The DOM CDR is a SaaS-based, modular component that is coupled with the DOM MCI. The DOM CDR allows for the storage and utilization of Medicaid-only clinical data from internal DOM systems and external DOM trading partners. Medicaid clinical data is first validated using a process within the DOM MPI to ensure the data is for an active Medicaid beneficiary, and then stored in the DOM CDR. The existing MMIS provides on a scheduled basis data to the DOM CDR (and validated via the MPI process as previously detailed) for Medical Claims files, Pharmacy Claims files, and other files (detailed in Figure 14. Clinical Data Interoperability Project). These MMIS files are then transformed into clinical data and stored in the DOM CDR. Terminology and Sensitivity services are applied to all incoming and outgoing clinical data, via the CDR. Upon request (demand) from a DOM system or external trading partner (such as UMMC), the CDR generates a CCDA in XML (in real time) for a single or multiple Medicaid beneficiaries.

- DOM Medicaid Clinical Portal (Provider Access): The DOM Provider Access portal is a secure portal where Medicaid providers can log in and search, view and download clinical data on Medicaid beneficiaries. MU Stage 2 compliant C-CDAs in XML can be downloaded from Provider Access as clinical summaries, for import into MU Stage 2 certified EHRs.

- DOM Medicaid Clinical Data Analytics: The DOM Medicaid Clinical Data Analytics provides DOM the ability to run custom reports on clinical data, claims data, and clinical data with claims data. Business users within DOM depend on this Analytics solution for program decision making, care and cost review, and responding to Legislative and CMS requests for information, etc.

In 2016, DOM added additional capabilities to the existing MCI when DOM implemented the first real-time EHR interface in the United States between a State Medicaid Agency and a Provider EHR (Clinical Integration). This Clinical Integration allows for the real-time query-exchange of C-CDAs clinical data summaries between the DOM MCI and the Provider’s EHR, and is occurring with the largest Medicaid provider in the State, the University of Mississippi Medical Center (UMMC).

Specifically, this Clinical Integration allows the UMMC Epic EHR to query the DOM MCI (and the DOM CDR) within seconds, and allows for a Medicaid clinical summary to instantly be sent back to the provider’s EHR. By having real-time Clinical Integration and providing the Medicaid C-CDAs directly in the provider EHR, providers can now instantly view, import, and utilize the DOM Medicaid clinical data from the DOM CDR. The Clinical Integration is bi-directional, meaning the Medicaid clinical data in the DOM CDR is updated at the end of each encounter at UMMC, thereby further enhancing the rich clinical data in the DOM CDR with every encounter by a Medicaid beneficiary at UMMC.

DOM and UMMC are exchanging over 4,000 clinical summaries (C-CDAs in XML) daily, and have surpassed one million C-CDAs exchanged in one year, affecting the care and quality of care for thousands of Medicaid beneficiaries, as they seek care, in real-time and within the provider’s native EHR workflow. Additionally, DOM has completed Clinical Integrations with the
Hattiesburg Clinic, the second largest Medicaid provider in the State, and has also completed a Clinical Integration with the Singing River Health System, a large Medicaid health system on the Mississippi Gulf Coast.

The Clinical Integrations with the Hattiesburg Clinic and the Singing River Health System have added approximately 4,000 additional clinical summaries (C-CDAs in XML) exchanged daily with DOM, bringing the total (with UMMC) to over approximately 8,000 clinical summaries exchanged daily between these three health systems and DOM. This daily exchange of clinical summaries (C-CDA in XML) supports approximately 8,000 Medicaid beneficiaries as they seek health care each day.

DOM is currently working on a Clinical Integration with the Delta Health Alliance (DHA) a large network of Ambulatory providers and several connected FQHCs in the Mississippi Delta, and is in discussions with other large health systems for future Clinical Integrations to support Medicaid beneficiaries as they seek health care. The DHA Clinical Integration is planned to go live with DOM in the winter of 2017.

The DOM MCI, including all Clinical Integrations, is currently live and is supported by Mede/Analytics, as the primary vendor until September 30, 2021. The DOM MCI is one component of the DOM Clinical Data Interoperability Program (CDIP), described below.

### 3.5 DOM Clinical Data Interoperability Program (CDIP)

As outlined in the Executive Summary of this document, DOM has outlined four goals to accomplish to improve the coordination of care and quality of care of Medicaid Beneficiaries in the State of Mississippi using clinical data and HIT/HIE: 1) Achieve greater interoperability with Medicaid providers and provider clinical systems (EHRs, other clinical systems) to aggregate provider-based Medicaid clinical data and store/utilize this data in the existing DOM Clinical Data Repository; 2) Utilize the aggregated provider-based Medicaid clinical data in the DOM Clinical Data Repository for Agency goals and programs including clinical data analytics and clinical data population health; 3) Offer tools and interfaces to providers so that providers may access and utilize the aggregated clinical data in the DOM Clinical Data Repository, including such tools as a Medicaid clinical data provider portal and real-time, bi-directional clinical data interfaces to support the sharing and updating of Medicaid clinical data interoperability within provider EHRs and provider EHR workflows; and 4) promote adoption of CEHRT for DOM providers with the goal of using CEHRT and HIT/HIE for promoting coordinated health care for DOM beneficiaries, better health care outcomes, and improvements in care quality.

The DOM Clinical Data Interoperability Program includes the infrastructure and personnel for DOM to support these four goals, including the aggregation of Medicaid clinical data from Medicaid providers, DOM utilization of the aggregated Medicaid clinical data for Agency goals and programs, DOM offered tools and interfaces to allow for the sharing of the aggregated Medicaid clinical data with provider clinical systems (EHRs, LIS, and other clinical systems) and current clinical workflows, and promoting and supporting the adoption of CEHRT and HIT/HIE technologies by Medicaid providers.

There are several benefits from the aggregation of Medicaid provider clinical data by DOM, including but not limited to:
- Medicaid beneficiary care coordination and improved care management;
- Agency goals and programs, such as Medicaid clinical data analytics and Medicaid clinical data population management; and
- Aggregated and up-to-date Medicaid beneficiary clinical summary documents, clinical reports, clinical data, and decision-making available in real-time and integrated directly into the provider EHRs and clinical system for real-time provider utilization in a care setting.

The Clinical Data Interoperability Project consists of three subprojects, two of which are existing and currently functioning:

- **Subproject 1, The Mississippi Health Information Network (MS-HIN) Integration** – This integration subproject has not been implemented at this time, and, in the future will support Medicaid clinical data exchange between DOM’s MCI and MS-HIN. More information about this subproject can be found in the To-Be section of this document, as this is not a functioning subproject.

- **Subproject 2, Medicaid Clinical Infrastructure (MCI)**  DOM has an existing, functional MCI with core clinical components of a Clinical Data Repository (CDR), Master Patient Index (MPI), Medicaid Provider Portal, Medicaid Analytics, and Medicaid Provider Clinical Integrations (EHR Interfaces), as explained in detail in the MEHRS section of this document. The existing MCI has been integrated with the existing DOM Interoperability Platform, and currently supports bi-directional clinical data from providers via the DOM Interoperability Platform. The MCI subproject is required to be interoperable with the other two subprojects.

- **Subproject 3, Interoperability Platform** – DOM has procured and implemented an Interoperability Platform from the vendor DXC Technology (formerly known as Hewlett-Packard Enterprise Services, or HPE) as a single point of connectivity. The DOM Interoperability Platform is a SOA-based, SaaS module, allowing interoperability between DOM components such as the existing MMIS and the future MES, the modernized Eligibility system, the DOM MCI and Clinical Integrations with Medicaid providers, other DOM internal systems and services, as well as external DOM trading partners (such as MS-HIN, other State Agencies, etc.). The DOM Interoperability Platform has been integrated with the DOM MCI as well as the Clinical Integrations, and will support a future integration with MS-HIN. The DOM Interoperability Platform is a modular service director that assists DOM in connecting all of the modular components of the internal DOM ecosystem, as well as DOM’s external trading partners. The Interoperability Platform is key component in DOM’s strategy for SOA, modularity, COTS, and MITA 3.0 compliance. The two major components of the Interoperability Platform include an Enterprise Service Bus (ESB) and an eHealth Exchange Gateway. The DOM Interoperability Platform subproject is required to be interoperable with the other two subprojects.

**Additional Integration Points and Systems for the CDIP and MITA alignment:**
• MMIS: Currently, DOM’s existing MMIS is in the final stages of its natural lifecycle with a contract expiration date of 6/30/2020. DOM received approval by CMS for a new MES contract in early 2017, with the vendor DXC Technology (DXC) providing the overall modular solution. The new, replacement MES is nearing the beginning stages of implementation, and will be integrated with the existing DOM CDIP components, including the existing DOM MPI and DOM CDR, and will utilize the existing DOM Interoperability Platform via the ESB as the modular component for connectivity between all the modules (MES, CDR, MPI, etc). By utilizing these modular, SaaS-based COTS components, DOM continues to align with the goals of SOA and MITA for modularity and COTS. It is not DOM’s objective to integrate the MMIS with the MS-HIN at this time. DOM expects to continue the current process of feeding the claims data from the new MES into the Medicaid CDR on a weekly basis. An interface will be required for the new MES similar to the one that currently exists with the MMIS. It is not anticipated that the new MES will house clinical data for DOM, rather, the new MES will access the DOM MCI components (the MPI and CDR), via the Interoperability Platform (as a service director) to share data as needed. Medicaid clinical data will reside in the existing CDR, but will be made available as a service to the new MES via the Interoperability Platform. For the time being, Identity Management will remain at the existing individual modules for each program (MPI module for clinical data and at the MES and E&E modules for administrative data); however, DOM’s long-term strategy is to achieve a single Medicaid identity across the SMA.

• Clinical Quality Measures (CQMs): As DOM aggregates rich clinical data in C-CDA (XML) format from providers and various other clinical data sources, DOM utilizes the existing Analytics capabilities of the existing DOM MCI to analyze this clinical data. DOM is also currently investigating a possible CQM pilot, with a CQM evaluation tool and process, which would utilize the aggregated rich clinical data (C-CDA in the DOM CDR). This potential pilot and CQM evaluation tool would allow DOM to support use-cases such as the analysis of at-risk populations, costs, quality among providers and quality of care, and other CQM-related use-cases, utilizing the DOM aggregated clinical data. Currently, DOM is not analyzing CQM data submitted with yearly EHR Attestations, and DOM is only capturing the data that is reported. Mississippi simply reports this data aggregately to CMS through our Annual Report. DOM continues to evaluate how best to utilize the rich clinical data that DOM is aggregating, including electronic submission of CQMs.

• Outreach and Training Resources: DOM has two full-time outreach and training resources that are responsible for adoption of the DOM Provider Access clinical data portal across the state. DOM’s Provider Access portal is populated with clinical data from the DOM Clinical Data Repository and is available at no cost to every Medicaid provider in the state. The outreach resources assist with adoption of the portal and provide education about clinical data exchange activities to support Medicaid beneficiaries (improving and coordinating care, etc.). Mississippi currently conducts extensive training and support each year. This is done through a one-to-one or small group approach as needed. DOM also hosts a number of informative, training webinars. We use our program mailing list to notify providers of yearly changes and webinar invitations.
DOM also uses contact information data from CMS registration for the EHR Incentive program to send reminders. DOM focuses on those that created a program registration but never created a State Level Registry (SLR) Account. Mississippi is in the 80th percentile of EP Registration to Paid (as of February 2017).

CMS requires all Meaningful Use participants to meet the Public Health Reporting Objective, Measure 1 – Immunization Registry Reporting by uploading the evidence of Active Engagement to submit data from an EHR to a Public Health Agency. DOM strongly encourages all providers to complete the Registration of Intent to Onboard with the Immunization Registry survey. Most providers adhere to this recommendation and complete this Registration Survey. However, Providers that claim an exclusion to the measure (due to provider type or specialty) must submit a brief memo describing the reason for the exclusion. DOM has asked for this since 2013 and has seen a large increase in registration with the Immunization Registry.

Figure 14: Clinical Data Interoperability Project (CDIP)
3.6 Current MITA Status

MITA is a CMS initiative designed to assimilate business and IT transformation across the Medicaid enterprise to improve the administration of the Medicaid program. MITA is a business-centric architectural framework that provides planning guidelines for states to define strategic business goals and objectives, define business processes, and assess current capabilities as a baseline to measure progress towards these goals.

A key activity within the MITA initiative is performing a MITA SS-A. Requests for FFP for MMIS enhancements must include a formal SS-A which describes the extent to which current MMIS systems reflect MITA and how requested changes will advance their transformation into the new architecture.

HIT, like MITA aims to improve quality of care through an open architecture that supports the integration of clinical and administrative data, promotes interoperability, and coordination with partners to improve health outcomes.

Mississippi completed a MITA SS-A in 2008 and a subsequent Gap Analysis was completed in June, 2010. The purpose of a completed SS-A is to identify the “As Is” state and “To Be” (target) state of a state’s Medicaid business enterprise. The Mississippi Medicaid enterprise has many business processes that are currently in the Level 1 maturity with some business processes in Level 2.

Since 2010, DOM formalized the business process narratives and mapping as a part of the Mississippi MITA goals. DOM plans to convert these business processes into MITA 3.0 standards in the coming year as a part of the ongoing update process that aligns current MITA status with MITA goals.

DOM has advanced in its use of technology to supply information to providers in the following ways:

- Implementation of Mede/Provider Access product. This assists providers in the appropriate treatment of beneficiaries and reduces unnecessary testing.
- Receipt of a large percentage of claims through electronic data interchange (EDI).
- Electronic funds transfer (EFT) is used to payment nearly all providers.
- Widespread usage of the Envision Web portal.

Although DOM has advanced its use of technology towards the MITA standard, challenges remain. For example, some business processes must still be performed and/or validated manually. DOM has not developed all of the business processes necessary to utilize the DSS to its full capacity. Care planning and care management are fields open to the State for increased gains in population health and cost savings.

The MITA SS-A results are a valuable resource in planning for the MMIS replacement with a modularized Medicaid Enterprise Solution. DOM is planning to perform a new SS-A using the new MITA 3.0 requirements. DOM will assimilate the results of the MITA 3.0 SS-A into the timing
and scope plans for the modularized MES architecture. Future plans will be coordinated to ensure the new MES modules will support the Medicaid EHR Incentive Program.

3.7 Current Broadband Initiatives

The State of Mississippi has had a public mandate to improve access to broadband technology since 2003 when the Mississippi Broadband Technology Development Act was passed (Miss. Code Ann. § 57-87-1 et. seq.). The Mississippi Broadband Task Force was founded in 2004 to promote citizen use of the Internet with a plan and broadband strategy. Since that time, the State has been moving forward with planning and implementation of improved access to broadband services. Over $77 million in grant funding was awarded to the Office of the Governor through federal broadband stimulus programs. The funding is to be used to expand broadband access and adoption in communities across the State of Mississippi. Specifically, the State is participating in the national broadband mapping and planning initiative through the Broadband Technology Opportunities Program (BTOP) administered by the Department of Commerce (DOC).

In April 2009, Governor Haley Barbour charged the Mississippi Broadband Task Force with the development of strategies to enhance the broadband infrastructure in Mississippi. The National Telecommunications and Information Administration (NTIA) awarded the State of Mississippi a grant as part of the BTOP, under the ARRA. With this funding, Mississippi is eager to deploy the Long Term Evolution (LTE) broadband network to better serve the citizens of the State. LTE is a next generation mobile broadband technology designed to support data applications that are currently too bandwidth-intensive for the existing technology. Additionally, on August 18, 2010, the State received a $7.1 million grant through the Broadband Initiatives Program (BIP) to design, engineer, and construct a broadband network in rural northeastern Mississippi.

In September 2010, the Office of the Governor received an additional award from NTIA of nearly $5 million for broadband planning and mapping activities under the State Broadband Data and Development Program, a matching grant program that implements the joint purposes of the ARRA and the Broadband Data Improvement Act. This is a supplement to the original $2 million award the State received in January 2010, allowing the extension of its current two-year broadband data collection program for an additional three years and allowing the State to identify and implement best practices in broadband mapping. The State of Mississippi will utilize a portion of the funding to support the creation of the Mississippi Broadband Connect Coalition, a non-profit, public-private partnership focused on producing a comprehensive statewide strategic plan for improving digital literacy, increasing access to broadband and enabling greater adoption of broadband in the State.

The Mississippi Broadband Connect Coalition (MBCC) began partnering with the Mississippi State University Extension Service in 2011 to develop the statewide strategic plan. This 125-member public-private partnership met for almost 9 months to create the statewide strategic plan titled, “Mapping Mississippi’s Digital Future,” a long-term plan that addresses recommendations on how to improve broadband usage across several policy areas. The policy areas included education, healthcare, workforce development, government performance and public safety. The long-term plan identifies barriers to further broadband deployment in Mississippi as well as why broadband is not more widely adopted. Finally, the plan looks at ways to improve broadband access specifically with the Delta and Tribal communities in Mississippi.
As previously documented, providers who participated in the 2017 Environmental Scan stated that Broadband access is not a challenge, and Broadband is widely available in the State of Mississippi. Although Mississippi ranks low on broadband availability (34% of the population without broadband access ranks Mississippi 50th among all states per the Federal Communications Commission), broadband is not a concern nor a limiting factor among the provider community, per the 2017 Environmental Scan.

### 3.8 Coordination with Medicare and Federally Funded, State Based Programs

DOM is participating with CMS to pay providers and is using the CMS Registration & Attestation System and MS SLR to coordinate Provider incentive payments with Medicare.

### 3.9 Coordination with the Statewide Health Information Exchange

DOM participated in the Mississippi Health Information Network (MS-HIN) SOP effort as a member of the Technical Infrastructure and Finance Domain Groups. The Statewide HIE SOP was submitted to the ONC in September 2010.

The structure for MS-HIN is set forth in Miss. Code Ann. §§ 41-119-1, et seq., entitled Health Information Technology Act, included as Appendix F. The governing body is the MS-HIN Board of Directors. The Board of Directors was appointed at the end of September and the first meeting was held on October 20, 2010. The overall structure for MS-HIN is shown in Figure 6: MS-HIN Organization Structure in Section 4.7.2.

MS-HIN has a broad representation of stakeholders, including Hospitals, clinics, individual providers, and service providers. After hurricane Katrina in 2008, five Mississippi integrated health systems (Singing River Health System, Memorial Hospital of Gulfport, Hancock Medical Center, Biloxi Regional Medical Center and the Coastal Family Health Center clinic system) partnered to create the Mississippi Coastal HIE, or MSCHIE, and MSCHIE acting as the pilot project for HIE in Mississippi. Subsequently, in 2009, HITECH passed, and thereafter, state legislation passed forming MS-HIN and its governance structure. MSCHIE expanded and became part of MS-HIN. MS-HIN currently has 41 connected facilities, which are primarily Hospitals. 19 more are being on-boarded with approximately 12 that will be live by the summer of 2017. There has been a 72% increase from 2015 in the number of MS-HIN connected hospitals. MS-HIN offers a community health record, electronic referrals, Direct Secure Messaging, medication history, clinical results delivery, MU reporting, bi-directional immunization gateway, bi-directional CCD/C-CDA gateway, HISP services, orders and results services, and event notification services.

The MS-HIN Board of Directors maintains oversight responsibility for all HIE activities in the State of Mississippi. DOM is a member of the MS-HIN Board of Directors and works in partnership with the MS-HIN, providing leadership to assure that Medicaid beneficiaries are best represented and served by the MS-HIN. DOM is providing leadership to assure funding for MS-HIN in accordance with the fair share principles and cost allocation defined in guidance from
CMS provided as part of the State Medicaid Director Letter dated May 18, 2011, along with subsequent State Medicaid Director Letters, such as the Letter dated February 29, 2016.

DOM continues to coordinate efforts with MS-HIN to support interoperability and a non-duplication of efforts. As a part of this coordination, DOM submitted an HIE IAPD as a part of the 2014 SMHP and IAPD submission process. CMS subsequently approved the HIE IAPD, including budget for FFY 2015, FFY 2016, and FFY 2017, however funding has not been expended as of this point in time. DOM is working with MS-HIN on an updated HIE IAPD for the FFY 2018 and 2019 timeframe, and will utilize the February 29, 2016 CMS Medicaid Director’s Letter for ongoing guidance.

DOM and MS-HIN are discussing implementation of technologies and interfaces, per the HIE IAPD, to support interoperability for Medicaid clinical data between MS-HIN and DOM. Data supported in this bi-directional exchange with MS-HIN includes Medicaid specific clinical data including the Consolidated-Clinical Document Architecture (C-CDA) patient summary on Medicaid beneficiaries.

Candice Whitfield was the previous HIT Coordinator for the State and was simultaneously the acting Executive Director of the Mississippi Health Information Network (MS-HIN). Candice assumed a role with UMMC approximately two years ago and Jeremy Hill took her position at MS HIN. The Department of Health (Public Health) is the entity that provides administrative oversight for MS HIN. Currently, the role of State HIT Coordinator is not filled.

3.10 Current Public Health Initiatives

The Mississippi State Department of Health (MSDH) has implemented a Health Data System (HDS) designed to improve data quality and efficiency of collection, as well as improve the ease of submission. The system is comprised of Rhapsody Connect, an integration broker that includes a rules engine, database, and secure messaging product. The primary goal of the HDS is to establish and maintain a centralized reporting system by collecting hospital discharge data from each licensed health care facility in Mississippi. In addition to the hospital discharge data collection and evaluation, the MSDH’s Office of Epidemiology interfaces to the HDS. The HDS will also be used to support disease registry information relating to heart disease, stroke, and asthma. With the future expansion of HDS, the MSDH is planning to interface the system with the State’s Trauma Registry, as well as conduct syndrome surveillance and participate in electronic laboratory reporting. As of July 2010, the system will perform automatic reporting of reportable diseases and conditions to the Centers for Disease Control and Prevention (CDC).

At this time, the MSDH communicates with CDC through the PHIN MS Rhapsody. MS-HIN is providing the MS-HIN infrastructure for all MS-HIN stakeholders to connect to MSDH to support these Public Health initiatives. MSDH has expressed the interest and forthcoming ability to exchange data with DOM (via the DOM-MS-HIN connection).

MSDH provides administrative oversight for MS-HIN, the State HIE. Through coordination and collaboration, MSDH has appointed MS-HIN as the single gateway for providers to access the State Immunization Registry which requires that the providers onboard to MS-HIN as the only way to submit their immunization information. DOM currently does not have access to the State Immunization Registry data.
DOM, MSDH and MS-HIN are continuing to coordinate on plans for additional future connections with other federal public health and welfare programs (Health Resources and Services Administration, Substance Abuse and Mental Health Services Administration or Indian Health Services), and will continue to collaborate and coordinate, so as not to create a duplication in efforts (connectivity, interoperability, etc.)

3.11 Federally Qualified Health Centers /Rural Health Clinics

Mississippi has 21 FQHCs. The FQHCs and RHCs are working together and exchanging health care information via shared systems. For example, at least four FQHCs are working with the Delta Health Alliance (DHA) and sharing data via the DHA’s cloud-based Allscripts EHR implementation.

Before the EHR Incentive Payment Program, Delta Health Alliance (a 501C3 Corporation) received a Beacon Grant to improve the quality of health in the rural farming counties in Mississippi. Delta Health Alliance continues to provide support and services to seventeen counties. Various funding sources are used to improve the quality of life, education and health care in their counties.

Ryan White Grants (HRSA Funding) provide funding for the treatment of patients with HIV/AIDS. Various Mississippi FQHCs, RHCs, and FQHC Lookalikes receive Ryan White grants.

None of the above grants are funded through the State of Mississippi. The Delta Health Alliance Beacon Grant was reviewed with CMS and it was determined not to impact the Mississippi Medicaid EHR Incentive Payment Program.

FQHC, RHC, IHS, and FQHC Lookalikes do not receive any funding from the State of Mississippi outside of the Division of Medicaid fee for service payments. FQHCs, RHCs, and FQHC Lookalikes’ base rates were set in 2001 and adjusted in 2002. The rates are adjusted annually based on the Medicare Economic Index. Rates for after-hours visits and Telemedicine have been added to the fee for service payments.

3.12 Department of Defense and Veteran’s Administration

There are three major military installations in the State of Mississippi: two are Air Force bases near Columbus and Biloxi and the third is a Navy facility near Meridian. While the military has expressed an interest in receiving information about off-base treatment of military personnel, they have been unable to connect to the State to retrieve the information due to severely restrictive security constraints.

In addition to the two large Veterans hospital facilities in Mississippi – one in Biloxi and one in Jackson, the Board of Veterans Affairs is located in Jackson, Mississippi. The DoD and the Veterans Administration (VA) are currently running the VLER EHR, however, recent developments have the DoD and VA moving away from VLER to a different EHR system. At this time there have been no further activities to integrate the VA and DoD with DOM.
3.13 Indian Health Services

Choctaw Indians are the most prevalent minority of the American Indian population in the State of Mississippi. Members of the Mississippi Indian Tribe receive basic health care through a community health service. Representatives of the Tribe indicate they are participating with Indian Health Services and anticipate connecting to DOM through MS-HIN in the future.

Presently, the Mississippi Choctaw Reservation has eight communities: Bogue Chitto, Bogue Homa, Conehatta, Crystal Ridge, Pearl River, Red Water, Tucker, and Standing Pine. Currently, there has been no further progress to integrate or share data with the IHS or tribes.
4 To-Be HIT Landscape

This section aligns the current As-Is HIT Landscape with the vision of DOM for adoption, promotion, and enhancement of EHR technology for Medicaid providers and for promotion of electronic exchange of Medicaid clinical data with DOM. This section also describes the goals and objectives and additional functionality that is planned to promote interoperability, providing the greatest benefit from the MMIS data and participation in the exchange of data with the MS-HIN, Medicaid providers, and DOM, using the DOM Interoperability Platform.

4.1 Future Vision for Providers

A key component of the Mississippi HIT strategy is continual meeting and yearly attestation of EHR Meaningful Use (Stage 3) by providers in the Medicaid EHR Incentive Program. To that end, DOM will continue offering a Web-based system for provider incentive payment attestations. The MS SLR is a public-facing application available over the Internet where providers supply registration and attestation data related to the incentive program. The Website can be reached directly or from a link on the current Mississippi MMIS Envision Web portal and the DOM Website. The MS SLR, described in further detail in Section 4.1.1, below, provides an easily accessible, easy to use system for the providers participating in the MPIP.

DOM will continue providing outreach and training to the provider community to enhance CEHRT updates and understanding of Stage 3 Meaningful Use through 2021. Further information on these efforts can be found in Section 6 – HIT Roadmap, of this document.

Table 4-1 shows DOM’s goals for provider adoption and MU of CEHRT in Mississippi:

<table>
<thead>
<tr>
<th>Provider Type</th>
<th>FFY 2011 - 2017</th>
<th>FFY 2018</th>
<th>FFY 2019</th>
<th>FFY 2020</th>
<th>FFY 2021</th>
<th>FFY 2022</th>
<th>Totals to Date</th>
<th>Totals to Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adopt Certified EHR</td>
<td>MU of EHR</td>
<td>Adopt Certified EHR</td>
<td>MU of EHR</td>
<td>MU of EHR</td>
<td>MU of EHR</td>
<td>Adopted Payments</td>
<td>MU Payments</td>
</tr>
<tr>
<td>Hospitals</td>
<td>95</td>
<td>186</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>95</td>
<td>186</td>
</tr>
<tr>
<td>Physicians</td>
<td>1835</td>
<td>2983</td>
<td>0</td>
<td>1200</td>
<td>1300</td>
<td>1300</td>
<td>1300</td>
<td>1835</td>
</tr>
<tr>
<td>Dentists</td>
<td>233</td>
<td>32</td>
<td>0</td>
<td>30</td>
<td>33</td>
<td>36</td>
<td>40</td>
<td>44</td>
</tr>
<tr>
<td>Nurse Practitioners</td>
<td>1167</td>
<td>1374</td>
<td>0</td>
<td>450</td>
<td>495</td>
<td>550</td>
<td>550</td>
<td>1167</td>
</tr>
<tr>
<td>Certified Nurse Midwives</td>
<td>14</td>
<td>31</td>
<td>0</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>14</td>
</tr>
</tbody>
</table>

Table 4-1: Total Payment Counts (Actual and Projected)
The following table shows the Performance Measures that DOM will use to gauge progress against the goals listed above:

**Table 4-2: Performance Measures for EH/EP and EHR Goals**

<table>
<thead>
<tr>
<th>Performance Measure</th>
<th>Method and Data Sources</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of EPs who received an EHR Incentive Payment for MU by the end of FFY 2016</td>
<td>Obtain a report from the MS SLR with the number of unique EP's by individual NPI, not Group, that received at least one EHR Incentive Payment for MU</td>
<td>2,983</td>
</tr>
<tr>
<td>Number of EHs who received an EHR Incentive Payment for MU by the end of FFY 2016</td>
<td>Obtain a report from the MS SLR with the number of unique EH's that received at least one EHR Incentive Payment for MU.</td>
<td>95</td>
</tr>
</tbody>
</table>

### 4.1.1 Mississippi State Level Registry Application

The core functions of the MS SLR Web application that are currently active in the MS SLR are categorized into the following five groups:

- MS SLR registration and view of CMS Registration & Attestation System data;
- Verification of Medicaid eligibility;
- Attestation to Meaningful Use under Modified Stage 2 criteria for Program Year 2017 and under Stage 3 criteria for Program Years 2018, going forward
- Review and approval; and
- Submission of payments.

The Current MS SLR functionalities are further detailed in Section 5 – Provider Incentive Program Blueprint.

Conduent continued to enhance functionalities within the MS SLR, including three major areas of development:

- Appeals – detailed appeals tracking and status reporting;
- Audits – initiation, tracking and reporting of provider audits; and
• Recoupments/adjustments – creation of the payment file (positive or negative) for total recoupments or payment adjustments.

These functional areas were released in August 2013. Audits and Appeals are processed through an external system, in accordance with state law and reported to the State Level Registry and to CMS.

Stage 2 changes were incorporated into the MS SLR during 2013 for hospital attestation beginning October 2013 and eligible professional attestation beginning January 2014. These changes included allowing providers to use a 90-day reporting period, regardless of the stage of MU, for 2014 only. In addition, Stage 2 changes included modifications to the Core and Menu Objectives and the Clinical Quality Measures as required in the Final Rule. Mississippi implemented CMS-mandated program changes known as the 2014 Flexibility Rule. Under the 2014 Flexibility Rule, eligible professionals were required to meet 17 core objectives, 3 menu objectives, and they would select 9 clinical quality measures (CQMs) from a list of 64. Eligible Hospitals and CAHs were required to meet 17 core objectives, 3 menu objectives, and 16 CQMs from a list of 29. Participants were allowed to select CEHRT software that was certified at either the 2014 level, 2011 level or a combination of both certification levels.

The Modified Stage 2 platform was implemented on April 29, 2016 for Program Years 2015 – 2017. All participants were expected to use only CEHRT software that was certified at the 2014 level and were given some alternate measure exclusion opportunities for those that were expected to demonstrate MU (years 1 and 2) during the 2015-2016 Program Years. There were no alternate measure exclusions available during the 2017 Program Year. Modified Stage 2 criteria created a bridge between the Previous Stage 1/Stage 2 criteria and the upcoming Stage 3 reporting requirements (beginning January 1, 2018). All participants were given a 90-day EHR Meaningful Use reporting period for Program Years 2015-2017, in accordance with CMS regulations. All CQM reporting requirements remain the same as the requirements from Program Year 2014. Although functionality currently exists in the State Level Registry for electronic CQM reporting, none of Mississippi’s providers took advantage of this functionality. All CQMs were manually entered.

The CQMs that DOM has been collecting will be aligned with the CQMs the three Medicaid Managed Care Organizations (MCOs) are required to report to DOM, from the MCOs day to day business of managing and coordinating the care for DOM beneficiaries. All three MCOs are required to report common metrics for Quality to DOM, which will be coordinated with the other DOM collected Quality metrics. In the DOM Quality Strategy Report (which will be delivered to CMS later this year), DOM outlines an overall Quality Strategy, and includes a roadmap to monitor and implement quality improvement (while allowing for periodic updates to strengthen and improve the effectiveness of the program). This DOM Quality Strategy provides a framework to communicate the State's vision, objectives and monitoring strategies addressing issues of health care cost, quality and timely access. The Quality Strategy contains the following domains: Maternal Health, Child Health, and Disease-Based Initiatives for Diabetes, Influenza, Hepatitis, and Hemophilia. As a part of the DOM Quality Strategy, the following quality measures will be monitored and published on DOM’s website annually beginning in 2018 for the 2017 measurement period.
## ADULT CORE SET MEASURES

### Primary Care Access and Preventive Care

- Cervical Cancer Screening (CCS-AD)
- Chlamydia Screening in Women Ages 21–24 (CHL-AD)
- Flu Vaccinations for Adults Ages 18 to 64 (FVA-AD)
- Screening for Clinical Depression and Follow-Up Plan (CDF-AD)
- Breast Cancer Screening (BCS-AD)
- Adult Body Mass Index Assessment (ABA-AD)

### Maternal and Perinatal Health

- PC-01: Elective Delivery (PC01-AD)
- PC-03: Antenatal Steroids (PC03-AD)
- Contraceptive Care – Postpartum Women Ages 21–44 (CCP-AD)*
- Prenatal and Postpartum Care: Postpartum Care (PPC-AD)

### Care of Acute and Chronic Conditions

- Controlling High Blood Pressure (CBP-AD)
- Comprehensive Diabetes Care: Hemoglobin A1c (HbA1c) Testing (HA1C-AD)
- Comprehensive Diabetes Care: Hemoglobin A1c (HbA1c) Poor Control (>9.0%) (HPC-AD)
- Diabetes Short-Term Complications Admission Rate (PQI01-AD)
- Chronic Obstructive Pulmonary Disease (COPD) or Asthma in Older Adults Admission Rate (PQI05-AD)
- Heart Failure Admission Rate (PQI08-AD)
- Asthma in Younger Adults Admission Rate (PQI15-AD)
- Plan All-Cause Readmissions (PCR-AD)
- HIV Viral Load Suppression (HVL-AD)
- Annual Monitoring for Patients on Persistent Medications (MPM-AD)

### Behavioral Health Care
### Initiation and Engagement of Alcohol and Other Drug Dependence Treatment (IET-AD)
- Medical Assistance with Smoking and Tobacco Use Cessation (MSC-AD)

### Antidepressant Medication Management (AMM-AD)
- Follow-Up After Hospitalization for Mental Illness: Age 21 and Older (FUH-AD)

### Diabetes Screening for People With Schizophrenia or Bipolar Disorder Who Are Using Antipsychotic Medications (SSD-AD)
- Follow-Up After Emergency Department Visit for Mental Illness or Alcohol and Other Drug Dependence (FUA-AD)*

### Diabetes Care for People with Serious Mental Illness: Hemoglobin A1c (HbA1c) Poor Control (>9.0%) (HPCMI-AD)*
- Use of Opioids at High Dosage in Persons Without Cancer (OHD-AD)

### Adherence to Antipsychotics for Individuals with Schizophrenia (SAA-AD)

### Experience of Care
- Consumer Assessment of Healthcare Providers and Systems (CAHPS®) Health Plan Survey 5.0H, Adult Version (Medicaid) (CPA-AD)

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### CHILD CORE SET MEASURES

#### Primary Care Access and Preventive Care
- Weight Assessment and Counseling for Nutrition and Physical Activity for Children/Adolescents – Body Mass Index Assessment for Children/Adolescents (WCC-CH)
- Chlamydia Screening in Women Ages 16–20 (CHL-CH)
- Childhood Immunization Status (CIS-CH)
- Well-Child Visits in the First 15 Months of Life (W15-CH)
- Immunizations for Adolescents (IMA-CH)a
- Developmental Screening in the First Three Years of Life (DEV-CH)
- Well-Child Visits in the Third, Fourth, Fifth and Sixth Years of Life (W34-CH)
- Children and Adolescents’ Access to Primary Care Practitioners (CAP-CH)
<table>
<thead>
<tr>
<th>Category</th>
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<tr>
<td><strong>Adolescent Well-Care Visit (AWC-CH)</strong></td>
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<td><strong>Maternal and Perinatal Health</strong></td>
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<td>Pediatric Central Line-Associated Bloodstream Infections (CLABSI-CH)</td>
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<td>PC-02: Cesarean Section (PC02-CH)</td>
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<td>Audiological Evaluation No Later Than 3 Months of Age (AUD-CH)</td>
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<td>Live Births Weighing Less Than 2,500 Grams (LBW-CH)</td>
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<tr>
<td>Contraceptive Care – Postpartum Women Ages 15–20 (CCP-CH)*</td>
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<tr>
<td>Frequency of Ongoing Prenatal Care (FPC-CH)</td>
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<td>Prenatal and Postpartum Care: Timeliness of Prenatal Care (PPC-CH)</td>
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<td><strong>Care of Acute and Chronic Conditions</strong></td>
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<td>Ambulatory Care: Emergency Department (ED) Visits (AMB-CH)</td>
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<td>Medication Management for People with Asthma (MMA-CH)</td>
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<td><strong>Behavioral Health Care</strong></td>
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<td>Follow-Up Care for Children Prescribed Attention-Deficit/Hyperactivity Disorder (ADHD) Medication (ADD-CH)</td>
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<td>Follow-Up After Hospitalization for Mental Illness: Ages 6–20 (FUH-CH)</td>
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<td>Child and Adolescent Major Depressive Disorder: Suicide Risk Assessment (SRA-CH)</td>
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<td>Use of First-Line Psychosocial Care for Children and Adolescents on Antipsychotics (APP-CH)*</td>
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<td>Use of Multiple Concurrent Antipsychotics in Children and Adolescents (APC-CH)</td>
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<tr>
<td><strong>Dental and Oral Health Services</strong></td>
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<td>Dental Sealants for 6–9 Year-Old Children at Elevated Caries Risk (SEAL-CH)</td>
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<tr>
<td>Percentage of Eligibles Who Received Preventive Dental Services (PDENT-CH)</td>
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<tr>
<td><strong>Experience of Care</strong></td>
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<tr>
<td>(CAHPS®) Health Plan Survey 5.0H – Child Version Including Medicaid and Children with Chronic Conditions Supplemental Items (CPC-CH)</td>
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</tbody>
</table>
The MS State Level Registry will implement the Stage 3 requirement set for Program Year 2017 reporting for those EPs that have successfully upgraded their CEHRT software to the 2015 certification level. Full Stage 3 Implementation is expected to be available for all providers (EP and EH) that remain in the EHR Incentive Program for Program Year 2018, going forward. All participants will have a 90-day EHR Meaningful Use reporting period for Program Year 2018, in accordance with CMS regulations. Program Year Attestation Submission windows will open each year in January and will close each year on the last day of April. This more closely follows the Medicare or MIPS attestation season and allows additional time for state staff to work with the increased number of Medicaid attesters in Mississippi.

4.2 Future MES Capabilities

The State of Mississippi completed final contract negotiations to procure a new solution referred to as the Mississippi Medicaid Enterprise System (MES) to include a state-of-the-art MMIS, including pharmacy claims processing, a DSS / DW solution and Fiscal Agent services to meet the business needs of DOM. As a result of recent CMS directives for modular solutions, DOM and the awarded vendor will evaluate the vendor’s solution sets for an implementation that will meet the new CMS requirements by defining components of the proposed solution that will be modularized during implementation or that can facilitate future modular procurements. The ambition is to preserve as much of the procurement effort to-date possible to upgrade the DOM core systems and services while accommodating the new CMS modularity requirements and respecting state and federal procurement guidelines. Due to limited state resources, a four-year implementation is still planned for the new system and services but a phased approach will be used where possible.

It is the goal of DOM MES procurement to:

- Improve communication and administration of the Medicaid Program;
- Provide timely and accurate adjudication of Medicaid claims;
- Increase data storage and improve data retrieval and reporting capabilities for Medicaid and the CHIP;
- Replace proprietary systems (e.g., clearinghouse and DSS/DW) with more technologically advanced and less costly products, which will result in more efficient operation of the Medicaid Program;
- Meet the requirements of MITA 3.0 standards, such as Service Oriented Architecture (SOA) using Enterprise Service Bus (ESB) infrastructure; and
- Allow for interface with the DOM Interoperability Platform.

The State request for proposals (RFP) is designed to move DOM forward in its vision of a Medicaid Enterprise that:
• Meets CMS certification requirements;
• Is aligned with the current MITA framework and future MITA frameworks\(^1\);
• Is aligned with CMS Enhanced Funding Requirements: Seven Conditions and Standards\(^2\);
• Implements all MITA business processes with the maximum business capability level possible – identifying any business processes that are at Level 1 or Level 2 and moving progressively to Level 3 or higher;
• Provides support for an open, flexible, and cost effective Medicaid Enterprise architecture;
• Utilizes an ESB for interfaces, including to the DOM Interoperability Platform, the MMEDS and/or new eligibility system, MS SLR, DOM CDIP and associated clinical systems, and other associated systems and environments, SOA, and Web services technology to allow for disparate system communication;
• Implements the latest technology standards - International Classification of Diseases (ICD-10), NCCI edits, Health Level Seven (HL7 – including offering increased support for the HL7 C-CDA), HIPAA version 5010 transactions, including the HIPAA 278 transaction, and the National Council for Prescription Drug Programs (NCPDP) Version 3.0 pharmacy claims;
• Uses a rules-based engine for ease of definition and update of eligibility and operational rules processing;
• Presents a browser-based Medicaid Enterprise system for minimal desktop footprint, ease of application update, and ubiquitous access for all users;
• Provides interface to the SLR, including support for the current and future SLR implementations;
• Provides an interface to the remediated MMEDS eligibility system. The new MMIS will require a future interface to a new eligibility system when MMEDS is re-procured; and
• Provides architecture for future interface to the DOM Interoperability Platform with the support of both clinical and administrative transactions with DOM trading partners, including MS-HIN.

\(^1\) MITA Framework 3.0 was released in 2012 and includes final policies on everything but eligibility and enrollment. Enhanced funding requirements – Seven Conditions and Standards will be incorporated into MITA 3.0.

\(^2\) CMS has issued new standards and conditions that must be met by the states in order for Medicaid technology investments (including traditional claims processing systems, as well as eligibility systems) to be eligible for the enhanced match funding, details can be found on the document Medicaid IT Supplement (MITS-11-01-v1.0), [https://www.cms.gov/Medicaid-Information-Technology-MIT/downloads/Enhanced-Funding-Requirement-Seven-Conditions-and-Standards.pdf](https://www.cms.gov/Medicaid-Information-Technology-MIT/downloads/Enhanced-Funding-Requirement-Seven-Conditions-and-Standards.pdf)
4.3 Future Vision for DOM Clinical Data Interoperability Program (CDIP)

As described in the As-Is, DOM has implemented the DOM MCI and DOM Interoperability Platform, as core subprojects in the DOM Clinical Data Interoperability Program (CDIP). DOM intends to support the MS-HIN subprojects, as well as interoperable exchange of Medicaid clinical data with DOM Medicaid providers, Medicaid trading partners, and Medicaid stakeholders, while improving care for Medicaid beneficiaries.

The DOM Strategy and Vision is an ecosystem of connected, interoperable Medicaid Providers, Medicaid trading partners and Medicaid stakeholders in the State of Mississippi. The expectation of DOM is to fully align with the SMHP and IAPD, as well as federal HIT-enabled health reform(s), including CMS Medicaid Information Technology Architecture (MITA) missions, goals and objectives.

The DOM CDIP includes the infrastructure and personnel for DOM to support the stated four goals in the Executive Summary section of this document, including, the aggregation of Medicaid clinical data from Medicaid providers, DOM utilization of the aggregated Medicaid clinical data for Agency goals and programs, DOM offered tools and interfaces to allow for the sharing of the aggregated Medicaid clinical data with provider clinical systems (EHRs, LIS, and other clinical systems) and current clinical workflows, and promoting and supporting the adoption of CEHRT and HIT/HIE technologies by Medicaid providers.

There are several benefits from the aggregation of Medicaid provider clinical data by DOM, including but not limited to:

- Medicaid beneficiary care coordination and improved care management;
- Agency goals and programs, such as Medicaid clinical data analytics and Medicaid clinical data population management; and
- Aggregated and up-to-date Medicaid beneficiary clinical summary documents, clinical reports, clinical data, and decision-making available in real-time and integrated directly into the provider EHRs and clinical system for real-time provider utilization in a care setting.

The Clinical Data Interoperability Project consists of three subprojects, as detailed in the As-Is section of this document. These three subprojects, two existing and one a future subproject, each could have upgrades, modifications, enhancements, and integrations as part of the To-Be environment, as described below.

- **To-Be Subproject 1, Mississippi Health Information Network (MS-HIN)** – This integration subproject will support Medicaid clinical data exchange between DOM’s MCI and MS-HIN. While not currently implemented at this date, this subproject will establish connectivity between DOM and MS-HIN to allow for Medicaid clinical data exchange, as well as potentially facilitate DOM’s connectivity and interoperability needs to external stakeholders. A primary use-case for the MS-HIN subproject is to support the flow of clinical data from MS-HIN’s Medicaid providers to DOM, thereby allowing DOM to populate the existing DOM Clinical Data Repository (CDR) with clinical data on Medicaid
beneficiaries. This Medicaid clinical data, typically in Health Level-7 (HL-7) format, includes the Consolidated Clinical Document Architecture (C-CDA) document. The MS-HIN subproject will be interoperable with the other two subprojects.

The MS-HIN To-Be environment required to achieve interoperability with DOM includes several future (To-Be) components, interfaces, and integration, including:

- A future MS-HIN Master Patient Index (MPI) that will be harmonized to the DOM (existing) MPI;
- A future MS-HIN Enterprise Service Bus (ESB) that will be connected to the DOM (existing) Interoperability Platform to support bi-directional clinical data query and exchange;
- Future integration between the existing MS-HIN eHealth Exchange Gateway (Sequoia Project) and the DOM existing eHealth Exchange Gateway (Sequoia Project) for connectivity to external stakeholders;
- Future connectivity, integration, and testing services between DOM and MS-HIN.

**Existing Subproject 2, Medicaid Clinical Infrastructure (MCI)** - DOM has an existing, functional MCI with core clinical components of a Clinical Data Repository (CDR), Master Patient Index (MPI), Medicaid Provider Portal, Medicaid Analytics, and Medicaid Provider Clinical Integrations (EHR Interfaces), as explained in detail in the As-Is section of this document. The existing MCI has been integrated with the existing DOM Interoperability Platform, and currently supports bi-directional clinical data from providers via the DOM Interoperability Platform. The MCI subproject will be interoperable with the other two subprojects.

The MCI To-Be Environment includes additional Clinical Integrations, and harmonization of systems with other State Agencies, including:

- Future additional Clinical Data Integrations, to support clinical data exchange between DOM and Medicaid providers, Payers, and State Agencies;
- Future harmonization between DOM systems and other State Agency systems to allow for data exchange;
- Future connectivity and integration services.

**Existing Subproject 3, Interoperability Platform** – DOM has procured and implemented an Interoperability Platform from the vendor DXC Technology (formerly known as Hewlett Packard Enterprise Services, or HPE) as a single point of connectivity. The DOM Interoperability Platform is a SOA-based, SaaS module, allowing interoperability between DOM components such as the existing MMIS and the future MES, the modernized Eligibility system, the DOM MCI and Clinical Integrations with Medicaid providers, other DOM internal systems and services, as well as external DOM trading partners (such as MS-HIN, other State Agencies, etc.). The DOM Interoperability Platform has been integrated with the DOM MCI as well as the Clinical Integrations, and will
support a future integration with MS-HIN. The DOM Interoperability Platform is a modular service director that assists DOM in connecting all of the modular components of the internal DOM ecosystem, as well as DOM’s external trading partners. The Interoperability Platform is a key component in DOM’s strategy for SOA, modularity, COTS, and MITA 3.0 compliance. The two major components of the Interoperability Platform include an Enterprise Service Bus (ESB) and a eHealth Exchange Gateway. The DOM Interoperability Platform subproject will be interoperable with the other two subprojects.

The MCI To-Be Environment includes connectivity to additional trading partners, including providers, payers, State Agencies, and other stakeholders, in coordination with the To-Be environment as outlined in the MS-HIN and MCI To-Be sections.

### 4.4 Future Alignment with MITA

As noted in Section 4.2 above, the State of Mississippi is currently in the process of procuring an MES to upgrade the systems and services that meet the business needs of the DOM. The ultimate goal is to define an Enterprise Architecture encompassing all Medicaid systems for the State of Mississippi that aligns to and advances increasingly in MITA maturity for business, architecture, and data and that includes MITA 3.0 standards, such as SOA using ESB infrastructure.

The new MES will interface with the DOM CDIP to allow for data interoperability between the MES and the CDIP subprojects, such as the MPI and CDR. This interoperability between the clinical and administrative systems will allow DOM to advance towards MITA 3.0, and specifically the care coordination components of MITA 3.0.

The MITA-enabling guidelines, processes, and tools provide a framework for the continuous improvement of service delivery and business processes based on efficient technology utilization. The MITA framework depicts this evolution as a progression of maturity levels that reflect DOM’s ability to execute business functions in the rapidly changing health care environment. DOM will use the MITA framework as a tool to assist in the strategic application of technology and enhancements that provide value and contribute to a continuous improvement in the Medicaid program’s maturity.

The selected MES vendor will employ SOA to take advantage of system components reuse across business functions as services. SOA is an approach to loosely coupled, protocol independent, standards-based distributed computing where software resources expose their functionality as services and are available on the network. SOA requires the use of business services in addition to technical services. The business services support business functions within the MES and map all applicable MITA business processes within the MITA Business Process Model, unless they are Mississippi-specific business processes. Each business service must meet the MITA definition of a business service. The SOA architecture must also enable the agency business units to build business applications quickly and efficiently in the future by reusing resident SOA infrastructure and application service components.

CMS requires a MITA roadmap that delineates how the proposed system enhancements for eligibility and enrollment functions will fit into the states’ greater MITA framework. Such a
requirement will align CMS’ expectations to see states continuing to make measurable progress in implementing their MITA roadmaps.

DOM has completed remediation of the eligibility system under amendment to the existing contract with the current fiscal agent. DOM has retired the MEDS and MEDSX systems with a new rules based system, Modernized MEDS (MMEDS), that determines MAGI based eligibility. DOM is now developing a rules based system that will combine MAGI and Non-MAGI eligibility determinations into one system. This system will be integrated to use the Federal Data Services Hub for needed verifications and referrals.

DOM’s roadmap will be aligned with MITA maturity target levels as follows:

- **As-Is:**
  - State Medicaid Agency complies with State regulations to maintain an adequate Provider network and pay claims promptly to encourage Provider participation and ensure access to care;
  - Many steps require manual intervention;
  - Data Content is nonstandard; and
  - Appropriateness of care is assessed retrospectively.

- **Target MITA Maturity Levels 3 & 4 (5 years):**
  - State Medicaid Agency coordinates with other payers to offer one-stop shop entry points to applicants for service and provider enrollment, provider reimbursement, and coordination of benefits;
  - Patients make personal healthcare decisions;
  - State Medicaid Agency accommodates cultural, linguistic, and health needs;
  - Clinical and Administrative systems (MES and DOM CDIP) interoperate and share data for improved, and where possible, automated decision making for improved care coordination;
  - State Medicaid Agency uses national standards for data content and exchange; and
  - Coordination and collaboration across healthcare programs intrastate contributes to improved outcomes.

The SOA will feature:

- **Technology Independence**: The service components will be invoked from multiple platforms and utilize standard protocols.

- **Standards-Based Interoperability**: The system will support multiple industry standards, including, at a minimum: HL7; XML; Extensible Stylesheet Language Transformation (XSLT); Web Services Interoperability (WS-I); WSDL; SOAP1.1 or
2.0; Universal Description, Discovery, and Integration (UDDI); Web Services (WS)-BPEL; Representational State Transfer (REST) (in place of SOAP); and WS-Message Transmission Optimization Mechanism (MTOM) Policy.

- **Life Cycle Independence**: Each service component will operate in a separate life cycle.

- **Loose Coupling**: Service components will be defined independently, with the interface components bridging the gap between components. For example, the Service Consumer Component specification must be defined independent of the Service Provider Component. The alignment of the two specifications is defined in the interface component.

- **Invocable Interfaces**: The Service interfaces will be invoked locally or remotely.

- **Communication Protocol**: A Service will be invoked by multiple protocols. The choice of protocol must not restrict the behavior of the service. Binding to a specific protocol will take place at run-time/deployment-time, and not at the design or development time.

- **Flexibility**: The selected vendor will focus on the business processes that comprise the systems, with the following in mind:
  - Ability to adapt applications to changing technologies;
  - Easily integrate applications with other systems;
  - Leverage existing investments in desired legacy applications; and
  - Quickly and easily create a business process from existing services.

- **Metadata Management**: SOA commonly provides application and data integration via an abstraction layer. Given the requirements of interoperability and independence, the proper use and management of metadata is extremely important to the effective operation of the SOA. It will also allow for:
  - Separation of the data and structures and convert them to a data layer within the SOA architecture;
  - Development of a Common Data Model and Metadata using the MITA HL7 methodology; and
  - Achievement of the SOA loosely coupled “separation of concern” approach, by separating the data layer from the application layer to more effectively and easily manage the data without changing the application code. This will create the desired more loosely coupled SOA environment and enable the business to accelerate any system changes required in the future.

- **ESB**: The MES will include an ESB for data transport, messaging, queuing, and transformation. The ESB is a service layer that provides the capability for services to interoperate and to be invoked as a chain of simple services that perform a more complex end-to-end process. The service layer is designed to handle both normal conditions and respond to failures and adapt to changes.
• **MITA Alignment**: The MES will be aligned with MITA. This includes, but is not limited to:
  - Map of business processes to MITA business processes;
  - Alignment of proposed business processes to the MITA maturity level and capabilities;
  - Use of MITA standard interface definitions (expressed in WSDL) and messages (expressed as an XML/schema) for all services;
  - Use of the MITA/HL7 methodology for defining the information model and messages; and
  - Adherence to the MITA governance process for newly developed interfaces and messages.

Because DOM and the MS Department of Human Services (MDHS) have a great deal of overlap in the communities they serve, they have long shared a joint-vision to improve collaboration and introduce technology and programmatic solutions to improve client services. Working together, the agencies explored improved interoperability and integration in technology, business process and workflows, case management, privacy, security, analytics/business intelligence and proper governance across health and human services systems to support integrated and coordinated services. As a result of the planning effort, DOM and MDHS have jointly launched a new phase of eligibility and enrollment enhancement, the Health and Human Services Transformation Project or “HHSTP.” Federal funding for HHSTP is not within the HIT IAPD, but has been approved by CMS and Food and Nutrition Services (FNS) via an Eligibility and Enrollment IAPD.

DOM is developing an RFP to procure services to perform a revised State Self-Assessment (SS-A) using the new MITA 3.0 guidelines. DOM will update this section, as appropriate, in a subsequent SMHPU.

### 4.5 Future Broadband Initiatives

As described in Section 3.7 – Current Broadband Initiatives, Mississippi has received funding to expand statewide broadband services. Utilizing these funding sources, MBCC continues to move towards implementing broadband expansion using the strategies outlined in their long-term strategic plan, “Mapping Mississippi’s Digital Future.” As a part of this effort, MBCC has launched the Extension Broadband Education and Adoption Team (e-BEAT), which deployed Regional Coordinators throughout Mississippi to work with elected officials, businesses, educators and community leaders on developing tools to increase digital literacy and increase broadband adoption. For example, e-BEAT is currently working on developing a map of broadband availability for inclusion in a comprehensive plan aimed at moving Mississippi towards greater access.

In addition to the ARRA broadband funding for expansion of broadband services, the State of Mississippi continues to participate in broadband connectivity expansion specifically for telehealth initiatives through the Federal Communications Commission (FCC) funding of the University of Mississippi Medical Center (UMMC). UMMC also received a United States Department of Health and Human Services (HHS) Health Resources and Services Administration
The State of Mississippi Health IT Committee Recommendations for Broadband include:

1. Partnership of the Mississippi Broadband Connection Coalition with the MS-HIN Board to coordinate growth and identify regulatory barriers to health IT adoption. An outcome of this partnership may be to form a sustainable public-private partnership with MS-HIN to support policy development in the field. This partnership could document Mississippi’s efforts in EHRs, Health Exchanges, Telemedicine, and Medical Record Imaging. An additional function could be to identify regulatory obstacles that may be inhibiting expansion of Health IT.

2. Attention to privacy and security concerns, including establishing a NPI system for all participants. The Health Information Technology Policy Committee (HITPC) report can serve as a guide for establishing Health IT growth policy at the state level.

3. Identification of a dedicated spectrum for medical imaging. High costs are associated to medical imaging from the limited supply of spectrum, however, the medical cost savings that could be realized through utilization of this technology in clinical and preventative practices makes the effort to find spectrum important. Once spectrum is found and financed, it could be dedicated to use by hospitals or rural physicians, and managed centrally.

4. Map availability of broadband to hospitals and rural physician groups. Hospitals should be at the top of the list for access to high speed Internet. To accomplish this, existing advocacy groups should unite to prioritize needs for a State Level Rural Health Care application. The first step should be to map the availability of broadband to the State’s hospitals.

5. Provision of Health IT-related digital literary courses at rural hospitals by Mississippi State University Extension Service eBEAT Team. National and state research suggests that geographic location is closely correlated with adoption rates. The challenge is how to introduce citizens who may already be marginalized from broadband usage to the concept of receiving healthcare from the Internet.

Per the 2017 Environmental Scan, Provider access to Broadband is not an issue, however, DOM will continue to monitor Broadband access and connectivity issues via the DOM outreach personnel and other DOM Provider-facing personnel.

4.6 Future Vision for Medicare and Federally-Funded, State-Based Programs

4.6.1 Medicare

As Medicare and CMS are migrating towards utilizing standards such as the eHealth Exchange network (Sequoia), it is essential for Mississippi to have the potential for eHealth Exchange connectivity with Medicare and CMS. Therefore, DOM’s existing Interoperability Platform
supports a variety of communication and interoperability standards and protocols, including eHealth Exchange to enable the potential for connectivity with CMS/Medicare/CMS Agencies for both clinical and administrative transactions. DOM plans to utilize the Interoperability Platform to facilitate connectivity through MS-HIN as the preferred connectivity methodology, and then by the MS-HIN eHealth Exchange Gateway to CMS. Coordination and planning with MS-HIN is ongoing to ensure a non-duplication of efforts, as described in the To-Be CDIP section.

4.6.2 CDC Coordination

A national initiative of CDC is to facilitate real time, interoperable data exchange between organizations for the promotion of collaboration and rapid dissemination of critical information in the organizations associated with public health. The integration and alignment of DOM with the State of Mississippi, including Public Health, for Public Health related reporting and surveillance to the CDC is important to improving health care outcomes for all Mississippians. DOM will consider implementing the GIPSE profile and other CDC-based reporting formats for interoperable data exchange with CDC using connectivity to MS-HIN and eHealth Exchange, including clinical and required (immunizations, syndromic surveillance, etc.) reporting. DOM is working with MSDH and MS-HIN to collaborate on standards-based connectivity and interoperability to facilitate reporting to MSDH and to further assist MSDH in reporting to the CDC, including using such standards as GIPSE and eHealth Exchange.

4.6.3 CMS/ASPE Coordination

Integration with CMS will enable electronic quality reporting via the eHealth Exchange connection, as ordered by the ARRA. Based on the recommendation of ONC, DOM is migrating toward utilizing Federal Health Architecture (FHA) standards via the DOM Interoperability Platform to coordinate with Medicare and federally funded, inter/intra-state based programs as they become compliant with FHA standards. By implementing and integrating standards, profiles, and interoperable infrastructure/technologies (including IHE, Healthcare Information Technology Standards Panel (HITSP), and eHealth Exchange standards, profiles, and technologies through the DOM Interoperability Platform, DOM will drive towards and migrate upwards to the higher levels of MITA and MITA compliance, as well as administrative simplification. DOM intends to report any required quality data to CMS, such as QRDA (via coordination and connectivity with the statewide HIE, MS-HIN). Accordingly, DOM has implemented and plans to continue to incorporate standards, profiles, and interoperable infrastructure such as IHE, HITSP and eHealth Exchange.

4.6.4 HRSA Coordination

HRSA is the primary federal agency for improving access to health care services for low income and uninsured individuals. The CFHC in Biloxi received a HRSA grant to connect 21 FQHC’s in Mississippi together for the exchange of health care data. These FQHCs have been connected together via an Allscripts cloud-based EHR. To date, the CFHC has not received any additional HIT grants. Lessons learned in the CFHC study can be used to encourage EHR adoption in other Mississippi FQHCs. DOM, via connectivity to the Delta Health Alliance (DHA) Allscripts EHR integration will support connectivity and clinical data interoperability for C-CDA clinical data
exchange with multiple FQHCs, as described in the MCI As-Is Section (EHR Integrations) as well as the As-Is FQHC Coordination Section of this document.

4.7 Future Vision for the Statewide Health Information Exchange

DOM is planning to support bi-directional clinical data exchange with MS-HIN, the State of Mississippi HIE, as outlined in the MS-HIN subproject (in the To-Be Section of this document), as a part of the overall CDIP. DOM has implemented the DOM Interoperability Platform as a single connectivity methodology to allow for MS-HIN connectivity to support bi-directional Medicaid-specific clinical data exchange. The DOM Interoperability Platform will provide connectivity and interoperability between the internal DOM systems and services (such as the DOM MCI), and provide a standards-based DOM eHealth Exchange to MS-HIN eHealth Exchange connection. This single connection to MS-HIN will help facilitate DOM’s connectivity needs to outside agencies, stakeholders, other States, other HIEs, and Federal Agencies.

DOM has identified several use cases that the DOM to MS-HIN connectivity model can support with To-Be infrastructure as outlined in the MS-HIN To-Be Subproject section of this document, including:

- Medicaid Beneficiary ADT Feed interoperability with MS-HIN to support harmonization between the existing DOM Master Patient Index (MPI) and the To-Be MS-HIN MPI;
- Medicaid Beneficiary clinical data exchange with MS-HIN and MS-HIN connected Medicaid providers in the form of a Medicaid C-CDA patient summary document (XML format) utilizing the MS-HIN To-Be ESB and the existing DOM Interoperability Platform.

4.7.1 DOM Agency-wide Enterprise Master Patient Index (eMPI)

DOM is planning to deploy an Agency-wide (Source of Truth) Enterprise Master Patient Index (eMPI) for Master Data Management (MDM) to provide patient matching and coordination of patient records and clinical data throughout DOM and across the DOM infrastructure, including for connectivity and interoperability with external stakeholders, State Agencies, and MS-HIN. The DOM CDIP and the future MES will each utilize their own existing, specific matching system for Medicaid Beneficiary matching based upon their specific rules and logic, as well as identity management. Therefore, it is critical to have a single, master ‘source of truth’ patient identifier for DOM beneficiaries via an Enterprise Master Patient Index (eMPI) for Master Data Management (MDM) to support systems and programs without an MPI as well as to coordinate identities between the CDIP and MES MPIs, and across the Medicaid Agency.

The DOM eMPI will allow for a limitation of duplicate beneficiary records, duplicate beneficiary clinical data and administrative data, and allow for more structure in the organization and storage of beneficiary data across the DOM infrastructure (including multiple clinical and administrative systems). Systems that would interface and utilize the DOM eMPI include the new MES, the DOM CDIP, and other various services and systems.
4.7.2 MS-HIN Governance

The structure for MS-HIN is set forth in Miss. Code Ann. §§ 41-119-1, et seq., entitled Health Information Technology Act, included as Appendix F. The governing body is the MS-HIN Board of Directors. The Board of Directors was appointed at the end of September 2010 and the first meeting was held on October 20, 2010. The figure below shows the overall structure for MS-HIN.

![Figure 145: MS-HIN Organization Structure](image)

The MS-HIN Board of Directors adopted the following statement to describe its vision for the HIE in Mississippi.

“The trusted source for secure, quality healthcare information – anywhere, anytime – for a healthier Mississippi.”

In addition, the Board adopted the following mission statement for HIE in Mississippi.

“To provide sustainable, trusted exchange of health information to improve the quality, safety, and efficiency of healthcare for all Mississipians.”

The MS-HIN Board of Directors maintains oversight responsibility for all HIE activities in the State of Mississippi. MS-HIN has a broad representation of stakeholders. DOM is a member of the MS-HIN Board of Directors and will work in partnership with the MS-HIN, providing leadership, as appropriate, to assure that Medicaid beneficiaries are best represented and served by the MS-HIN. In addition, Mississippi ITS staff members work directly with the MS-HIN and are specifically chartered to ensure that MS-HIN is compliant with the State of Mississippi’s laws and policies.

DOM will work closely with MS-HIN to ensure that each system supports broad, standards-based, interoperable environments to maximize DOM’s investments in these efforts. Having this standards-based foundation allows DOM the greatest flexibility moving forward.
Policy development, including providing advice and counsel, is a function of the MS-HIN Board of Directors. The MS-HIN requires a majority of the total membership to approve all policy decisions. MS-HIN may form special advisory groups on an as-needed basis to address specific issues of importance.

The State HIT Coordinator is not a member of the MS-HIN Board, but coordinates the MS-HIN Board meetings and acts as a liaison between ONC and the MS-HIN Board. The State HIT Coordinator also works closely with the senior staff at DOM to coordinate activities across a wide range of issues.

4.8 Future Vision for the Public Health Initiatives

DOM is planning to utilize the DOM Interoperability Platform to connect to the MSDH, via a connection to MS-HIN, for such use cases as:

- Bi-directional Medicaid immunization data exchange between the MSDH MIIX and DOM;
- Medicaid admissions, discharge, transfer (ADT) Feeds from MSDH (if available) to DOM;
- Interoperability with the MSDH Patient Centered Medical Home (PCMH) and other such MSDH Programs.

4.9 Future Vision for Federally Qualified Health Centers/Rural Health Clinics

FQHCs and RHCs are already working together and exchanging health care information. A project connecting 14 of 21 FQHCs is already in place. The CFHC in Biloxi, Mississippi has connected all 21 locations via an Allscripts cloud-based EHR.

The Delta Health Alliance in Greenville, Mississippi received a Beacon Community Grant and has connected all the RHCs in the 18-county Delta region of the State via an Allscripts cloud-based EHR. DOM, via connectivity to the Delta Health Alliance (DHA) Allscripts EHR integration will support connectivity and clinical data interoperability for C-CDA clinical data exchange with multiple FQHCs, as described in the MCI As-Is Section (EHR Integrations) as well as the As-Is FQHC Coordination Section of this document. The goal for this project is to go live by December 2017.

4.10 Future Vision for DoD and VA

There are three major military installations in the State of Mississippi: two are Air Force bases near Columbus and Biloxi and the third is a Navy facility near Meridian. While the military has expressed an interest in receiving information about off-base treatment of military personnel, they have been unable to connect to the State to retrieve the information due to severely restrictive security constraints.
In addition to the two large Veterans hospital facilities in Mississippi – one in Biloxi and one in Jackson, the Board of Veterans Affairs is located in Jackson, Mississippi.

The DoD and the Veterans Administration (VA) are currently running the VLER EHR, however, recent developments have the DoD and VA moving away from VLER to a different EHR system. The DoD and the VA are currently using the VLER, however, they may be migrating to a new, commercial EHR. VLER and many commercial EHRs both support future connections to MS-HIN, and subsequently DOM, via the eHealth Exchange. By connecting to the VA and DoD, DOM can exchange clinical data and documents with the VA and DoD and coordinate care for the active duty military personnel or veterans, if need be. DOM will continue to examine DoD and VA use-cases and coordination of clinical data and care coordination.

4.11 Future Vision for Indian Health Services

Choctaw Indians are the most prevalent minority of the American Indian population in the State of Mississippi. Members of the Mississippi Indian Tribe receive basic health care through a community health service. Representatives of the Tribe indicate they are participating with Indian Health Services and can connect to DOM through MS-HIN, via the DOM to MS-HIN eHealth Exchange connectivity. Therefore, the exchange of health care information can be accomplished through MS-HIN and by connecting with Indian Health Services using secure protocols and standards.
5 Provider Incentive Program Blueprint

5.1 Introduction

5.1.1 Overview

The Medicaid EHR Incentive Program, defined by the ARRA of 2009, was established to provide incentive payments to eligible providers for their efforts to meaningfully use certified EHR technology, including adoption, implementation, or upgrade (A/I/U). Through August, 2017, DOM has paid $207,746,612 in incentive payments to 3,127 unique eligible professionals (EPs) and 98 eligible hospitals (EHs) for attesting to A/I/U or Meaningful Use (MU).

This Provider Incentive Program Blueprint (Blueprint) describes the high-level requirements, process flows, and technical requirements of the Mississippi Provider Incentive Program (MPIP) to interface with the CMS Registration & Attestation System to enable providers to register for Medicaid incentives, attest to their eligibility requirements in each year of the program, and allow DOM to pay incentive payments in 2018 and subsequent years. The software application supporting the MPIP is the Conduent solution, currently being offered to multiple states as a software as a service (SaaS) solution. DOM's decision to implement a SaaS solution has helped the MPIP leverage resources across the participating states.

DOM has branded the Conduent solution as the Mississippi State Level Registry (MS SLR) to be specific to the MPIP and DOM policies.

This Blueprint has liberally borrowed from efforts in other states and documentation from CMS.

5.1.2 Purpose

The purpose of this program is to capture and track provider attestations, evaluate eligibility, and collect information in order to make timely incentive payments to qualifying providers for A/I/U and MU of certified EHR technology. The goal of the program is to ensure the right payment was made to the right provider at the right time.

The MS SLR has interfaced with the CMS Registration & Attestation System and is configured to capture and document information regarding the following:

- Eligibility history;
- Payment history;
- Audit (implemented in 2013);
- Appeals (implemented in 2013); and
- Recoupment and/or Adjustment (implemented in 2013).

DOM utilizes the MS SLR for storing, tracking and reporting on attestation data including all the information listed above.
Figure 16 depicts the high level overview of the necessary components of the MPIP:

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<tr>
<th>Medicaid Providers</th>
<th>CMS</th>
<th>Medicaid</th>
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<tbody>
<tr>
<td>Provider NLR Registration</td>
<td>National Level Repository (NLR)</td>
<td>CMS NLR Registration Module</td>
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<tr>
<td>Provider SLR Registration</td>
<td>NLR Registration Status</td>
<td>NLR – States, Provider Registration Data Interface (B-6)</td>
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<tr>
<td>Provider Attestation</td>
<td>States – NLR Registration Confirmation Data Interface (B-7)</td>
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<tr>
<td>Medicaid Provider Incentive Program Solution</td>
<td>States – NLR, Final Registration Confirmation Data Interface (B-7)</td>
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<td>Provider Web or PIP Support Inquiries</td>
<td>MS SLR Eligibility Status</td>
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<td>Provider Appeals</td>
<td>Dually Eligible Hospital Attestation Data (C-8)</td>
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<td>Provider Audits</td>
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<td>Medicaid Payment Request Response Interface (D-16)</td>
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Figure 156: Mississippi Provider Incentive Program Solution
5.2 Eligibility: Provider Type, Eligibility Period, and Patient Volume

Providers must meet the eligibility requirements for provider type (EP or EH) and patient volume to receive EHR Incentive Payments.

5.2.1 EH Eligibility Criteria

EHs must meet the following criteria for the EHR Incentive Payment program. Please note that criteria have been updated to reflect changes to eligibility as stated in the CMS Stage 2 Final Rule (2012).

5.2.1.1 EH Provider Type

To be eligible for the MPIP, EHs must fall into one of the following hospital types:

- **Acute Care Hospital:**
  - The CCN has the last four digits in the series 0001 – 0879; and
  - The average length of patient stay is 25 days or fewer; or

- **Critical Access Hospital (CAH):**
  - The CCN has the last four digits in the series 1300 – 1399; and
  - The average length of patient stay is 25 days or fewer; or

- **Children’s Hospital: (None in Mississippi)**
  - The hospital is separately certified as a children’s hospital - either freestanding or a hospital within hospital and the CCN has the last four digits in the series 3300-3399; or
  - The hospital is separately certified, either freestanding or hospital within a hospital, which predominately treats individuals 21 years of age or younger and does not have a CCN because they do not serve any Medicare beneficiaries but has been provided an alternative number by CMS for purposes of enrollment in the Medicaid EHR Incentive Program.

5.2.1.2 EH Eligibility Period

For the purposes of calculating hospital patient volume the eligibility period is defined as:

- A representative, continuous 90-day, 3-month, 6-month or full year period from the preceding fiscal year; or

- A representative, continuous 90-day period in the 12-month period directly preceding the attestation date.
DOM requires that the eligibility period start on the first day of the month to ensure that patient volume data self-reported in the eligibility period selected by the provider aligns with the reporting periods of the data available in the MMIS. Once an eligibility period is used for the purposes of calculating Medicaid patient volume, the same eligibility period may not be used in subsequent attestation years for the purposes of proving Medicaid patient volume.

5.2.1.3 EH Patient Volume

Acute Care and CAHs must have at least a 10 percent Medicaid patient volume based on both the inpatient and emergency room discharges. Children’s hospitals are not required to meet a minimum Medicaid patient volume. To calculate Medicaid patient volume, an EH must divide total Medicaid encounters (numerator) by total patient encounters (denominator) using the same eligibility period for both numerator and denominator.

For purposes of calculating hospital patient volume, a Medicaid encounter means services rendered to an individual per inpatient discharge and/or in an emergency department on any one day where:

- Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) paid for part or all of the service; or
- Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) paid for all or part of the individual’s premiums, co-payments, and/or cost sharing; or
- The individual was enrolled in a Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act), regardless of payment liability, in accordance with CFR §495.306.

As noted above, the optional EHR Hospital Patient Volume Calculator can be found at [http://msehrpip.wordpress.com](http://msehrpip.wordpress.com). Also, see Appendix G attached hereto. Hospitals may use the EHR Hospital Patient Volume Calculator as a worksheet; however it will no longer be required for submission with the attestation.

Hospitals are allowed to count a maximum of one encounter per patient per day. Hospitals will be required to use their discharges from both the inpatient facility (POS 21) and the emergency room (POS 23) to determine their patient volumes.

The authorized data source documents (detailed below) are required documentation to be submitted with EH attestations. Only MS DOM authorized data sources as described below will be used to calculate the Medicaid share percentage.

- The authorized data source for the total Inpatient Discharges (POS 21) will be the annual cost report for the hospital's fiscal year ending in the prior federal fiscal year.
- The authorized data source for the total Medicaid Primary Inpatient Discharges (POS 21) will be the annual cost report for the hospital's fiscal year ending in the prior federal fiscal year.
• The authorized data source for the total Medicaid Secondary Payer Inpatient Discharges will be the hospital's inpatient accounting/billing system. Only Medicare and Third party claims with Medicaid as the secondary payer showing that the individual was enrolled in Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) will be used to determine the Medicaid Secondary Payer Inpatient Discharges, regardless of payment liability by Medicaid. Summary data supporting each discharge amount will be attached to the hospital's application. Upon request, the hospital may be required to provide detailed reports including the payer (primary and secondary), patient ID, claim number, date, revenue and procedure codes, and paid amounts.

• The authorized data source for the total Medicaid Primary Payer Emergency Room Discharges will be the hospital's inpatient accounting/billing system. Summary data supporting each discharge amount will be attached to the hospital's application. Each emergency room visit will be considered a single discharge. Emergency room visits that result in transfer to the inpatient unit for other than observation will not be included in the emergency room discharges. Upon request, the hospital may be required to provide detailed reports including the payer (primary and secondary), patient ID, claim number, date, revenue and procedure codes, and paid amounts.

• The authorized data source for the total Medicaid Secondary Payer Emergency Room Discharges will be the hospital's emergency room accounting/billing system. Only Medicare and Third party claims with Medicaid as the secondary payer showing that the individual was enrolled in Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) will be used to determine the Medicaid Secondary Payer Emergency Room Discharges, regardless of payment liability by Medicaid. Medicare and Third party claims will be reported separately. Summary data supporting each discharge amount will be attached to the hospital's application. Upon request, the hospital may be required to provide detailed reports including the payer (primary and secondary), patient ID, claim number, date, revenue and procedure codes, and paid amounts. Each emergency room visit will be considered a single discharge. Emergency room visits that result in transfer to the inpatient unit for other than observation will not be included in the emergency room discharges.

As noted above, hospitals have the option to complete the EHR Hospital Patient Volume Calculator. The EHR Hospital Patient Volume Calculator will no longer be required for a hospital’s attestation but may be uploaded with the hospital’s attestation as needed. However, all other authorized data sources must be attached to the hospital’s attestation as supporting documentation.

5.2.2 EP Eligibility Criteria

Medicaid EPs must meet the following criteria to be eligible for the MPIP. Please note that criteria have been updated to reflect changes to eligibility as stated in the CMS Stage 2 Final Rule (2012).
5.2.2.1 EP Provider Type

To be eligible for attestation to the MPIP, EPs must be licensed as one of the following:

- Doctor of Medicine;
- Doctor of Osteopathy;
- Doctor of Dental Medicine or Surgery;
- Optometrist;
- Nurse Practitioner;
- Certified Nurse Mid-Wife; or
- Physician assistant (PA) when working at a Federally Qualified Health Clinic (FQHC) or Rural Health Clinic that is so led by a PA.

EPs working in a FQHC or RHC will be determined based on prior year claims history for “predominately” status. EPs with at least 50 percent of their encounters (claims) provided through or in a FQHC or RHC environment will qualify as working “predominately” in a FQHC or RHC. Professionals must also be currently performing services in a FQHC or RHC.

5.2.2.1.1 Physician Assistant Criteria

PAs are considered to be EPs if the PA is practicing predominately in an FQHC or RHC that is “so led” by a PA. An FQHC or RHC is considered to be “so led” under the following circumstances:

- A PA is the primary provider in a clinic (for example, when there is a part-time physician and full-time PA, the PA is the primary provider);
- A PA is a clinical or medical director at a clinical site of practice; or
- A PA is an owner of an RHC.

A PA practicing predominately in a FQHC or RHC is eligible to use Needy Individual patient volume. A PA is considered to be practicing predominately if over 50 percent of his or her total patient encounters over a period of six months in the most recent calendar year occur at a FQHC or RHC.

5.2.2.1.2 Pediatricians

Pediatricians must be board certified or board eligible and must have the appropriate taxonomy code in the MS SLR Provider Master File (PMF). Pediatricians may qualify for a reduced payment if they have greater than 20 percent Medicaid patient volume, but less than 30 percent Medicaid patient volume. Pediatricians may receive the full incentive payment amount if they can demonstrate 30 percent Medicaid patient volume in a given program year. Pediatricians working in an FQHC or RHC that choose to use Needy Individual patient volume must have at least 30 percent Needy Individual patient volume.
5.2.2.1.3 Hospital Based EPs

Hospital based EPs are determined on the EP’s services provided in service code areas 21 and 23. In accordance with the CMS Stage 2 Final Rule (2012), hospital based EPs are now eligible to attest for individual incentive payments if they can demonstrate that they have funded, acquired, implemented and maintained certified EHR technology, including supporting hardware and any interface necessary to meet MU, without reimbursement from an EH or CAH.

EPs will be deemed to be hospital based if 90 percent or more of total Medicaid encounters are provided in service code areas POS 21 and POS 23. Total Medicaid encounters include both Medicaid and Medicaid Managed Care encounters. The formula for the computation will be 

\[(\text{Total Medicaid encounters provided in service code areas POS 21 and POS 23}) / (\text{Total Medicaid encounters for all areas})\].

The MS SLR assists DOM in identifying non-hospital based EPs by requiring that EPs attest to the fact that they do not perform greater than 90 percent of their services in an inpatient or emergency room setting.

5.2.2.2 EP Eligibility Period

For all program years, EPs may use an eligibility period that falls under the following criteria:

- A 90-day period, 3-month period, 6-month period or a full year period from the preceding calendar year; or
- A 90-day period from the 12-month period directly preceding the EP’s attestation date.

The length of the period will be identified during attestation in the MS SLR. The numerator and denominator of the Medicaid patient volume equation must use the same eligibility period. Once an eligibility period is used for the purposes of calculating Medicaid patient volume, the same eligibility period may not be used in subsequent attestation years for the purposes of proving Medicaid patient volume.

DOM requires that the eligibility period start on the first day of the month to ensure that self-reported patient volume data in the eligibility period selected by the provider aligns with the reporting periods of the data available in the MMIS.

5.2.2.3 EP Patient Volume

DOM opted to offer the Medicaid fee for service (standard) calculation for EP Medicaid patient volume. Patient volume can be aggregated from multiple locations or states.

EPs must demonstrate at least 30 percent Medicaid patient volume based on Medicaid encounters and total encounters during a chosen eligibility period. To calculate Medicaid patient volume, an EP must divide total Medicaid encounters (numerator) by total patient encounters (denominator) using the same eligibility period for both the numerator and denominator. An encounter includes concurrent care or transfer of care visits, consultant visits, or prolonged physician service without direct (face to face) patient contact (telehealth),
regardless of financial liability. Providers are allowed to count a maximum of one encounter per recipient per day. No financial obligation is necessary for encounters to be included in Medicaid patient volume calculations.

For purposes of calculating patient volume a Medicaid encounter is defined as services rendered to an individual on any one day where:

- Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) paid for part or all of the service; or
- Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act) paid for all or part of the individual’s premiums, co-payments, and/or cost sharing; or
- The individual was enrolled in Medicaid (or a Medicaid demonstration project approved under section 1115 of the Act), regardless of payment liability, in accordance with CFR §495.306.

The EHR Professional Patient Volume Calculator can be found at Provider Resources (Calculators, Security, etc...). There are two versions of the EHR Professional Patient Volume Calculator, one for EPs using Medicaid patient volume only and the other for EPs practicing in FQHCs, RHCs, and IHS. Also, see Appendix G attached hereto. A copy of the EHR Professional Patient Volume Calculator may be attached with the MS SLR application as optional supporting documentation.

All providers are required to attach summary reports from their practice management or billing systems supporting their encounter calculations for their online application. Summary reports must separate the eligible encounters by the primary and secondary payer. Managed Care patient encounters must be identifiable in the Medicaid and all payer encounter counts. DOM will verify that all providers have attached this required documentation with applications submitted.

All Medicaid encounter counts are compared to the provider’s practice management or billing reports (regardless of financial obligation) for verification of encounters claimed on their application. Both the total and Medicaid primary and secondary encounters are verified. Medicaid claim counts are available in the MS MMIS as a secondary source of verification or Medicaid encounters.

The MS SLR provides for statistical data to be entered by State and can accept multiple states. Mississippi Medicaid encounters will be compared to the EP’s and/or Group’s claims data for the appropriate period of time. Out of state claims data will be subject to written verification from the other state at the option of the DOM audit staff. All applications are subject to both prepayment and post-payment audits.

**5.2.2.3.1 Needy Individual Patient Volume**

EPs practicing predominately in a FQHC or RHC may choose to use Needy Individual Patient volume in lieu of Medicaid patient volume for the purposes of meeting the 30 percent threshold. Needy Individual patient volume is calculated using the following formula:

\[
((\text{Needy Individual Patient Encounters} + \text{Medicaid Encounters})/\text{Total Patient Encounters}) \times 100 = n\
\]
To be considered a Needy Individual patient, a patient must meet one of the following criteria:

- Receives medical assistance from Medicaid;
- Receives medical assistance from the Children’s Health Insurance Program;
- Receives uncompensated care by the Provider; or
- Receives services at either no cost or reduced cost based on a sliding scale determined by the individual’s ability to pay.

5.2.2.3.2 MississippiCan

Because MississippiCan was initiated in 2011, applications can include encounters for Managed Care patients in the eligible professional encounters. Managed Care Encounters must be included in the numerator and denominator during attestation in the MS SLR. Additionally, encounters for Managed Care patients should be shown on a separate line in the EHR Professional Patient Volume Calculator (if included in the attestation documentation).

5.2.2.3.3 Group Medicaid Patient Volume

EPs may opt to use Group patient volume as proxy for their individual patient volume. An EP may use Group patient volume as a proxy for their own under the following conditions:

- The clinic or group practice’s patient volume is appropriate as a patient volume methodology calculation for the EP (for example, if an EP only sees Medicare, commercial, or self-pay patients, this is not an appropriate calculation);
- There is an auditable data source to support the clinic or group practice’s patient volume determination; and
- The clinic or group practice and EPs decide to use one methodology in each year (in other words, clinics could not have some of the EPs using their individual patient volume for patients seen at the clinic, while others use the clinic-level data).

The clinic or group practice must use the entire clinic or group practice’s patient volume and not limit it in any way. EPs may attest to patient volume under the individual calculation or the clinic or group practice proxy in any participation year.

If the EP works in the clinic as well as outside the clinic (or with and outside a group practice), then the clinic or group practice level determination includes only those encounters associated with the clinic or group practice.

In order to meet the requirements to use Group patient volume, including the requirement of an auditable data source, Mississippi will require the clinic or group practice to include all servicing providers’ claims regardless of the payer or whether or not they are eligible for the incentive payment.
For purposes of calculating Group patient volume for EPs, the clinic or group should divide:

- The total eligible Medicaid encounters for all EPs in the clinic or group practice in the continuous 90-day period, 3 month period, 6 month period, or full year period, in the preceding fiscal year; or

- The total eligible Medicaid encounters in the clinic or group practice in the continuous 90-day period in the 12-month period directly preceding the attestation date; by

- The total encounters for the clinic or group practice for all servicing providers not limited in any way for the same eligibility period.

For Mississippi, a Group will be defined as having the same NPI, TIN and Payee Medicaid ID. All individual EPs and clinics or group practices must be registered with the DOM with a current license, must be in good standing with CMS, the DOM, and the State of Mississippi and must have an NPI and Mississippi Medicaid provider number. Both the individual EP and Group must have an active status in the DOM PMF, including active licenses, and all individual EP’s seeking an EHR incentive payment which is assigned to the Group must dual-affiliation with the Payee Group in the MMIS.

5.3 Provider Registration and Verification

5.3.1 CMS Registration & Attestation System Registration

CMS has ownership of all processes concerning registration at the national level. A brief description is provided here. More detailed information can be found in the document entitled “HITECH Interface Control Document.” The most important aspect of the registration process for the MPIP concerns the interface transaction sent from the CMS Registration & Attestation System to the MS SLR once a provider has registered with CMS. More detail on this interface is contained in this Blueprint in Section 5.2.2.1 – CMS Registration & Attestation System – States, Provider Registration Data Interface (B-6) Process.

Regardless of the provider’s intent to attest with the Medicare or Medicaid EHR Incentive Program, all providers applying for incentives must first register with CMS Registration and Attestation System. The CMS Registration and Attestation System will capture basic information such as provider type (EP or EH) and whether the provider is applying for Medicare, Medicaid, or both (allowed for certain EHs). To eliminate duplication, CMS has restricted EPs to a single Web account that requires EPs to use their Social Security Number (SSN)/Tax Identification Number (TIN) to establish their registration and has restricted the issuance of the Web accounts to one per SSN/TIN.
If a provider chooses Medicaid, or both Medicaid and Medicare (EHs only), the provider must identify the state selected for attestation. The CMS Registration and Attestation System will check for a valid National Provider Identifier (NPI), TIN (if on record), and for any federal level sanctions. For EHs only, the CMS Registration and Attestation System will also check for a valid CMS Certification Number (CCN). Providers opting for Medicaid who are not included in the Social Security Administration (SSA) Death Master File will be passed through to the Medicaid state selected by the provider. If registration checks complete successfully, the new provider information will be written to the CMS Registration & Attestation System and sent to the State for validation in a data transaction defined by CMS named the “CMS Registration & Attestation System – States Provider Registration Data Interface (B-6).”

Hospitals registering for both the Medicaid and Medicare EHR Incentive Program at the same time that are approved by CMS as a meaningful user will also be deemed a meaningful user by Medicaid. The CMS Registration & Attestation System will send a C-5 record to confirm that CMS has determined the hospital to be a meaningful user of EHR technology. The hospital must still submit their attestation to Medicaid in order to receive their Medicaid MU incentive payment. This is the recommended pathway for dually eligible hospitals that apply for an MU incentive payment.

The CMS Registration & Attestation System communicates the registration status back to the provider.

5.3.2 CMS Registration & Attestation System/MS SLR Data Validation Process

This process will accept and parse the B-6 Interface. The purpose of the B-6 Interface is to inform the states of new, updated, and inactivated Medicaid registrations. The CMS Registration & Attestation System will send batch feeds to the states of new EPs and EHs that registered for the EHR Incentive Program and selected or switched to Medicaid. The data also includes any updates/changes to the EP or EH entries and any registration inactivation events. A detailed description of this interface can be found in the document entitled “HITECH Interface Control Document.”

This process will perform the following actions:

- Accept new transactions;
- Handle duplicate transaction exception; and
- Send back the Provider Registration Confirmation Interface (B-7 Interface) immediately after the first time a B-6 Interface is received, parsed, and stored for a given provider. The B-7 Interface will contain an Eligibility Status of

---

3 Please note that the CCN was previously known as the Medicare Provider number.
“Pending” and allow CMS to record the fact the B-6 Interface was received by DOM before DOM determines the provider’s registration status with the State.

Processes to manage transactions that do not pass Exception Handling are not described because the HITECH Interface Control Document states that CMS does not expect any exceptions from the B-6 Interface.

If the transaction passes Exception Handling and Duplicate Check processing, the process named “CMS Registration & Attestation System/MS SLR Data Validation” (described in this section) is executed.

The CMS Registration & Attestation System/MS SLR data validation process supports the requirements that provider data in the B-6 Interface be verified by the provider. Process execution logic depends on several different scenarios:

- **NPI from a B-6 Interface transaction being processed does not match a MS SLR Provider Registration transaction:** The B-6 transaction is stored in the MS SLR awaiting MS SLR Provider Registration using the same NPI.

- **NPI from a B-6 Interface transaction being processed does match a MS SLR Provider Account transaction:** The data from the B-6 transaction is matched against the data input by the provider during MS SLR provider account creation.

- **NPI from a MS SLR Registration transaction being processed does not match a B-6 Interface transaction:** The MS SLR provider can create an account and can complete the “About You” step. The provider will receive a hard stop after the “About You” step and will be notified that he/she must complete his/her CMS Registration and Attestation System application before proceeding in the MS SLR. The receipt of the matching B-6 transaction will allow the provider to proceed in the MS SLR.

- **NPI from a MS SLR Registration transaction being processed does match a B-6 Interface transaction:** The data from the MS SLR Provider Registration is matched against the B-6 transaction. If all data matches, the provider can proceed with the completion of their attestation.

In the event that the information entered by the provider and transmitted through the B-6 Interface cannot be validated, the provider may be asked to correct information entered at the CMS Registration & Attestation System. The MS SLR will not allow any changes to the NPI, SSN, CCN or TIN entered at CMS Registration & Attestation System. If an EP or EH needs to change any of this information to proceed, the Help Desk staff will refer them to CMS Registration & Attestation System where the EP or EH will be responsible for correcting the information. Upon completion and update at the CMS Registration & Attestation System, the information will be sent to and incorporated in the MS SLR electronically as an update.

State Reason Codes received on the B-6 transaction will also be interrogated to determine if the provider eligibility should be rejected based on code values sent to the MS SLR from the CMS Registration & Attestation System. The following table lists the codes. The codes designated by a “Hard Stop” will cause the provider’s eligibility to be rejected. If the B-6 transaction includes
one of the “Soft Stop” codes, it means the provider’s eligibility was rejected by another state. This will not exclude the provider from being eligible in Mississippi. Normal eligibility determination processes will still be performed.

Table 5-1: State Reason Codes

<table>
<thead>
<tr>
<th>State Reason Code Description</th>
<th>Reason Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eligible Hospitals</td>
<td></td>
</tr>
<tr>
<td>Excluded / Federal</td>
<td>EH01</td>
</tr>
<tr>
<td>Excluded / State</td>
<td>EH02</td>
</tr>
<tr>
<td>Not Licensed / Credentialed</td>
<td>EH03</td>
</tr>
<tr>
<td>Failed Patient Volume</td>
<td>EH04</td>
</tr>
<tr>
<td>No Certified EHR</td>
<td>EH05</td>
</tr>
<tr>
<td>Failed A/I/U</td>
<td>EH06</td>
</tr>
<tr>
<td>Failed MU</td>
<td>EH07</td>
</tr>
<tr>
<td>Excluded / Federal / 2nd Check</td>
<td>EH08</td>
</tr>
<tr>
<td>Excluded / State / 2nd Check</td>
<td>EH09</td>
</tr>
<tr>
<td>Eligible Professionals</td>
<td></td>
</tr>
<tr>
<td>Excluded / Federal</td>
<td>EP01</td>
</tr>
<tr>
<td>Excluded / State</td>
<td>EP02</td>
</tr>
<tr>
<td>Dead</td>
<td>EP03</td>
</tr>
<tr>
<td>Not Licensed / Credentialed</td>
<td>EP04</td>
</tr>
<tr>
<td>Hospital Based</td>
<td>EP05</td>
</tr>
<tr>
<td>Failed Patient Volume</td>
<td>EP06</td>
</tr>
<tr>
<td>Failed Practices predominantly at a FQHC / RHC with 30% needy individual patient volume</td>
<td>EP07</td>
</tr>
<tr>
<td>No Certified EHR</td>
<td>EP08</td>
</tr>
<tr>
<td>Failed A/I/U</td>
<td>EP09</td>
</tr>
<tr>
<td>Failed MU</td>
<td>EP10</td>
</tr>
<tr>
<td>Excluded / Federal / 2nd Check</td>
<td>EP11</td>
</tr>
<tr>
<td>Excluded / State / 2nd Check</td>
<td>EP12</td>
</tr>
<tr>
<td>Dead / 2nd Check</td>
<td>EP13</td>
</tr>
</tbody>
</table>

The B-7 Interface will be sent back to the CMS Registration & Attestation System the second time as the Provider Final Registration Status Interface (B-7). At this time, the B-7 transaction will contain an Eligibility Status of “Accepted” or “Rejected” notifying the CMS Registration & Attestation System of the provider’s registration status with the MPIP. The rejection reason will be communicated back to the CMS Registration & Attestation System using one of several codes. Please refer to Table 5-1: State Reason Codes above. The Hard Stop/Soft Stop designation has no meaning in this context; they all signify that provider eligibility was rejected. Mississippi may use any of the State-specific codes to specify the reason the provider was rejected.
5.3.3 MPIP MS SLR Registration

The MS SLR registration process will only accept registration requests from Mississippi Medicaid Providers. A provider is considered a Mississippi Medicaid Provider if the provider has an active Mississippi Medicaid Provider number. Providers who work in an FQHC or a Coordinated Care Organization must also have a Mississippi Medicaid Provider number. Any provider who attempts to register in the MS SLR without a Medicaid Provider number will be prohibited by the application from proceeding with registration. DOM has emphasized the fact that the Medicaid Provider number is a requirement for eligibility in the MPIP training for providers.

This process supports provider registration with the MS SLR. The provider verifies information obtained via the CMS Registration & Attestation System interface and supplies additional information the State may require for determining eligibility before the attestation process. Areas of focus within the MS SLR for Mississippi registration and eligibility verification include:

- Mississippi Medicaid Provider number;
- Professional license number – for providers with licenses in multiple states, the MS MMIS will search for a Mississippi license, regardless of the number of other state licenses associated with a given provider;
- Provider type and any hospital, FQHC, or RHC affiliation; and
- Provider sanctions/exclusions; those checked at the State level by the MS SLR include terminated licenses, expired licenses, State terminations, deceased providers, legal actions, and voluntary terminations by the provider. Based on the CPI Informational Bulletin, CPI-B11-05, issued on 05/31/2011, Mississippi will not permit individuals or entities that are currently terminated or sanctioned under Medicare or any other State Medicaid program to apply for or receive payment.

A Provider Master File (PMF) is generated weekly from the MMIS and holds information on all EPs and EHS that are potentially eligible for the MPIP. This file is sent from the MMIS to the MS SLR each week. The MS SLR Registration Validation from the MMIS and PMF includes the following checklist:

- Provider and Payee NPI are valid;
- Provider is not deceased;
- Medicaid Provider number is valid, including clinic or group practice Medicaid Provider numbers;
- Providers have current licenses issued by the State of Mississippi;
- Provider is not sanctioned by Mississippi DOM; and
- Provider type is included in the attestation and is a valid code.
• Evidence of a previously paid Medicaid claim and date of last paid Medicaid claim.
5.4 MPIP MS SLR Attestation

Once registration is complete, the provider’s next step in applying for the MPIP is to access the MS SLR and answer a variety of questions attesting to the A/I/U or MU of certified EHR technology. EP and EH attestations are subject to eligibility verification processes as described.
in Section 5.2 above. As stated, DOM will verify this information using practice management reports for EPs as a part of required documentation to be attached to an attestation. EHs will be verified by a review of cost reports and data sources prior to payment.

DOM will continue using established lines of communication between the SMA and individual providers. Providers are notified via email and phone call when validation issues occur (e.g. missing or incorrect supporting documentation, incorrect data entered into the MS SLR, license expiration, payee affiliation issues, etc.) and when the pre-payment verification steps have been completed and that payment is forthcoming.

5.4.1 Adoption, Implementation, or Upgrade

Along with the attestation information described above for provider type, eligibility period, and patient volume, providers also may attest to the A/I/U of certified EHR technology in the first year. Providers must enter the CMS EHR Certification code from its EHR vendor to identify their EHR software. The MS SLR will validate the CMS EHR Certification code against the current ONC database of valid CMS EHR Certification codes. Please note that there is no EHR reporting period required for A/I/U attestations.

The definition of Adopt/Implement/Upgrade (A/I/U) in 42 CFR 495.302 allows a provider to demonstrate A/I/U through any of the following: (a) acquiring, purchasing or securing access to certified EHR technology; (b) installing or commencing utilization of certified EHR technology capable of meeting meaningful use requirements; or (c) expanding the available functionality of certified EHR technology capable of meeting meaningful use requirements at the practice site, including staffing, maintenance, and training, or upgrade from existing EHR technology to certified EHR technology per the EHR certification criteria published by ONC.

Program Year 2016 was the last year new participants could begin the MS Medicaid EHR Incentive Program. June 30, 2017 was the end of the MS EHR Attestation Submission collection season for Program Year 2016. After that date, no new providers would be allowed to join the program. This coincided with enhancements at the NLR which noted new registrations without previous participation years.

During the attestation process in the MS SLR, the provider is required to supply the following attestation information to qualify for an A/I/U incentive payment: (The following AIU description is for historical purposes only – detailing the process of first-year AIU attesters)

- Select Adoption, Implementation, or Upgrade;
- Provide a brief textual description of how the provider meets the criteria for Adoption, Implementation, or Upgrade of certified EHR technology;
- Attach external documents supporting Adoption, Implementation, or Upgrade of certified EHR technology. DOM prefers that a signed contract is uploaded demonstrating proof of a fiscal relationship between the vendor and the EP/EH. The Division of Medicaid expects the following components to be addressed within a vendor contract: Names or Vendor and Name of clinic; CEHRT product name; description; CEHRT ID; Version numbers; Business Associates Agreement;
End User Licensing Agreement; Dates of contract execution; contractual terms; contractual updates; nature and scope of updates; Number of end user licenses available; evidence that contract was in effect during specified periods; Issues relating to HIPAA Compliance; Etc... In instances in which a signed contract is not applicable DOM will accept other documentation, including but not limited to, a vendor invoice, an End-User License Agreement (EULA), or other evidence that sufficiently demonstrates A/I/U.

- Certified EHR Technology: Enter ONC certification code. CMS publishes a list of codes identifying all ONC certified EHR technology products. During attestation the provider must enter the code from its EHR vendor to identify the EHR or obtain the certification number from the current ONC CHPL list.

- Attestation Agreement: Sign and attach an Attestation Agreement indicating A/I/U. Attestation Agreement must be executed by the Eligible Provider or the designated representative of an Eligible Hospital. The EHR Incentive Payment will be made to the designated payee as referenced on the Attestation Agreement. It is the responsibility of the provider to verify accuracy of information contained on the Attestation Agreement, including the designated Payee.

5.4.2 Meaningful Use

Providers are eligible to receive EHR Incentive Payments for demonstrating they are meeting Meaningful Use criteria. Meaningful Users must meet the same certified EHR technology and patient volume criteria as described for A/I/U. In addition, Meaningful Users must meet required Core and Menu objectives and Clinical Quality Measures (CQM).

Meaningful User is defined in 42 CFR 495.4 as a provider that meets the EHR Incentive Payment program eligibility criteria that, for an EHR reporting period for a payment year or payment adjustment year, demonstrates meaningful use of certified EHR technology and meets the objectives and associated measures specified in the regulation and reports CQMs selected by CMS.

By definition, certified EHR technology must include the capability to electronically record the numerator and denominator and generate a report including the numerator, denominator, and the resulting percentage for all percentage-based MU measures (specified in the certification criterion adopted at 45 CFR 170.302(n)).

Please note that providers cannot use a non-certified system to calculate the numerators, denominators, and exclusion information for CQMs. The numerator, denominator, and exclusion information for CQMs must be reported directly from certified EHR technology.

As defined by 45 CFR 170.302(n), MU and CQM measures are a product of a provider’s certified EHR technology software. The MS SLR will allow providers to directly enter MU reporting and CQM attestation data or upload CQM measures from their .xml files created in their certified EHR technology. (The upload function was not required until 2014. MS SLR will validate that the requirements for MU have been met.)
DOM does not plan to propose any changes to the MU definition. Mississippi will follow the CMS regulations for defining a Meaningful User as outlined in future rule-making.

5.4.2.1 MU Reporting Period

The MU EHR reporting period is a continuous period where the provider successfully demonstrates all the MU objectives of certified EHR technology according to CMS requirements.

In the first year of MU attestation (generally the second year of MPIP participation) all providers including EPs and EHs must meet MU requirements during a single 90-day reporting period within the current calendar year in order to receive the second payment. In subsequent years of participation, the MU EHR reporting period will be a full year, unless specified by future CMS Rule-Making with attestation and payment occurring directly after the close of the calendar (EPs) or federal fiscal year (EHs). In some cases, EPs and EHs may have attested to MU with the Medicare EHR Incentive Program prior to their attestation with the MPIP; EPs and EHs falling under this category would be required to follow the CMS timeline for the MU EHR reporting period. EHs filing for both Medicare and Medicaid in the same payment year must follow the Medicare guidelines for determining MU.

Beginning with Program Year 2017, Medicaid EHR Meaningful Use participants, that also serve Medicare Part B patients (according to the guidelines and thresholds set forth by CMS), will be expected to report under the Medicare/MIPS ruling, in addition to yearly Medicaid EHR Incentive Program submissions. The Medicare/MIPS Program replaced the Medicare EHR Incentive Program, and will impact all Medicare providers that see more than 100 Medicare patients per year or bill Medicare more than $30,000 per year. The MPIP staff will work diligently to inform all current Eligible Professionals (EPs) in the MS Medicaid EHR Incentive Program of these changes and will help providers as they transition into this new (additional) way of reporting. Plans to host webinars, send out emails and update our website will be ongoing through the first few years of this reporting, starting in 2017.

5.4.2.2 Meaningful Use - EHs

As described above, after attesting to A/I/U in the first program year of the MPIP, EHs will be required to attest to MU to receive incentive payments. For EHs and CAHs, the program year now means the calendar year.

For Modified Stage 2, EHs are required to meet a total of 9 MU objectives, including one consolidated public health reporting objective. They must attest to objectives and measures using EHR Technology certified to the 2014 edition.

DOM will not require any additional MU criteria for EHs. Additionally, as a part of MU, EHs are required to submit Clinical Quality Measures (CQM) data electronically to CMS. Appendix I contains the listing of Modified Stage 2 MU core and menu set objectives.

During the attestation process in the MS SLR for Modified Stage 2, the provider is required to supply the following attestation information to qualify for Meaningful Use incentive payment:

- Select MU (first MU submission only);
- Attach external documents supporting Meaningful Use of certified EHR technology. DOM prefers that a signed contract is uploaded demonstrating proof of a fiscal relationship between the vendor and the EH. In instances in which a signed contract is not applicable DOM will accept other documentation, including but not limited to, a vendor invoice, an End-User License Agreement (EULA), or other evidence that sufficiently demonstrates MU.

- Certified EHR Technology: Enter ONC certification code. CMS publishes a list of codes identifying all ONC certified EHR technology products. During attestation the provider must enter the code from its EHR vendor to identify the EHR.

- Using certified EHR technology, respond to the Meaningful Use Core, and Clinical Quality Measures (CQM) objectives.

- Attestation Agreement: Sign and attach an Attestation Agreement indicating Meaningful Use. Attestation Agreement must be executed by the designated representative of an Eligible Hospital. The EHR Incentive Payment will be made to the designated payee as referenced on the Attestation Agreement. It is the responsibility of the provider to verify accuracy of information contained on the Attestation Agreement, including the designated Payee.

5.4.2.2.1 Dually Eligible Hospitals

Note that the CMS Registration & Attestation System is sending Medicare hospital attestation data to the State for dually eligible EHs via the Dually Eligible Hospital Attestation Data (C-5). The State must receive attestation data for core and menu objectives. The State must also receive attestation data for electronically submitted Clinical Quality Measures (CQM). Once both C-5 data transmissions have been received by the State, the Eligible Hospital is able to use the MS SLR to submit their Meaningful Use Attestations for a Medicaid incentive payment.

If the hospital is eligible for Medicare payment, then the hospital will be deemed eligible to meet Medicaid MU requirements and will not have to complete the MU validation questionnaire. As a result, the attestation agreement will show that the hospital has been deemed a meaningful user by CMS. CMS still requires the State to send the Medicaid Payment Request Response Interface (D-16) transaction prior to issuing payment. EHs that are dually eligible will still have to meet the Medicaid patient volume requirements.

5.4.2.3 Meaningful Use - EPs

After attesting to A/I/U with the MPIP, EPs will be required to attest to MU in subsequent program years to receive incentive payments. For EPs, “year” means calendar year.

Beginning with Program Year 2015, all Providers (regardless of participation years) were considered to be classified as Modified Stage 2 participants. All EPs will be required to meet a total of 10 Meaningful Use Objectives. Appendix I contains the listing of MU Modified Stage 2 Objectives.

Some MU objectives are not applicable to every provider’s clinical practice, eliminating any eligible patients or actions for the measure denominator. In these cases, the EP would be
excluded from having to meet that measure. Examples of exclusions include dentists that do not perform immunizations and chiropractors that do not e-prescribe.

EP’s must also report on 9 of 64 approved CQMs. The option to submit CQM data electronically through the MS State Level Registry will be available during the online attestation process.

During the attestation process in the MS SLR for Meaningful Use Modified Stage 2, the provider is required to supply the following attestation information to qualify for Meaningful Use incentive payment:

- Select MU.
- Attach external documents supporting Meaningful Use of certified EHR technology. DOM prefers that a signed contract is uploaded demonstrating proof of a fiscal relationship between the vendor and the EP. In instances in which a signed contract is not applicable DOM will accept other documentation, including but not limited to, a vendor invoice, EULA, or other evidence that sufficiently demonstrates MU.
- Certified EHR Technology: Enter ONC certification code. CMS publishes a list of codes identifying all ONC certified EHR technology products. During attestation the provider must enter the code from its EHR vendor to identify the EHR.
- Using certified EHR technology, respond to the Modified Stage 2 Meaningful Use Core measures and CQMs.
- Attach the following supporting documentation (required by the MS Division of Medicaid):
  - Security Risk Analysis – SRA (Meaningful Use Objective 1 – Protect Health Information) Mississippi requires that all participants complete a Security Risk Analysis tool similar or equivalent to the tool downloadable from the healthIT.gov website and upload a copy of the final summary report (generated from the tool). Providers may use a third party vendor to complete their annual Security Risk Analysis. Participants may use the same SRA for an entire group or clinic. New SRAs are required each year.
  - Full Meaningful Use Summary Report, All MU and CQM (generated by CEHRT), independently for each participating provider, objectives (to be attached on the CPOE screen)
  - Evidence of Level of Active Engagement with a Public Health Agency to submit data taken from their EHR. (Evidences may include, but will not be limited to 1) Immunization Registry – Acknowledge Letters for Registration of Intent to Onboard 2) Evidence of Testing and Validation or 3) Evidence of Production level status
- Attestation Agreement: Sign and attach an Attestation Agreement indicating Meaningful Use. Attestation Agreement must be executed by the Eligible Professional. The EHR
Incentive Payment will be made to the designated payee as referenced on the Attestation Agreement. It is the responsibility of the provider to verify accuracy of information contained on the Attestation Agreement, including the designated Payee.

5.4.3 Changes to Exclusions

Beginning in 2015, EPs that were scheduled to demonstrate Stage 1 (Meaningful Use year 1 or Meaningful Use year 2) will be given additional alternate measure exclusions. These alternate measure exclusions will decrease from 2015 to 2016. Program Year 2017 will have no alternate measure exclusions available.

5.5 MPIP MS SLR Payment Calculation/Verification

At the successful completion of the registration and attestation verification of eligibility process, DOM began to disburse incentive payments. The payment process involves a number of important activities:

- Calculating the payment;
- Verifying with CMS, via the CMS Registration & Attestation System, that the provider should not be denied payment; and
- Tracking the payment and verifying that the right payment was made to the right provider at the right time.

5.5.1 Payment Calculation

Payments are calculated differently for EPs and EHs.

5.5.1.1 EP Payment Calculation

In the MS SLR, EPs will attest that the data they enter is correct and the MS SLR will automatically determine eligibility for the incentive payment. The EP Medicaid EHR incentive payment (a fixed amount), based on the EP’s year of participation, is specified in the table below. The table includes payment for A/I/U. The preliminary payment amount is subject to DOM verification. In the event of an audit, the EP must have auditable supporting documentation, such as reports from their practice management system, for each included line item. Providers will be given the option of uploading or faxing the supporting information with their attestation.

EPs may not receive EHR incentive payments from both the Medicare and Medicaid programs in the same year. In the event an EP qualifies for EHR incentive payments from both the Medicare and Medicaid programs, the EP must elect to receive payments from only one program. After an EP qualifies for an EHR incentive payment under one program before 2015, an EP may switch between the Medicare and Medicaid programs one time. Upon switching programs, the EP will be placed in the payment year the EP would have been in had the EP not switched programs. For example, if an EP decides to switch after attesting to MU of certified EHR technology for a
Medicare incentive payment for the second payment year, then the EP would be in the third payment year for purposes of the Medicaid incentive payments.

As EPs reach their sixth or final participation year and it is discovered that previous Medicare and Medicaid (combined) payments exceed the aggregate amount of $63,750.00, the state will modify the D16 payment authorization amount to reflect the actual payment, guaranteeing that it does not, in fact exceed the specified $63,750.00. These modified D16s will be submitted (re-submitted) through a manual request process (outside the existing SLR/CMS D16 interface process) coordinated by the state and the SLR.

Table 5-2: Medicaid EP Payment Table

<table>
<thead>
<tr>
<th>Medicaid EHR Incentive Payment Schedule for Eligible Professionals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Payment Amount in 2011</td>
</tr>
<tr>
<td>Payment Amount in 2012</td>
</tr>
<tr>
<td>Payment Amount in 2013</td>
</tr>
<tr>
<td>Payment Amount in 2014</td>
</tr>
<tr>
<td>Payment Amount in 2015</td>
</tr>
<tr>
<td>Payment Amount in 2016</td>
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<tr>
<td>Payment Amount in 2017</td>
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<tr>
<td>Payment Amount in 2018</td>
</tr>
<tr>
<td>Payment Amount in 2019</td>
</tr>
<tr>
<td>Payment Amount in 2020</td>
</tr>
<tr>
<td>Payment Amount in 2021</td>
</tr>
<tr>
<td>TOTAL Incentive Payments</td>
</tr>
</tbody>
</table>

Note: The total for pediatricians who meet the 20 percent patient volume but fall short of the 30 percent patient volume is $14,167 in the first year and $5,667 in subsequent years. This adds up to a maximum Medicaid EHR incentive payment of $42,500 over a six-year period.

5.5.1.1 Medicaid EHR Incentive Payment Assignment

The following process applies only when an EP is assigning their EHR incentive payment. Such assignment of payments must be entirely voluntary for the EP. When registering for the MPIP, EPs may assign their incentive payments to their Medicaid Group account provided the EP is affiliated with the Group in the MMIS. To verify this, the payee must be a hospital or designated as a Group in the MMIS and the payee’s NPI, SSN, TIN, or Medicaid Provider Number must match with the CMS Registration & Attestation System and the PMF file. The payee must register with the CMS Registration & Attestation System using a NPI, SSN, TIN, or Medicaid Provider Number that matches the PMF file. This data cannot be changed at the State level.
As part of the annual attestation process, DOM requires that all EPs who are assigning their payment attest that the assignment is voluntary and is being made to an established Medicaid provider.

Once a payment has been disbursed by DOM to the designated payee, as assigned by the EP, the payee cannot be changed, removed or revoked. DOM expects that once a payment is assigned and an EP submits an attestation for approval, the EP authorizes payment to be made to the payee as indicated.

### 5.5.1.2 EH Payment Calculation

Hospitals need to supply several factors that go into the EH Medicaid EHR incentive payment calculation. All factors for calculating the payment amount are derived directly from the current and prior cost reports. Only CMS pre-approved data sources will be used in calculating the payment amount. These factors are based on the hospital fiscal year that ends during the federal fiscal year prior to the hospital fiscal year that serves as the first payment year, and are listed below:

- Total Medicaid Discharges (most recent four years);
- Medicaid Discharges for the Current Year;
- Medicaid Acute Inpatient Bed Days;
- Medicaid Managed Care Acute Inpatient Bed Days;
- Total Acute Inpatient Bed Days;
- Total Hospital Charges; and
- Total Hospital Uncompensated Care Charges.

DOM will verify the EH’s calculation of their overall EHR amount. The overall amount is the sum over four years of (a) the base amount of $2,000,000 plus (b) the discharge related amount defined as $200 for the 1,150 through the 23,000 discharge for the first payment year then a pro-rated amount of 75 percent in year 2, 50 percent in year 3, and 25 percent in year 4. For years 2-4 the rate of growth is assumed to be the previous 3 years' average. Note that if a hospital’s average annual rate of growth is negative over the three year period, it will be applied as such. Transition factors are applied to years one through four in the following amounts: Year One – 100 percent; Year Two - 75 percent; Year Three - 50 percent, and Year Four - 25 percent.

Auditable data sources will be used to calculate the Medicaid aggregate EHR hospital incentive amounts, as well as determining Medicaid incentive payments to these EHs. Auditable data sources for the calculation of the Medicaid EHR incentive amounts are the EH’s Medicare/Medicaid cost reports.

For the purpose of calculating the Medicaid discharges for determining the annual Medicaid patient volume percentage, DOM will allow EHs to count discharges when Medicaid is the primary or secondary payer, regardless of payment liability on the discharge. This method is in
accordance with the instructions from CMS’s Facts, Answers, and Questions section published on the CMS Website.

The “Medicaid Share,” which is applied against the aggregate EHR incentive amount, is essentially the percentage of an EH’s Medicaid inpatient days divided by the total inpatient non-charity care days. This method is in accordance with the instructions from CMS’s Facts, Answers, and Questions section published on the CMS Website.

The estimated total charges and charity care charges used in the formula must represent inpatient hospital services only and exclude any professional charges associated with the inpatient stay.

In any given payment year, no annual Medicaid EHR incentive payment to an EH may exceed 50 percent of the EH’s aggregate EHR incentive amount. Likewise, over a two-year period, no Medicaid EHR incentive payment to an EH may exceed 90 percent of the aggregate EHR incentive amount. A hospital cannot receive payments after 2016 unless the hospital received a payment for the previous year. Prior to 2016, Medicaid EHR incentive payments to EHs can be made on a non-consecutive annual basis.

Due to the high cost of hospital software and to encourage the early adoption of the EHR technology in hospitals, DOM is choosing to pay the Overall EHR Amount over the minimum three-year period at the maximum allowable percentages in each year that the EH qualifies for payment (Year 1 - 50 percent, Year 2 – 40 percent, Year 3 – 10 percent). The entire EH payment calculation is defined in the worksheet included in Appendix G.

Calculation of the Overall EHR Amount is a one-time calculation based on the following steps:

- Calculate the average annual growth rate over three years using the Medicare/Medicaid Cost Reports prior to the most current Cost Report.

- Calculate the total Medicaid discharges using the Medicaid discharges in the Medicare/Medicaid Cost Reports plus the discharges where Medicaid is the secondary payer. Only discharges between 1149 and 23,000 per CCN will be allowable discharges.

- Calculate each of the next four year’s total discharges by multiplying the previous year’s discharges times the average computed growth rate.

- Calculate the Medicaid Aggregate EHR Incentive Amount for each year by adding (total discharges times $200) to the $2,000,000 base.

- Apply the appropriate transition factor to each year’s Aggregate EHR Incentive Amount. (Year One – 100 percent, Year Two – 75 percent, Year Three – 50 percent, Year Four – 25 percent).

- Calculate the total Overall EHR Incentive Amount by adding the total of each year with the transition factor applied.
• Apply the Medicaid Share percentage to the Overall EHR Incentive Amount.  
  (See Medicaid Share calculation below).  This is the hospital’s Medicaid 
  Aggregate EHR Incentive amount.

Calculation of the Medicaid Share percentage:

• Total Medicaid days includes both the total Medicaid Days and total Medicaid 
  HMO days from the Medicare/Medicaid Cost Report.

• Calculate the non-charity percentage.  Divide the (total hospital charges less 
  uncompensated care) by the total hospital charges.

• Calculate the non-charity days by multiplying the non-charity percentage times 
  the total hospital days.

Calculate the Medicaid Share percentage by dividing the Medicaid days by the non-charity days.  
DOM has created a calculation worksheet for EHs that mirrors the calculation in the MS SLR 
application.  The calculation worksheet is included as Appendix G: EHR Hospital PIP Calculator 
and will be available on DOM’s Websites and made available through its outreach program.

Hospitals must use their filed and accepted cost report data only in the onetime calculation of 
the EH’s incentive payment amount.  EHs are required to use the last four (4) consecutive years’ 
cost reports in the calculation of the onetime payment.  Any deviation will result in the rejection 
of the EH’s application.  All cost reports are subject to audit by Medicare and Medicaid.  Any 
audit adjustments to the cost report used to calculate the onetime payment may result in a 
payment adjustment or denial of Medicaid payment at the discretion of the DOM.  Data sources 
below are in accordance with CMS FAQ 10771.

For hospitals filing the 2552-96 cost report, the authorized data sources are:

• Total Discharges - Worksheet S-3 Part 1, Column 15, Line 12
• Medicaid Days - Worksheet S-3, Part I, Column 5, Line 1 + Lines 6-10
• Medicaid HMO Days - Worksheet S-3, Part I, Column 5, Line 2
• Total Inpatient Days - Worksheet S-3 Part 1, Column 6, Line 1, 2 + Lines 6 -10
• Total Hospital Charges - Worksheet C Part 1, Column 8, Line 101
• Charity Care Charges - Worksheet S-10, Column 1, Line 30
  
  o DOM does not expect that any 2552-96 cost reports will be submitted 
  due to the change to 2552-10.  However, DOM will accept the PDF 
  version of the 2552-96 cost reports for EHR Incentive Payments or the 
  hospital can use zero for the Charity Care Charges.

For hospitals filing the 2552-10 cost report, the authorized data sources are:

• Total Discharges - Worksheet S-3 Part 1, Column 15, Line 14
• Medicaid Days - Worksheet S-3, Part I, Column 7, Line 1 + Lines 8-12
• Total Inpatient Days - Worksheet S-3 Part 1, Column 8, Line 1, 2 + Lines 8 - 12
• Total Charges - Worksheet C Part 1, Column 8, Line 200
• Charity Care Charges - Worksheet S-10, Column 3, Line 20

For new hospitals or hospitals that have a change of ownership with a new CCN, CMS is allowing states to decide when a new hospital can apply for the EHR incentive program. MS DOM has determined that a hospital must have four years of history (four cost reports) before they can apply. Cost report years containing more or less than 12 months must be excluded from the growth calculation. Only years with 12 months can be used in the calculation. The hospital must use the previous year’s cost report. For example, if cost report year 2008 contained 13 months, the hospital would have to use the cost reports for 2010, 2009, 2007, and 2006.

DOM will utilize the applicable statistics and financial data from the hospitals’ Medicare/Medicaid Cost Reports for the last four years to validate the initial calculation of the incentive payment amount and to validate that the average length of stay does not exceed the 25-day maximum. This means that the hospital must submit four cost reports on their initial application for the first payment. For subsequent years, the hospital’s cost report ending during the previous federal fiscal year will be used, and only the most recent cost report will be required.

5.5.2 CMS Verification

Before payment can be distributed, a final CMS check must be performed to validate that the provider can receive payment. The validation is done via the Medicaid Payment Request Response Interface (D-16) to the CMS Registration & Attestation System. The CMS Registration & Attestation System will return a batch interface transaction via the Medicaid Payment Request Response Interface (D-16) authorizing the payment or denying it with a Denial Reason, such as a duplicate payment or federally excluded reason.

5.6 MPIP Payment Entry/Processing

DOM will use the existing MMIS system to make provider payments. The automated payment interface from the MS SLR to the MMIS system is now operational and facilitates a streamlined payment process for the MPIP. EHR incentive payments will follow the established rules for all provider payments and will use the existing payment rules built into the current and future MMIS systems. The MMIS will notify the MS SLR that a payment was made; allowing the MS SLR to create the batch interface transaction notifying the CMS Registration & Attestation System that payment is complete.

DOM is making EHR incentive payments from the MMIS on a weekly basis. DOM makes the incentive payments to the provider, the employer, or a facility assigned the payments without any reduction or rebate. DOM does not make incentive payments to any entities promoting the adoption of certified EHR technology since none exist in Mississippi.

DOM will use existing MMIS capability to take advantage of existing reconciliation, accounting, tracking, and reporting capability supporting provider reimbursement. Reporting capabilities of...
the existing MMIS and Decision Support System/Data Warehouse (DSS) will be utilized to facilitate the CMS-37 and CMS-64 report information. Utilization of the MMIS and the DSS will allow the EHR incentive payment information to be available to the current and future audit and analysis tools built into the MMIS and DSS. DOM anticipates that the current MMIS system will be replaced during the life of the EHR incentive program.

5.7 **MPIP MS SLR Payment Complete**

As stated above, the MS SLR must send a Medicaid Payment Completion Interface transaction (D-18) to the CMS Registration & Attestation System when the payment is distributed to the Provider. The D-18 will be sent five business days after the payment is issued. Mississippi may submit an updated D18 transaction as needed to report future adjustments and possible payment recoupment.

5.8 **MPIP MS SLR Inquiry**

The MS SLR allows inquiry processes for providers to track the progress of their incentive payments, including if their attestation has been received, sent to CMS, or approved for payment. Inquiry processes may also be used by Conduent Help Desk Support Representatives to answer providers’ questions or provide guidance to providers to correct information. In addition to contacting the Conduent Help Desk, providers have the option to call DOM staff to inquire about specific information contained outside of the MS SLR.

5.9 **MPIP MS SLR Update and Risks**

DOM is participating in a multi-state SaaS solution to allow providers to attest online for their EHR incentive payment. Version 1 of the MS SLR was implemented to allow providers to apply for and submit the required documentation needed for A/I/U approval. Version 1 of MS SLR also enabled verification of most of the pre-payment audit requirements for approval of payment and captures the required documentation for additional manual review and/or audit of the attestation.

Version 2 of the MS SLR was implemented in the 1st quarter of 2012. Version 2 allows providers to attest to MU online with an immediate response that indicates whether they meet the MU requirements. Supporting documentation may include the patient volume calculators found at www.medicaid.ms.gov, contractual documents, reports from the EHR system and other documents. See the CMS-approved screenshots pertaining to Modified Stage 2 attached hereto as Appendix K.

The MS SLR also includes a Dashboard component that is an internal tool used by DOM for verification, review, internal audits, submission of audits to CMS, and processing payments. The Dashboard allows the DOM payment approver to see the attestation and all supporting documentation. The Dashboard includes expanded tools and reporting to support the additional pre- and post-payment audits, payment tracking and analysis of provider attestation.
statuses. Conduent is phasing in online post-payment audit tools and tracking of audit, appeals, and recoupment/adjustment. DOM expects that they will fully implement the audit, appeals, and recoupment/adjustment functionality available in the MS SLR once all phases are made available by Conduent.

DOM is making a best effort to apply MITA principles to all future development and deployments of the MS SLR. One challenge for DOM is using a SaaS model with multiple states, with each state having different workflows and needs. This multi-stakeholder approach has created many challenges, including configuration and customization of the application for Mississippi DOM-specific needs. For example, DOM has chosen to forgo implementing the post-payment auditing function within the MS SLR until it is more robust. Although many states are satisfied with the current functionalities available within the Conduent solution, DOM continues to perform audit, recoupment and adjustment, and appeals processes manually outside of the MS SLR due to the limited functionality.

Conduent has updated the system to incorporate Stage 1 2013 and 2014 changes related to the Final Rule. Xerox has developed and implemented changes required by the Stage 2 Final Rule from 2012. These were implemented in the State for EPs on June 25, 2014. 2014 implementation for EHs will be available on October 1, 2014.

One potential risk specific to the MS SLR relates to CMS’s changes to the definition of a Medicaid encounter from 2013. DOM foresees many challenges in verifying encounters that do not have an associated claim searchable within the MMIS. This change requires more robust post-payment audit requirements and increases the need for resources and potentially creates a larger burden upon providers to demonstrate proof through auditable data sources.

SLR Release 5.1 included functionality approved by CMS through the Addendum for Program Year 2017 allowing providers to select the desired level of attestation – Modified Stage 2 (with Program Year 2017 updates) or Stage 3. Clinical Quality Measure selection and reporting was changed to allow providers to select six CQMs that best reflected their scope of practice and removed Quality Standard domain restrictions. This was done to better align CQM reporting requirements for participants in the Quality Payment Program (QPP) and the Medicaid EHR Incentive Program. Release 5.1 was moved into production in the summer of 2017.

SLR Release 5.2 is scheduled for May 2017. Release 5.2 is only cosmetic changes and holds no impact on operation or cost of the SLR.

SLR Release 5.3 is anticipated to implement in January 2019, which is the start of Program Year 2018. This release is pending CMS details for CQM reporting and functionality.
5.9.1 SMA Hosted Website

DOM has launched a public-facing website that includes links to the MS State Level Registry as well as program resources for providers. This can be accessed by the public at https://MSEHRPIP.wordpress.com.

5.10 Program Oversight

5.10.1 MPIP MS SLR Prepayment Verification

DOM is conducting a robust and comprehensive prepayment oversight program. The prepayment oversight activities are led by the Office of Information Technology Management (iTECH). The levels of prepayment oversight and monitoring include the review, tracking and verification of provider attestations, including all of the information and documents necessary for a Medicaid provider to receive an incentive payment for each program year. This process ensures each provider meets provider registration, attestation, and eligibility criteria prior to receiving their incentive payment. Prepayment verifications are primarily performed by the MS SLR through configurable items within the application; however, iTECH staff members also perform some manual verification prior to releasing providers for payment.

5.10.1.1 Automated Prepayment Verification Process

As a part of the prepayment verification process, the automated MS SLR functions and the CMS Registration and Attestation System are leveraged to assure that no duplicate Medicaid EHR incentive payments are paid by more than one state or between the Medicaid and Medicare programs. The MS SLR automated processes and manual stops will also ensure that the incentive payments are made accurately, without reduction or rebate and will be made directly to a provider or to an eligible third party entity to which the provider has assigned payments.

DOM has created a PMF that consists of all EPs and EHs to compare to B-6 Interface information during MS SLR Registration. The PMF excludes all providers whose licenses have expired, as well as all OIG excluded providers and State of Mississippi exclusions. The PMF also includes those EPs who qualify as “non-hospital” based and excludes EPs listed on the State death registry. The PMF is automatically generated weekly from the MMIS provider master and claims data files. The PMF file will be the control file used by the MS SLR for approval of all EP and EH attestations. The CMS and OIG sanctions are updated monthly; the State of Mississippi sanctions are updated daily.

In addition to verification against the PMF, the MS SLR has been configured to automate several prepayment verifications on information entered by the provider during attestation. The MS SLR incorporates hard stops to verify that all information entered by providers aligns with program rules and that required documents are attached.

The MS SLR will automatically verify the following items during the attestation process:

- Eligibility reporting period using dates entered by the provider;
• (EHs only) – Average Length of Stay is less than 25 days;
• Medicaid patient volume (or Needy Individual Patient Volume) using numerator and denominator;
• ONC EHR certification number by matching the provider certification number with the ONC Certified HIT Product List;
• A/I/U criteria or MU criteria, depending upon the attestation type; and
• Provider NPI and SSN/TIN and payee NPI and SSN/TIN with the PMF.

Providers will be required to upload documentation in support of many of these items prior to proceeding in the MS SLR as well. If any one item cannot be verified, then the attestation will stop and the provider will not be able to proceed until corrected.

In the final step of attestation in the MS SLR, providers are required to submit an attestation agreement document. DOM currently uses a comprehensive attestation document that ensures DOM and CMS that the provider meets the requirements for eligibility and incentive payment. The attestation agreement will be automatically generated from the information entered into the MS SLR by the provider and will vary based on provider type. The attestation agreement includes the following statements that the provider:

• Is voluntarily participating in the Mississippi Medicaid EHR Incentive Payment Program;
• Has met all of the eligibility requirements for the program for the payment year;
• Has created a binding legal or financial obligation to acquire, implement or upgrade to the CMS Certified EHR software identified by the CMS EHR Certification identification;
• Agrees that any assignment of the EHR Incentive Payment is made voluntarily;
• Understands that their application is subject to review and/or audit by the State of Mississippi and that all supporting data must be maintained for a minimum of seven years;
• Understands that any falsification or concealment of material information may result in the provider being declared ineligible to participate in this program or any other Mississippi Medicaid program;
• Understands that any incentive payments found to have been made based on fraudulent information or attestation may be recouped by DOM, including all collection costs and penalties that may be assessed by the State of Mississippi;
• Understands that the EHR incentive payments are treated like all other income and are subject to federal and state laws regarding income tax, wage garnishments, and debt recoupment;
• Certifies that information contained in the MS SLR and attestation agreement is true, accurate, and complete; and

• Understands that Medicaid EHR incentive payments submitted under this provider number will be from federal funds and that any falsification or concealment of a material fact may be prosecuted under federal and state laws.

Moreover, given that this is a legally binding document, DOM requires the following:

• The above statement will appear directly above the provider’s signature or, if they are printed on the reverse of the form, a reference to the statements must appear immediately preceding the provider’s signature;

• The provider’s signature;

• The provider and provider’s name, NPI, SSN, and TIN appears on the attestation agreement;

• The provider is responsible for verifying both the provider and provider’s payee information is correct on the attestation agreement; and

• The provider attestation must be resubmitted upon any change in the provider’s attestation and/or representative.

As a final step in the prepayment verification process, the MS SLR will work to prevent multiple payments to providers by:

• Indexing files using the CCN, NPI, and TIN as the key for EHs;

• Indexing files using NPI and SSN for all other providers; and

• Requiring an NPPES Web account through the CMS Registration and Attestation System before an attestation can be complete.

  o EPs – the Web account is only issued using the Provider’s SSN. The individual Provider is only issued one account per SSN.

  o EHs – the Web account is only issued using the hospital’s CCN. The hospital is only issued one account per CCN.

5.10.1.2 iTech Staff Prepayment Verifications

DOM iTech staff members are responsible for conducting manual prepayment verifications and provider outreach. To ensure that staff levels are appropriate for the MPIP program, quarterly reports are reviewed to assess attestation-to-payment time and provider outreach efficiency. Over time, staff levels have been increased to support paying incentives in a timely manner.

Conduent offers a HelpDesk call center for all providers covered by this application. However, in Mississippi we encourage all EPs and EHs to contract our program staff directly with questions.
or concerns. This information is posted on our website (https://MSEHRPIP.wordpress.com/Contract-Us)

5.10.1.2.1 Manual Prepayment Verification Process

iTECH staff review every attestation prior to releasing for payment. Given that the MS SLR cannot automatically verify all information, the iTECH manual verification process for all providers includes:

- Ensuring that all documentation attached is correct and accurate as described by the MS SLR;
- Verifying that CEHRT standards are met by the submission of currently required certification numbers from the ONC (i.e. 2014 and beyond);
- Verifying that the certified EHR technology contract is valid within the last 12 months;
- Ensuring that the attestation agreement is signed and valid according to DOM regulations; and
- (For MU only) verifying required documents are attached and appropriate for chosen MU measures.

All attestations found without proper documentation attached will be pended and a notice identifying the missing or incorrect information will be sent to the provider’s e-mail address with instructions on how to correct.

In addition to verifying documentation, iTECH performs several other manual verifications on EPs prior to payment. These verifications include:

- Verifying that the EP is affiliated with the assigned payee in the MMIS and that the EP payee has a group indicator, if applicable; and
- Verifying that the SLR payment report matches the SLR request for approval to pay file.

Any exceptions are noted and researched for the reason for non-approval. The following is a “checklist” of items that will be used by iTECH staff to verify attestations prior to payment.

**Table 5-3: Checklist of Items for Pre-Payment Verification**

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Automated State Level Registry System / Manual Process</th>
</tr>
</thead>
<tbody>
<tr>
<td>Collect and verify basic information to assure Provider enrollment eligibility upon enrollment or re-enrollment to the Medicaid EHR payment incentive program.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td>Requirement</td>
<td>Automated State Level Registry System / Manual Process</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------</td>
<td>-------------------------------------------------------</td>
</tr>
<tr>
<td>Collect and verify basic information to assure patient volume in the numerator. Both the Medicaid and total patient volumes will be verified.</td>
<td>Automated - MS SLR</td>
</tr>
<tr>
<td></td>
<td>Manual – Provider management reports and Review of Provider supporting documentation</td>
</tr>
<tr>
<td>Collect and verify basic information to assure that PA EPs are practicing predominantly in a FQHC or RHC and are so led by the PA.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td>Assure that Medicaid providers who wish to participate in the EHR incentive payment program have or will have a NPI and will choose only one program from which to receive the incentive payment using the NPI, a TIN, and CMS’ national provider election database.</td>
<td>Automated – CMS Registration &amp; Attestation System and MS SLR</td>
</tr>
<tr>
<td></td>
<td>Manual – Review NPI, TIN and active license for validity</td>
</tr>
<tr>
<td>Based on provider type, assure that the provider meets all requirements to be eligible to participate in the EHR Payment Incentive Program as a Medicaid Provider. “All requirements” means all requirements that can be verified using external data sources available to DOM.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td></td>
<td>Manual - Review of provider supporting documentation</td>
</tr>
<tr>
<td>To eliminate long-term care hospitals, ensure that a hospital eligible for incentive payments has demonstrated an average length of stay of 25 days or less.</td>
<td>Automated – MS SLR will calculate the average length of stay for all hospitals. The calculation will be the total number of inpatient days divided by the total number of discharges. The application has a hard stop and will not allow the application to proceed if the average length of stay is greater than 25 days.</td>
</tr>
<tr>
<td>Ensure all eligibility information is verified at least on an annual basis. Provider eligibility information is only going to be verified when the Provider requests a payment via the MS SLR.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td></td>
<td>Manual - Review of Provider supporting documentation</td>
</tr>
<tr>
<td>Requirement</td>
<td>Automated State Level Registry System / Manual Process</td>
</tr>
<tr>
<td>---------------------------------------------------------------------------</td>
<td>--------------------------------------------------------</td>
</tr>
<tr>
<td>Verify the Provider has met the certified EHR requirements, through use of the ONC - certified EHR code and attached vendor contracts, purchase order, EULA or license agreement.</td>
<td>Automated - MS SLR&lt;br&gt;Manual verification is required to ensure the document attached is the type to which attestation is made.</td>
</tr>
<tr>
<td>Based on Provider type, assure the MU Core requirements have been attested to and are accurate.</td>
<td>Automated - MS SLR&lt;br&gt;Manual – review specific objectives, including CPOE, problem list and DOM security risk analysis questionnaire&lt;br&gt;*The DOM security risk analysis questionnaire can be found at <a href="http://www.medicaid.ms.gov">www.medicaid.ms.gov</a></td>
</tr>
<tr>
<td>Based on Provider type, assure the proper number of MU Menu Item requirements have been attested to and are accurate.</td>
<td>Automated - MS SLR</td>
</tr>
<tr>
<td>Capture and verify clinical quality measures from each Provider.</td>
<td>Automated –MS SLR</td>
</tr>
<tr>
<td>Based on Provider type, assure the first year payment is accurately calculated.</td>
<td>Automated - MS SLR</td>
</tr>
<tr>
<td>Based on Provider type, assure the payment for years two through six are accurately calculated.</td>
<td>Automated - MS SLR</td>
</tr>
<tr>
<td>Assure a Provider does not receive incentive payments for more than six years.</td>
<td>Automated – CMS Registration &amp; Attestation System and MS SLR</td>
</tr>
<tr>
<td>Assure a Provider does not receive duplicate payments for any given year.</td>
<td>Automated – CMS Registration &amp; Attestation System and MS SLR</td>
</tr>
<tr>
<td>Requirement</td>
<td>Automated State Level Registry System / Manual Process</td>
</tr>
<tr>
<td>-------------</td>
<td>------------------------------------------------------</td>
</tr>
<tr>
<td>Ensure that each Provider that collects an EHR incentive payment has collected an incentive payment from only one state, even if the Provider is licensed to practice in multiple states.</td>
<td>Automated – CMS Registration &amp; Attestation System and MS SLR</td>
</tr>
<tr>
<td>Assure payments are not made for any year starting after the year of 2015 unless the Provider has been provided payment for a previous year within the active program period.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td>Assure that Medicaid EHR incentive payments are made without reduction or rebate have been paid directly to a Provider or to an employer, a facility, or an eligible third-party entity to which the Medicaid Provider has assigned payments.</td>
<td>Automated – MS SLR</td>
</tr>
<tr>
<td>Ensure that any existing fiscal relationships with providers to disburse the incentive payments through Medicaid managed care plans does not result in payments that exceed 105 percent of the capitation rate, in order to comply with the Medicaid managed care incentive payment rules at §438.6(v)(5)(iii).</td>
<td>Does not apply to MS providers. Incentive payments are made directly to the provider.</td>
</tr>
<tr>
<td>Ensure that only appropriate funding sources are used to make Medicaid EHR incentives. DOM apportions money from the proper account, via existing DOM accounting processes, before the money is disbursed.</td>
<td>Manual - MMIS and State accounting processes.</td>
</tr>
</tbody>
</table>
5.10.1.3 MMIS Automated Audits

The MMIS conducts automated audits before payment is generated in the MMIS. MMIS audits include:

- Verifying that the provider is affiliated with the payee in the MMIS Provider File to make a payment to the payee listed in the MS SLR. If this affiliation is not present, the provider will be notified of the error and will be given instructions on how to correct the problem;

- Verifying that the provider’s Mississippi Medicaid ID is active; and

- (For EPs only) – Verifying that the EP’s license is active and valid.

5.10.2 Financial Reporting

The Office of Finance and Performance Review (OFPR) conducts audits, handling all compliance audits. OFPR reports through Finance to the Executive Director. Program Integrity handles all provider billing audits and fraud identified by the OFPR. Program Integrity reports through Health Services to the Executive Director.

MPIP Financial Reporting is conducted through iTECH and OFPR by leveraging functions available in the MS SLR. The MS SLR incorporates reporting capabilities for the incentive payment program, including pre-payment verification activities, post-payment auditing activities, and incentive payment amounts by provider type. iTECH and the OFPR utilize these reporting capabilities, in addition to guidance from the Final Rule, to report to CMS on oversight activities and financial activities.

DOM claims federal reimbursement in accordance with all applicable federal laws, regulations, and policy guidance. More specifically, the OFPR has a process in place to ensure that its expenditures for administration of the MPIP will not be claimed at amounts higher than 90 percent of the cost of such administration. A separate reporting category, 039 SLR Incentive Payments, has been established to identify all direct costs related to the Medicaid EHR incentive payment program. This category of service is tracked throughout the following reports produced from the MMIS:

- RX045 – Final Payment Summary
- RX047 – Financial Transaction Summary
- RX048 – Medicaid Register by Provider Type
- RX051 – Preliminary Payment Summary
- RX053 – Remittance Activity Control Totals
- RX054 – Remittance Advice (RA)
- RX100 – Final Payment Estimation by Billing Provider
• RX124 – Weekly Category of Service Summary
• RX134 – New Financial Transactions Report
• RX141 – Financials by Category of Service
• RX241 – Monthly Financials by Category of Service
• RX245 – Monthly Final Payment Summary
• RX341 – Quarterly Financials by Category of Service
• RX345 – Quarterly Final Payment Summary

Administrative costs are determined based on our agency accounting records. Expenses related to HIT are designated with distinct reporting codes within the accounting system. Monthly and quarterly account reconciliations and preparation of the quarterly CMS-64 reports identify all administrative expenditures related to the Medicaid EHR incentive payment program, including any expenditure erroneously claimed at an amount higher than 90 percent. The Office of Finance and Performance Review would take corrective action immediately if erroneous expenditures are identified.

The Office of Finance and Performance Review also has a process in place to ensure that it does not claim amounts higher than 100 percent of the cost of such payments to providers. This control process will be supported by reports based on data extracted from MMIS and the MPIP MS SLR solution, which will be compared to estimated expenditures from the CMS-37.

Additional financial oversight reports include:

**Table 5-4: Additional Financial Oversight Reports**

<table>
<thead>
<tr>
<th>Report</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reports showing payments pending by Provider.</td>
<td>Weekly and Monthly</td>
</tr>
<tr>
<td>Reports showing payments made by Provider.</td>
<td>Weekly and Monthly</td>
</tr>
<tr>
<td>Payment reconciliation reports to track payment by NPI/Provider ID from MS SLR to MMIS to MS SLR to the CMS Registration &amp; Attestation System.</td>
<td>Weekly and Monthly. Dollars in the payment calculation of MS SLR by Provider. Dollars input in to the MMIS system by Provider. Payments made by MMIS to Provider. Payments reported to the MS SLR by Provider. Payments reported to the CMS Registration &amp; Attestation System by Provider.</td>
</tr>
<tr>
<td>Reports tracking the status of all applications in the redetermination or appeals processes.</td>
<td>Weekly and Monthly</td>
</tr>
<tr>
<td>CMS Report with number of providers by type and location using A/I/U.</td>
<td>Year One Report - Quarterly and Annually</td>
</tr>
</tbody>
</table>
DOM will create additional reports as necessary to administer, manage, and monitor MPIP.

5.11 Audit Strategy

DOM began making payments to providers in May 2011. Since that time, DOM has conducted an ongoing evaluation of its verifications and Audit Strategy. As a result of this ongoing evaluation, DOM has determined that it will conduct pre-payment verifications of 100 percent of all provider attestations and will follow a rigorous pre-payment verification process. As noted above, certain pre-payment verifications are automated through the MS SLR, while other pre-payment verifications are manually completed by iTech staff. The verification workflow begins after the provider completes registration and attestation. DOM has up to 60 days to verify the provider’s eligibility and an additional 45 days to distribute payment. This 45-day period starts after payment authorization is confirmed through the Medicaid Payment Request Response Interface (D-16).

DOM Office of Finance and Performance Review (OFPR) staff members are responsible for conducting post-payment audits on behalf of DOM. OFPR staff members will leverage all existing data sources for post-payment verifications, including MMIS claims data for comparison to a provider’s self-reported data.

Post-payment audits of providers that have attested to and been paid for A/I/U have already commenced. OFPR will begin conducting post-payment audits of providers that have attested to and been paid for MU in 2013. The post-payment MU audit strategy is included in Appendix J. Appendix J is marked as confidential and will not be released as part of the public document.

5.11.1 Pre-Payment Audits

DOM conducts pre-payment audits for A/I/U and MU on 100 percent of provider attestations using the process previously explained in Section 5.10.1.

5.11.2 Post-Payment Audits

DOM conducts post-payment audits for A/I/U and MU as outlined in Appendix J. Appendix J is a confidential document and will not be posted on public Websites.
DOM acknowledges that the Audit Strategy, including pre and post-payment verifications, for A/I/U and MU as outlined above and in Appendix J will need to be evaluated on a regular basis. In subsequent SMHP updates, DOM will include necessary revisions to the Audit Strategy, as a part of the Appendices, to reflect the level of risk encountered in attestation reviews and based on lessons learned as the MPIP proceeds.

### 5.11.3 Fraud and Abuse

Abuse is defined as provider practices that are inconsistent with sound fiscal, business or medical practices and result in unnecessary costs to DOM. Fraud is when the provider has the intent to deceive or misrepresent with knowledge that this deception could result in an unauthorized benefit. Fraud detection focuses on providers with intent to commit either a civil or criminal action for personal gain. Fraud and abuse prevention includes the previously described pre and post-payment verification and audit activities with additional investigation that starts at the conclusion of the initial pre and post-payment audit processes. When DOM determines that there is an issue related to payment that is more than a provider’s mistake or error or negligence then the provider is referred to the Attorney General’s Medicaid Fraud Control Unit (MFCU) for investigation. The MFCU has specific authority to investigate and prosecute Medicaid fraud and abuse using search warrants and administrative document request. The MFCU may determine settlements, obtain judgments and convictions and recover criminal and civil restitution, fines, penalties and costs.

#### 5.11.3.1 Recoupment

Conduent has completed and implemented all development work surrounding Audit, Appeals, Recoupment and Adjustment in the MS SLR. This functionality (ability to capture recoupment and adjustment information, including tracking recoupments/adjustments and flagging providers that have been paid improperly in previous program years) is currently available in the MS SLR. This was deployed into a Production environment in late November, 2013.

Recoupments and adjustments of Medicaid EHR incentive payments will be handled in the same fashion as all other Medicaid claims. DOM will use its current recovery process (MS Code 43-13-121) to take corrective action regarding any improper payments to providers through the MPIP. DOM recognizes the need to repay CMS all FFP received by providers in the event of an improper payment, regardless of whether or not DOM has actually received the recoupment.

DOM plans to use the current MMIS functionality to track overpayments and will utilize MMIS negative payment files to facilitate the recoupment or adjustment of incentive payments. To date, DOM has not completed a recoupment or adjustment for any incentive payments that have been distributed.

### 5.12 Administrative Redetermination and Appeal Plan

This section of the SMHP describes the DOM appeals process regarding the MPIP appeal rights, the valid reasons for an appeal, and types of provider eligible for an appeal. The redetermination and appeal processes will proceed in accordance with the Mississippi state law.
Specifically, Medicaid Providers can appeal if they believe that they have been incorrectly denied an incentive payment, or have received an incorrect payment amount because of an incorrect determination of eligibility, including but not limited to the following DOM decisions:

- Measuring patient volume;
- Demonstrating MU; and
- Efforts to adopt, implement, or upgrade to certified EHR technology.

The first step in the appeals process is for the provider to request an informal reconsideration prior to invoking a formal appeal. This can be achieved by contacting iTech or OFPR staff. iTech or OFPR staff may grant the provider the opportunity to make changes to their MS SLR information after the informal reconsideration process and discussion. If the reconsideration process results in a denial decision, MS DOM will provide a written notification of the denial action to the provider. The provider may then proceed in the appeals process by submitting a formal appeal to DOM at that time.

The provider may formally appeal the decision by filing a written notice for appeal with the Office of Administrative Appeals within 30 days of the written receipt of the adverse decision. State of Mississippi law requires that providers file a formal appeal in writing, detailing the reason for the appeal. DOM uses an internal system to track all appeals and all supporting documentation is stored on a secure server within DOM. The notice of appeal is considered filed when it is date stamped by the Office of Administrative Appeals. The notice must identify the issues being appealed, explain the reasons why the provider disagrees with the adverse decision, and include all supporting documentation.

DOM manually updates the status of all formal appeals in the National Level Repository (NLR). This process allows DOM to maximize the benefits of using the existing system for all appeals and minimizes administrative costs of the program. Redetermination is an informal process and documented within an internal system.

Appeals, audits, fraud and abuse administration and work will be supported by processes external to MS SLR and may take place at any point described above (Registration, Attestation, etc.). “Historical log” information will be stored in the MS SLR that documents the initiation, progress, and results of each appeal, audit, and recoupment or adjustment case. Mississippi has a substantial investment in staff training and systems designed to facilitate and track appeals, audits, fraud and abuse. Mississippi will leverage this investment to reduce the administrative cost of the EHR incentive payment program. Documentation generated during the process will be secure and readily available to DOM staff to assist in answering provider questions.

DOM has an existing relationship with the Mississippi Attorney General’s Office Medicaid Fraud Control Unit and has incorporated this process as part of the MPIP oversight responsibilities.

The provider will receive a fair hearing in accordance with the Division of Medicaid State of Mississippi’s Administrative Code Title 23, Part 300 – Appeals. DOM has not updated its appeals process since program inception, but may reserve the right to do so in subsequent SMHP.
updates based upon lessons learned and the number and type of appeals being filed and processed on an annual basis.

5.12.1 Miscellaneous Provider Issues and Complaints

DOM has established an e-mail address for provider issues and complaints. The e-mail account is monitored daily and distributed to the appropriate person to resolve the issue. Mississippi DOM assists providers in addressing all issues as quickly as possible. DOM will track the issue to its final resolution and will maintain a log of ongoing and resolved issues. DOM will summarize and categorize all provider issues received.

5.13 MPIP MS SLR Post Payment Processing

Whenever a provider’s incentive payment is adjusted due to an audit finding, the state will notify CMS via a CMS Registration & Attestation System Medicaid Payment Adjustment Interface (D18 – payment adjustment/recoupment) transaction.

5.14 Quarterly Reporting to CMS

CMS implemented a standard report format for quarterly reporting on EHR Incentive Payment program measures of progress. DOM submits these quarterly reports directly to CMS on or before the required deadlines on the required CMS template. The template includes the following items:

- State System - Dates
  - Registration Implementation
  - AIU Attestation Implementation
  - Payments Implementation
  - Audits Implementation
  - MU Attestation
  - IAPD Expiration
- Provider Outreach – Number and Dates
  - Outreach Events
  - Phone Calls
  - Emails
- Auditing – Planned and Actual Dates
  - EP AIU Audits
  - EP MU Audits
  - EH Audits
- State-Specific SMHP Tasks – Planned and Actual Dates
  - Conduct Year One post payment audits and analysis
  - Finalize audit plan for Year Two MU and other program requirements
  - Receive CMS APD approval for eligibility determination remediation
  - Develop requirements/release RFP for interface to the State HIE and Sequoia Project (eHealth Exchange)
o Create RFPs for Sequoia Project (eHealth Exchange) platform consulting, IV&V, and implementation vendors
  o Release MMIS system replacement RFP
  o Develop audit plan for MU and other program requirements
  o Start development of required changes to the MS SLR
  o Share limited Medicaid data with local HIEs as agreed and requested (e.g., MSCHIE)
  o Finalize audit plan for MU and other program requirements

• Staffing Levels and Changes – Planned and Actual
  o Operational Staff
  o IT Staff
  o Auditing Staff
  o New Staff This Quarter

• EP/EH Counts and Amounts Paid (Total since start of program)
  o EP AIU Count
  o EP AIU Paid Amount
  o EP MU Count
  o EP MU Paid Amount
  o EH AIU Count
  o EH AIU Paid Amount
  o EH MU Count
  o EH MU Paid Amount

• Other Information
  o Additional tasks
6 HIT Roadmap

6.1 Major Activities and Milestones Moving from “As-Is” to “To-Be”

The following table shows the major activities and milestones to move DOM from the “As-Is” to the “To-Be” status. There are several recurring activities shown within the table that should be pointed out. These activities show only one quarter, but continue throughout the Milestone Schedule on a quarterly basis. The recurring activities include:

- **Implementation of MU for EH and EP** – Starting in the third quarter of FFY 2012, the MS SLR began accepting MU attestations. Although this is shown as a milestone that ended in Q3 of FFY2012, the MU functionality remains active in the MS SLR;

- **Post Payment Audit Implementation** – In the fourth quarter of FFY2012, the post payment audit program was initiated. As noted in Section 5 – Provider Incentive Program Blueprint, post payment audits have commenced for A/I/U attestations, as well as MU attestations. Post payment audits will continue on a regular basis throughout the program; and

- **SMHP and IAPD Annual Updates** – Beginning in the second quarter of FFY2012, DOM has submitted annual updates of the SMHP and IAPD to CMS for approval. Annual SMHP updates include changes to the “As-Is” and “To-Be” landscape, policy changes to the MPIP, and a new HIT Roadmap. Annual IAPD updates outline the requested funds for implementing HIT initiatives outlined in the SMHP.

Table 6-1: Master Milestones/Schedule

<table>
<thead>
<tr>
<th>MILESTONE</th>
<th>START DATE</th>
<th>END DATE</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>State Level Registry (SLR) Upgrades</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meaningful Use UAT</td>
<td>Q2 FFY12</td>
<td>Q2 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>Implementation of Meaningful Use for EH and EP (On-going)</td>
<td>Q3 FFY12</td>
<td>Q3 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>First EP Payments for Meaningful Use</td>
<td>Q3 FFY12</td>
<td>Q3 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>Provider Training on Meaningful Use</td>
<td>Q4 FFY12</td>
<td>Q4 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>Post Payment Audit Implementation (On-going)</td>
<td>Q4 FFY12</td>
<td>Q4 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>MMIS / SLR Payment Electronic Interface Implementation</td>
<td>Q4 FFY12</td>
<td>Q4 FFY12</td>
<td>Completed</td>
</tr>
<tr>
<td>SMHP Update for Stage 2 Final Rule Changes</td>
<td>Q1 FFY13</td>
<td>Q1 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 2.4 - Stage 1 Changes for 2013 Implementation</td>
<td>Q1 FFY13</td>
<td>Q1 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 2.5</td>
<td>Q2 FFY13</td>
<td>Q2 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 2.6</td>
<td>Q3 FFY13</td>
<td>Q3 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Functionality for Audit, Recoupment &amp; Adjustment, and Appeals</td>
<td>Q3 FFY13</td>
<td>Q3 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 2.7</td>
<td>Q4 FFY13</td>
<td>Q4 FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>MILESTONE</td>
<td>START DATE</td>
<td>END DATE</td>
<td>STATUS</td>
</tr>
<tr>
<td>--------------------------------------------------------------------------</td>
<td>------------</td>
<td>----------</td>
<td>------------</td>
</tr>
<tr>
<td>SLR Release 3.0 - Stage 2 Meaningful Use Implementation for EH</td>
<td>Q1 FFY14</td>
<td>Q1 FFY14</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 3.1 - Stage 2 Meaningful Use Implementation for EP</td>
<td>Q2 FFY14</td>
<td>Q2 FFY14</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 3.2 – Stage 2 Meaningful Use Implementation for EH (additional e-CQM reporting interface from CMS)</td>
<td>Q4 FFY14</td>
<td>Q1 FFY15</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 3.3 - Response to CMS NPRM (effective October 1, 2014) Additional development needed to allow providers to take advantage of Flexibility Rule for CEHRT 2011, 2014 or combination 2011/14</td>
<td>Q4 FFY14</td>
<td>Q1 FFY15</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 4.0 – SLR Dashboard and Internal Reporting Enhancements</td>
<td>Q1 FFY15</td>
<td>Q3 FFY15</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 4.1 – Modifications to Program Year 2015 for Modified Stage 2 for EPs and EHs</td>
<td>Q3 FFY15</td>
<td>Q3 FFY16</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 4.1.2 – Modifications to Program Year 2016 for Modified Stage 2 for EPs and EHs</td>
<td>Q3 FFY16</td>
<td>Q2FFY17</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 5.0 and Release 5.1 - Modifications to Program Year 2017 for Modified Stage 2 and Stage 3 for EPs and EHs (implementing the requirements as outlined in the recent IPPS ruling – published August 2, 2017)</td>
<td>Q3 FFY17</td>
<td>Q3FFY18</td>
<td>Completed</td>
</tr>
<tr>
<td>SLR Release 5.2 - Cosmetic clean-up of SLR solution only</td>
<td>Q2FFY18</td>
<td>Q3FFY18</td>
<td>In Progress</td>
</tr>
<tr>
<td>SLR Release 5.3 – Regulatory updates based upon pending CMS new ruling</td>
<td>Q1FFY19</td>
<td></td>
<td>Pending</td>
</tr>
</tbody>
</table>

**Environmental Scan**

<table>
<thead>
<tr>
<th>MILESTONE</th>
<th>START DATE</th>
<th>END DATE</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Planning</td>
<td>Q1 FFY16</td>
<td>Q1 FFY17</td>
<td>Completed</td>
</tr>
<tr>
<td>Survey Development</td>
<td>Q1 FFY17</td>
<td>Q2 FFY17</td>
<td>Completed</td>
</tr>
<tr>
<td>Visits and Surveys</td>
<td>Q1 FFY17</td>
<td>Q2 FFY17</td>
<td>Completed</td>
</tr>
<tr>
<td>Collection of Data / Analysis of Information</td>
<td>Q1 FFY17</td>
<td>Q3 FFY17</td>
<td>Completed</td>
</tr>
<tr>
<td>Report / SMHP Update</td>
<td>Q2 FFY17</td>
<td>Q3 FFY17</td>
<td>Completed</td>
</tr>
</tbody>
</table>

**Outreach to providers in the EHR Incentive Program**

<table>
<thead>
<tr>
<th>MILESTONE</th>
<th>START DATE</th>
<th>END DATE</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Work on getting providers that previously attested to return to the program</td>
<td>Q3 FFY 18</td>
<td>Q3 FFY 19</td>
<td></td>
</tr>
<tr>
<td>Targeted Outreach to prepare providers for Stage 3 Meaningful Use Attestation</td>
<td>Q3 FFY 18</td>
<td>Q3 FFY 19</td>
<td></td>
</tr>
<tr>
<td>Targeted Outreach to help providers better understand the importance and workflow of Active Care Team Coordination; Submitting to Public Health Specialized Registries; Sharing electronic health records using the existing functionality of the EHR (moving away from the fax machine)</td>
<td>Q3 FFY 18</td>
<td>Q3 FFY 19</td>
<td></td>
</tr>
</tbody>
</table>

**SMHP and IAPD Annual Update**

<table>
<thead>
<tr>
<th>MILESTONE</th>
<th>START DATE</th>
<th>END DATE</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q3 FFY17</td>
<td></td>
<td>Q4 FFY17</td>
<td>In Progress</td>
</tr>
</tbody>
</table>

**DOM Interoperability Platform Acquisition and Implementation**

<table>
<thead>
<tr>
<th>MILESTONE</th>
<th>START DATE</th>
<th>END DATE</th>
<th>STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vendor analysis and review of offerings, including presentations, HIMSS meetings</td>
<td>Q2FFY14</td>
<td>Q2FFY13</td>
<td>Completed</td>
</tr>
<tr>
<td>Procure Interoperability Staff</td>
<td>Q2FFY14</td>
<td>Q4FFY14</td>
<td>Completed</td>
</tr>
<tr>
<td>Write RFP for Interoperability Platform</td>
<td>Q1FFY15</td>
<td>Q4FFY15</td>
<td>Completed</td>
</tr>
<tr>
<td>Open bids for vendors</td>
<td>Q2FFY16</td>
<td>Q2FFY16</td>
<td>Completed</td>
</tr>
<tr>
<td>Evaluate bids for vendors</td>
<td>Q2FFY16</td>
<td>Q2FFY16</td>
<td>Completed</td>
</tr>
<tr>
<td>Negotiate contract with vendor</td>
<td>Q2FFY16</td>
<td>Q2FFY16</td>
<td>Completed</td>
</tr>
<tr>
<td>Implement Interoperability Platform</td>
<td>Q3FFY16</td>
<td>Q2FFY17</td>
<td>Completed</td>
</tr>
</tbody>
</table>
### 6.2 Governing Law

The following is a summary of federal and state law and state administrative rules applicable to the SMHP. DOM is in compliance with all relevant law and rules.

- **45 C.F.R. Part 170**, entitled *Health Information Technology Standards, Implementation Specifications, and Certification Criteria and Certification Programs for Health Information Technology*.

These regulations implement parts of the Public Health Service Act regarding Health Information Technology. The standards, implementation specifications, and certification criteria adopted in these regulations apply to Complete EHRs and EHR Modules and the testing and certification of such Complete EHRs and EHR Modules. These requirements regarding certified EHRs include the requirement known as “meaningful use” which requires that the EHR possess among other things, “capabilities that are necessary to meet the objectives and associated measures [required of eligible professionals, eligible hospitals, and critical access hospitals] and successfully report the clinical quality measures selected by CMS in the form and manner specified by CMS (or the States, as applicable) for the stage of meaningful use that an eligible professional, eligible hospital, or critical access hospital seeks to achieve.” The Mississippi Division of Medicaid (DOM) electronic health record systems fell under this requirement between the dates of July 1, 2013 to June 30, 2014. These regulations do not apply to the Provider Portal.

During the applicable period of July 1, 2013 to June 30, 2014, DOM was in compliance with 45 C.F.R. Part 170 while offering a Certified EHR to Medicaid providers. Because DOM is no longer offering an EHR, these regulations no longer apply.


HIPAA applies to covered entities, which include health plans, health care clearinghouses, and health care providers who transmit any health information in electronic form in
connection with a transaction covered by HIPAA, as well as business associates of covered entities. It requires (1) certain security standards for the protection of electronic protected health information, (2) certain notification requirements if there is a breach of unsecured protected health information, and (3) certain privacy standards regarding individually identifiable information.

During the applicable period of July 1, 2013 to June 30, 2014, DOM was in compliance with HIPAA while offering a Certified EHR to Medicaid providers. For the follow-on product, DOM’s Provider Portal was compliant July 1, 2014 to current and will continue to be.


  These regulations impose restrictions upon the disclosure and use of alcohol and drug abuse patient records which are maintained in connection with the performance of any federally assisted alcohol and drug abuse program.

  Where applicable, DOM is compliant with 42 C.F.R. Part 2 through implementation of its sensitive data policy, which prohibited the display/disclosure of alcohol and drug abuse data in the former EHR and continues to prohibit the display/disclosure of such data in the Provider Portal.

- **Miss. Code Ann. § 41-21-97**, entitled *Confidentiality of Hospital Records and Information; Exceptions*, in regards to persons in need of or receiving mental treatment.

  This statute provides that hospital records of and information pertaining to patients in need of mental treatment at treatment facilities or patients being treated by physicians, certain psychologists, licensed master social workers, or licensed professional counselors be confidential, with certain exceptions.

  Where applicable, DOM is compliant with Miss. Code Ann. § 41-21-97 through implementation of its sensitive data policy, which prohibited the display/disclosure of data related to mental treatment in the former EHR and continues to prohibit the display/disclosure of such data in the Provider Portal.

- **Miss. Code Ann. §§ 41-30-1, et seq.,** entitled *Alcoholism and Alcohol Abuse Prevention, Control and Treatment*, and implemented under the Comprehensive Alcoholism and Alcohol Abuse Prevention, Control and Treatment Act of 1974.

  These statutes provide for confidentiality requirements regarding registration and other records of services by approved treatment facilities that provide treatment or rehabilitation services for alcoholics, whether in-patient, intermediate or out-patient.

  Where applicable, DOM is compliant with Miss. Code Ann. §§ 41-30-1, et seq., through implementation of its sensitive data policy, which prohibited the display/disclosure of alcohol abuse data in the former EHR and continues to prohibit the display/disclosure of such data in the Provider Portal.
Miss. Admin. Code 24-3:9.9, entitled **DMH Principles of Ethical and Professional Conduct**.

This rule provides standards of confidentiality and disclosure regarding information of mental health patients.

Where applicable, DOM is compliant with Miss. Admin. Code 24-3:9.9 through implementation of its sensitive data policy, which prohibited the display/disclosure of data related to mental treatment in the former EHR and continues to prohibit the display/disclosure of such data in the Provider Portal.

Miss. Admin. Code 23-100:3.5 (**Confidentiality of Information**), 3.6 (**Protected Information**), 3.7 (**Release of Information Without Client Consent**), 3.9 (**Safeguarding Confidential Information**), and 23-200:1.1 (**Disclosure of Confidential Information**), regarding the confidentiality of Medicaid beneficiary information.

DOM is in compliance with the above rules.

### 6.3 Assumptions and Dependencies

The following assumptions and dependencies may affect the SMHP as described in this document:

- **Assumptions** - this plan assumes that:
  - The DOM Interoperability Platform Acquisition and Implementation will be available for integration and testing per the schedule listed in the table “Master Milestones/Schedule” above;
  - Certification and implementation of EHR systems will be timely in keeping with the MPIP schedule; and

- **Dependencies** – this plan depends upon:
  - The SLR Upgrades activities listed in the table “Master Milestones/Schedule” above are dependent on Conduent’s ability to meet the timeline dictated by the proposed release schedule.

### 6.4 Participation in the State Health Information Exchange (MS-HIN)

The structure for MS-HIN is set forth in Miss. Code Ann. §§ 41-119-1, *et seq.*, entitled **Health Information Technology Act**, included as Appendix F. The governing body of MS-HIN is the Mississippi Health Information Network Board of Directors. DOM is a member of the MS-HIN Board of Directors and will work in partnership with MS-HIN, providing both leadership and funding support, as appropriate, to assure that Medicaid beneficiaries are best represented and served by MS-HIN.
DOM will work closely with MS-HIN to ensure that each system supports broad, standards-based, interoperable environments to maximize DOM’s investments in these efforts. Having this standards-based foundation allows DOM the greatest flexibility moving forward.

DOM expects the MPIP will encourage and advance the use and number of certified EHR systems available and functioning throughout the State. DOM will participate in MS-HIN and will closely coordinate with MS-HIN to align and leverage resources. Some of the anticipated activities include:

- Coordinating with the MS-HIN to use existing HIT infrastructure and services, when possible;
- Coordinating with MS-HIN to assist providers in achieving MU; and
- Coordinating with the State HIT Director, the Hinds Community College (Workforce Development), and Medicaid providers to disseminate information about MS-HIN, Provider adoption and incentive payments to providers.

### 6.5 Participation in the Sequoia Project (eHealth Exchange)

#### 6.5.1 Alignment with MITA Mission, Goals, and Objectives

CMS expects that the SMHP will be fully aligned with MITA’s mission, goals, and objectives that support the Medicaid mission and goals. MITA and Medicaid’s mission and goals include:

- Adopt industry standards for data exchange;
- Develop seamless, integrated systems;
- Promote flexible, reusable, and adaptable environment;
- Support interoperability, integration, and an open architecture;
- Provide data that is timely, accurate, useable, and easily accessible;
- Support integration of clinical and administrative data;
- Provide performance measurement;
- Promote an enterprise view and efficient/effective data sharing;
- Coordinate with Public Health and other trading partners; and
- Promote secure data exchange.

MITA and Medicaid’s mission and goals are also aligned with federal standards including the FHA and the Sequoia Project (eHealth Exchange) initiative. Furthermore, CMS expects that states will bring their business/technical capabilities in line with MITA 3.0 standards and will advance within the maturity model, at which time states will agree on common data standards, jointly developed business services, and adopt Sequoia Project (eHealth Exchange) standards for interoperability and data.
- **MITA Maturity Level 3 [Clinical Data]**: Data standards are adopted nationally. Shared repositories of data improve efficiency of access and accuracy of data used, resulting in better business process results.

- **MITA Maturity Level 4 [Clinical Data]**: Access to standardized Medicaid clinical data through regional data exchange enhances the decision-making process. With clinical evidence, decisions can be immediate, consistent, and decisive.

- **MITA Maturity Level 5 [National Interoperability / Sequoia Project (eHealth Exchange)]**: Data exchange on a national scale optimizes the decision-making capabilities of the state agency.

DOM has targeted achievement of MITA Maturity Levels 3, 4, and 5 by adopting and aligning with federal standards, including Sequoia Project (eHealth Exchange).

### 6.5.2 Sequoia Project (eHealth Exchange)

The Sequoia Project (eHealth Exchange) comprises the conventions, standards, and shared infrastructure necessary to facilitate the secure and interoperable exchange of electronic health information between organizations over the Internet. Much has already been accomplished to enable the exchange of clinical data, such as summaries between providers. Considerable infrastructure has already been defined at the national level to provide robust security, patient discovery, authentication and authorization, and auditing support. The Sequoia Project (eHealth Exchange) is a critical part of the national health IT agenda to improve population health by making it possible for health information to follow the consumer, be available for clinical decision making, and support appropriate use of health care information beyond direct patient care.

Technical and policy activities over the course of the next several years will expand the value of Sequoia Project (eHealth Exchange) standards, services, and trust fabric and extend the ability to securely exchange health information to a larger audience. This expansion will support providers wishing to achieve MU of CEHRT and qualify for incentives under the HITECH Act.

The ONC, along with federal agencies, state agencies, and HIEs, is facilitating the growth and connectivity to the Sequoia Project (eHealth Exchange). As such, compliance with the Sequoia Project (eHealth Exchange) is an important element of the HIT Roadmap for the State of Mississippi.

The Sequoia Project (eHealth Exchange) can facilitate the exchange of both clinical and administrative data between providers, payers, patients, and other health care professionals. Agencies involved in the Sequoia Project (eHealth Exchange) include CMS, CDC, SSA, DoD, and VA. The Sequoia Project (eHealth Exchange) supports a wide range of use cases for a wide range of users. A list of common use-cases is provided below:

- **Provider to Provider**: Providing the ability to locate providers, send referrals, exchange patient medical history, and send messages for the administrative coordination of care.
• **Provider to Patient**: Providing the ability to send patient reminders, send patient medical history to a Personal Health Record (PHR), and to provide patient medical summaries to patients.

• **Laboratory to Provider**: Providing the ability to send lab results to providers and submit reportable lab results to public health.

• **Provider to Federal Agencies**: Providing the ability to send quality reports, surveillance reports, and more to federal agencies.

• **Provider to Pharmacy**: Providing the ability to send electronic prescriptions for medications and implement drug-drug, drug-allergy, and drug-formulary checks.

• **Provider to Payer**: Providing the ability to check eligibility, submit claims, receive prior authorization, and submit patient information.

Standards-based connectivity initiatives include Sequoia Project (eHealth Exchange) and the Direct Project. The Sequoia Project (eHealth Exchange) and the Direct Project are separate sets of standards and protocols used for information exchange, while eHealth Exchange is a set of software designed to facilitate information exchange. The Sequoia Project (eHealth Exchange) is meant to facilitate inter-HIE data exchange, while the Direct Project is meant to facilitate Intra-HIE data exchange. The Sequoia Project (eHealth Exchange) is used for states or large Provider organizations to connect with the federal government and to communicate among HIEs.

The Direct Project is used for Provider-to-Provider messaging and communication among smaller health care organizations. eHealth Exchange is a federally funded, Open Source software solution that allows for the secure and private exchange of health information. The eHealth Exchange software, referred to as a eHealth Exchange Gateway, is the “on ramp” to the Sequoia Project (eHealth Exchange) network.

### 6.5.3 Sequoia Project (eHealth Exchange) Gateways

In order to connect to the Sequoia Project (eHealth Exchange) organizations can utilize a Sequoia Project (eHealth Exchange) certified Gateway.

DOM has implemented the DOM Interoperability Platform, supporting a Sequoia Project (eHealth Exchange) into the DOM ecosystem. This Interoperability Platform, with full support of standards such as the Sequoia Project (eHealth Exchange), as well as support for other standards and protocols, will ensure coordination with the federal initiatives and connectivity among the providers, stakeholders, HIEs (both in the State of Mississippi and in other states), other State Medicaid agencies, and other entities associated with DOM and the State of Mississippi. DOM is coordinating with MS-HIN to allow for DOM to MS-HIN connectivity, using standards such as the Sequoia Project (eHealth Exchange), to allow for other Mississippi agencies to connect to MS-HIN or DOM and have access to both MS-HIN and DOM.
6.5.4 Connectivity

DOM included requirements for implementing The Sequoia Project (eHealth Exchange) Gateway(s) in the DOM Interoperability Platform, in order to encourage connectivity between DOM, the statewide HIE (e.g., MS-HIN), neighboring HIEs and state agencies/departments, and federal agencies.

The DOM Interoperability Platform, and integrated Sequoia Project (eHealth Exchange) Gateway, can support connectivity and interoperability with MS-HIN and the Provider organizations within the HIE, including the Provider locations receiving EHR Incentive Payments from DOM. DOM has identified several use cases that this connectivity model can support, including:

- Interoperability with the MSDH MIIX System for Medicaid clinical data;
- Medicaid Clinical data exchange with MS-HIN and MS-HIN Medicaid Providers.

This DOM – MS-HIN connectivity can also be utilized to support Medicaid clinical data exchange with:

- Other Mississippi State agencies and stakeholder connectivity and interoperability needs, such as MSDH, the Mississippi Department of Human Services (MDHS), the Mississippi Department of Mental Health (DMH), the Mississippi Department of Rehabilitative Services (MDRS), the Mississippi Department of Corrections (MDOC), the Mississippi Department of Revenue, and the Mississippi Department of Employment Security (MDES);
- Neighboring HIEs such as the Louisiana Statewide HIE, the Arkansas Statewide HIE, the Alabama Statewide HIE;
- Neighboring state agencies such as state Medicaid agencies, State Departments of Health; and
- Federal agencies such as the CMS, the Social Security Administration, the DoD, the VA, the CDC.

The benefits of employing an Interoperability Platform with an integrated Sequoia Project (eHealth Exchange) Gateway(s) Module for DOM are:

- The ability to interact with the aforementioned trading partners (MS-HIN, states, federal agencies, HIEs);
- The ability to leverage a standards-based, modular platform with a compliant Gateway for communication and interoperability;
- The ability to utilize the Sequoia Project (eHealth Exchange) for both clinical and future administrative transactions with multiple trading partners; and
- A decrease in dependence on other entities to provide connectivity and interoperability with health care partners.
Integrating the Healthcare Enterprise Statement and Standards Integration to Drive MITA Compliancy

IHE was formed by the Healthcare Information and Management Systems Society (HIMSS) and the Radiological Society of North America (RSNA). IHE is an initiative by health care professionals to improve the way health care information is shared between systems and organizations around the world for the purpose of improving the overall quality of health care to patients. The mission of IHE is to achieve interoperability of systems through the precise definition of health care tasks, the specification of standards-based communication between systems required to support those tasks, and the testing of systems to determine that they conform to the specifications. IHE promotes the coordinated use of established standards such as DICOM and HL7 to address specific clinical need in support of optimal patient care.

IHE has developed a set of profiles (Integration Profiles) specifying a clear implementation path, including, but not limited to: IT infrastructure, Cardiology, Anatomic Pathology, Eye Care, Laboratory, Patient Care Coordination, Radiology, and Patient Care Devices. Integration Profiles describe how a workflow crossing multiple systems can be achieved using established standards. The Sequoia Project (eHealth Exchange) core services are developed based on IHE profiles, especially IT Infrastructure.

IHE, in general, is a standard way to share EHRs between providers and major HIT or EHR systems that already are IHE compliant. IHE provides a proven solution to resolve health IT interoperability challenges. The following are some core IHE Integration Profiles enabling data sharing among disparate health information systems:

- **PIX/PDQ (Patient Identifier Cross-Referencing and Patient Demographic Query):** Allows systems to query a central master patient index for patient demographics and visit information;
- **XDS (Cross-Enterprise Document Sharing):** Queries/retrieves a list of clinical documents located within a health care community such as RHIO;
- **XDR (Cross-Enterprise Document Reliable Interchange):** Provides document interchange using a reliable messaging system. This permits document interchange between EHRs, PHRs, and other healthcare IT systems in the absence of a document sharing infrastructure such as XDS Registry and Repositories;
- **XCPD (Cross-Community Patient Discovery):** Locates communities for patients and correlates patient identifiers (PID);
- **XCA (Cross-Community Access):** Queries and retrieves data from partner communities;
- **XUA (Cross-Enterprise User Authentication):** Provides a means to communicate claims about the identity of an authenticated principal (user, application, and system) in transactions that cross enterprise boundaries;
- **ATNA (Audit Trail and Node Authentication):** Secures access control via secure nodes and request and retrieve audit logs from external communities;
- CT (Consistent Time): Ensures that system clocks and time stamps of computers in a network are well synchronized; and
- BPPC (Basic Patient Privacy Content): Supports a mechanism to record the patient privacy consent.

EHR systems supporting IHE profiles generally work together better, are easier to implement, and help providers utilize information more efficiently. According to IHE.net, an IHE profile is a technical definition or standard that provides “a common language for purchasers and vendors to discuss the integration needs of healthcare sites and the integration capabilities of healthcare IT products.” To ensure that EHR systems comply with IHE profiles, the IHE hosts “connectathons” to permit vendors to showcase their systems and technology as an IHE compliant vendor.

Many EHR vendors and HIE vendors and suppliers worldwide, including foreign nations, are participating in the IHE workgroups and adopting IHE standards. As participation and adoption of IHE standards and profiles grow, so does the ability for disparate systems and infrastructures to interface, integrate, and communicate data freely.

The State of Mississippi has providers with multiple, diverse EHR systems; therefore, it is critical for DOM to adopt standards, profiles, and an overall interoperable infrastructure to support clinical and administrative data exchange between DOM and the State of Mississippi HIE (MS-HIN) stakeholders and other trading partners. By implementing and integrating standards, profiles, and interoperable infrastructure/technologies (including HL7/IHE/HITSP/Sequoia Project (eHealth Exchange) standards, profiles, and technologies), DOM will drive towards and migrate upwards to the higher levels of MITA and MITA compliance.

### 6.6 Sunset of Medicaid EHR Incentive Program

We plan to continue serving our existing provider population that remain in the EHR Incentive Program by addressing their concerns and questions as they submit their yearly EHR Attestations for Meaningful Use. At this point in time, Mississippi does not plan to increase current staffing requirements. However, as we begin to implement the following goals and objectives, we may find that staffing requirements may need to be adjusted.

#### 6.6.1 Educational Goals / Objectives

Mississippi hosts weekly webinars throughout the year. Each webinar focuses on specific topics relevant to achieving Meaningful Use. Some of our more popular webinars include: Patient Portal Integration, Secure Messaging, and Coordination of Care, which we repeat regularly. Our goal is to host one webinar each week or at least 50 sessions each year (taking into account holidays and slower periods). We find our highest number of attendees join us as Attestation season draws near. During those months, we will often host two webinars per week.

Another strategy we currently use is to publish by-monthly newsletters which emphasize program highlights and regulation updates. Our focus over the past few years has been on Modified Stage 2 requirements with a slight shift toward Stage 3 reporting. At this time we are offering more content that pertains to Stage 3. We plan to use information gained from our Advisory Panels as part of our monthly communications.
6.6.2 Provider Retention Goals and Objectives

The Mississippi Medicaid EHR Incentive Program will reach out to providers across the state and will establish a Provider Advisory Panel by September 30, 2018. This Panel will meet monthly (and more as needed). The purpose of this Advisory Panel is to gain insight and perception as to the needs of our state’s provider community. Our goal is to offer assistance to providers that are participating in both the Medicaid EHR Incentive Program and the Quality Payment Program (QPP). We will use this panel to discover best practices and to identify areas where additional support is needed.

In order to re-engage providers that have dropped out of the program, Mississippi plans to review all payment records. Surveys, much like those used in our recent Environmental Scan, will be sent out and focused email and webinar campaigns will be conducted. We also plan to use information garnered from our Provider Advisory Panel as part of our strategy. Our objective is to make contact with every EP that previously participated but dropped out for various reasons. We want to provide educational resources, best practices, support and encourage re-participation.

Our goals are 1) to re-engage providers that dropped out of our program and, 2) to help providers better utilize their existing technology and/or new technology. The targeted benchmarks are accumulative going forward:

- 25% by end of FFY 2018
- 40% by end of FFY 2019
- 60% by end of FFY 2020

6.6.3 Clinical Quality Measures (CQMs) Goals and Objectives

Currently all CQM reporting is collected from providers as they manually input their reporting data. We are in the beginning steps of talking through requirements that would allow for eCQM collection. We are working closely with our Interoperability team as they collect Clinical data or CCDs from our provider partner systems across the state. We are looking at ways to analyze this data and better use it to accomplish our state’s initiatives in healthcare via a FY2018-19 pilot using population health analytics.
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## Appendix A: Acronyms

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<thead>
<tr>
<th>Acronym</th>
<th>Stands For:</th>
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<tbody>
<tr>
<td>A/I/U</td>
<td>Adopt, Implement or Upgrade</td>
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<tr>
<td>ACO</td>
<td>Accountable Care Organization</td>
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<tr>
<td>ADT</td>
<td>Admission, Discharge, Transfer</td>
</tr>
<tr>
<td>ARRA</td>
<td>American Recovery and Reinvestment Act</td>
</tr>
<tr>
<td>ATNA</td>
<td>Audit Trail and Node Authentication</td>
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<tr>
<td>BPPC</td>
<td>Basic Patient Privacy Content</td>
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<tr>
<td>BIP</td>
<td>Broadband Initiatives Program</td>
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<tr>
<td>BTOP</td>
<td>Broadband Technology Opportunities Program</td>
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<tr>
<td>CAH</td>
<td>Critical Access Hospital</td>
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<tr>
<td>CCD/C-CDA</td>
<td>Continuity of Care Document; Consolidated-Clinical Document Architecture</td>
</tr>
<tr>
<td>CCHIT</td>
<td>Certification Commission for Health Information Technology</td>
</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention</td>
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<tr>
<td>CEHRT</td>
<td>Certified Electronic Health Record Technology</td>
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<td>CDI</td>
<td>Clinical Data Infrastructure</td>
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<tr>
<td>CFHC</td>
<td>Coastal Family Health Center</td>
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<tr>
<td>CMS</td>
<td>Centers for Medicare and Medicaid Services</td>
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<tr>
<td>COTS</td>
<td>Commercial Off the Shelf</td>
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<tr>
<td>CPOE</td>
<td>Computerized Physician Order Entry</td>
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<tr>
<td>CQM</td>
<td>Clinical Quality Measures</td>
</tr>
<tr>
<td>CT</td>
<td>Consistent Time</td>
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<tr>
<td>DMH</td>
<td>Mississippi Department of Mental Health</td>
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<tr>
<td>DOC</td>
<td>Department of Commerce</td>
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<td>DoD</td>
<td>Department of Defense</td>
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<tr>
<td>Acronym</td>
<td>Stands For:</td>
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<tr>
<td>DOM</td>
<td>State of Mississippi Division of Medicaid</td>
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<tr>
<td>e-BEAT</td>
<td>Extension Broadband Education and Adoption Team</td>
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<tr>
<td>EFT</td>
<td>Electronic Funds Transfer</td>
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<tr>
<td>EH</td>
<td>Eligible Hospital</td>
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<tr>
<td>EHR</td>
<td>Electronic Health Record</td>
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<tr>
<td>eMPI</td>
<td>Enterprise Master Patient Index</td>
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<td>EMR</td>
<td>Electronic Medical Record</td>
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<tr>
<td>EP</td>
<td>Eligible Professional</td>
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<tr>
<td>ESB</td>
<td>Enterprise Service Bus</td>
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<tr>
<td>EULA</td>
<td>End User License Agreement</td>
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<tr>
<td>FCC</td>
<td>Federal Communications Commission</td>
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<tr>
<td>FFP</td>
<td>Federal Financial Participation</td>
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<tr>
<td>FFY</td>
<td>Federal Fiscal Year</td>
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<tr>
<td>FHA</td>
<td>Federal Health Architecture</td>
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<tr>
<td>FQHC</td>
<td>Federal Qualified Health Center</td>
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<tr>
<td>HDS</td>
<td>Health Data System</td>
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<tr>
<td>HHS</td>
<td>Department of Health and Human Services</td>
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<tr>
<td>HIE</td>
<td>Health Information Exchange</td>
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<tr>
<td>HIPAA</td>
<td>Health Insurance Portability and Accountability Act</td>
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<tr>
<td>HIT</td>
<td>Health Information Technology</td>
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<tr>
<td>HITECH</td>
<td>Health Information Technology for Economic and Clinical Health</td>
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<tr>
<td>HIX</td>
<td>Health Insurance Exchange</td>
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<td>HL7</td>
<td>Health Level Seven</td>
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<tr>
<td>IAPD</td>
<td>Implementation Advanced Planning Document</td>
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<tr>
<td>ICD-10</td>
<td>International Classification of Diseases</td>
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<tr>
<td>Acronym</td>
<td>Stands For:</td>
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<tr>
<td>IHE</td>
<td>Integrating the Healthcare Enterprise</td>
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<tr>
<td>IT</td>
<td>information technology</td>
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<tr>
<td>iTech</td>
<td>Office of Information Technology Management</td>
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<tr>
<td>ITS</td>
<td>Information Technology Services</td>
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<tr>
<td>LTE</td>
<td>Long Term Evolution</td>
</tr>
<tr>
<td>MBCC</td>
<td>Mississippi Broadband Connect Coalition</td>
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<tr>
<td>MDES</td>
<td>Mississippi Department of Employment Security</td>
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<tr>
<td>MDHS</td>
<td>Mississippi Department of Human Services</td>
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<tr>
<td>MDM</td>
<td>Master Data Management</td>
</tr>
<tr>
<td>MDOC</td>
<td>Mississippi Department of Corrections</td>
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<tr>
<td>MDRS</td>
<td>Mississippi Department of Rehabilitative Services</td>
</tr>
<tr>
<td>MES</td>
<td>Mississippi Enterprise System</td>
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<tr>
<td>MHA</td>
<td>Mississippi Hospital Association</td>
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<tr>
<td>MID</td>
<td>Mississippi Insurance Department</td>
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<tr>
<td>MIIX</td>
<td>Mississippi Immunization Information Exchange System</td>
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<tr>
<td>MITA</td>
<td>Medicaid Information Technology Architecture</td>
</tr>
<tr>
<td>MMIS</td>
<td>Medicaid Management Information System</td>
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<tr>
<td>MPIP</td>
<td>Mississippi Provider Incentive Program</td>
</tr>
<tr>
<td>MS SLR</td>
<td>Mississippi State Level Registry</td>
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<tr>
<td>MSCHIE</td>
<td>Mississippi Coastal Health Information Exchange</td>
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<tr>
<td>MSDH</td>
<td>Mississippi Department of Health</td>
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<tr>
<td>MS-HIN</td>
<td>Mississippi Statewide Health Information Network</td>
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<tr>
<td>MTOM</td>
<td>WS Message Transmission Optimization Mechanism</td>
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<tr>
<td>MU</td>
<td>Meaningful Use</td>
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<tr>
<td>NCPDP</td>
<td>National Council for Prescription Drug Programs</td>
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<tr>
<td>Acronym</td>
<td>Stands For:</td>
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<tr>
<td>NLR</td>
<td>National Level Repository</td>
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<tr>
<td>NPI</td>
<td>National Provider Identifier</td>
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<tr>
<td>NTIA</td>
<td>National Telecommunications and Information Administration</td>
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<tr>
<td>OAT</td>
<td>Office for Advancement of Telehealth</td>
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<tr>
<td>OFPA</td>
<td>Office of Financial and Performance Audit</td>
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<tr>
<td>ONC</td>
<td>Office of the National Coordinator for Healthcare Information Technology</td>
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<tr>
<td>PHR</td>
<td>Personal Health Record</td>
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<tr>
<td>PIX</td>
<td>Patient Identifier Cross-Referencing</td>
</tr>
<tr>
<td>PDQ</td>
<td>Patient Demographic Query</td>
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<tr>
<td>REST</td>
<td>Representational State Transfer</td>
</tr>
<tr>
<td>RFP</td>
<td>Request for Proposals</td>
</tr>
<tr>
<td>RHC</td>
<td>Rural Health Clinic</td>
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<tr>
<td>RHIO</td>
<td>Regional Health Information Organization</td>
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<tr>
<td>SaaS</td>
<td>Software as a Service</td>
</tr>
<tr>
<td>SLR</td>
<td>State Level Registry</td>
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<tr>
<td>SMHP</td>
<td>State Medicaid Health Information Technology Plan</td>
</tr>
<tr>
<td>SOP</td>
<td>Strategic and Operational Plan</td>
</tr>
<tr>
<td>SRA</td>
<td>Security Risk Analysis</td>
</tr>
<tr>
<td>UDDI</td>
<td>Universal Description, Discovery and Integration</td>
</tr>
<tr>
<td>UMMC</td>
<td>University of Mississippi Medical Center</td>
</tr>
<tr>
<td>VA</td>
<td>Veterans Administration</td>
</tr>
<tr>
<td>VLER</td>
<td>Virtual Lifetime Electronic Record</td>
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<tr>
<td>WS-I</td>
<td>Web Services Interoperability</td>
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<td>Acronym</td>
<td>Stands For:</td>
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<tr>
<td>XCA</td>
<td>Cross-Community Access</td>
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<td>XCPD</td>
<td>Cross-Community Patient Discovery</td>
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<tr>
<td>XDR</td>
<td>Cross-Enterprise Document Reliable Interchange</td>
</tr>
<tr>
<td>XDS</td>
<td>Cross-Enterprise Document Sharing</td>
</tr>
<tr>
<td>XSLT</td>
<td>Extensible Stylesheet Language Transformation</td>
</tr>
<tr>
<td>XUA</td>
<td>Cross-Enterprise User Authentication</td>
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### Appendix B: Glossary

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>4010 Format</td>
<td>The current version of the HIPAA electronic transaction standards.</td>
</tr>
<tr>
<td>Adopt, Implement, or Upgrade (A/I/U)</td>
<td>Defined in CMS regulations at 42 CFR 495.302 as (1) Acquire, purchase, or secure access to certified EHR technology; (2) Install or commence utilization of certified EHR technology capable of meeting meaningful use requirements; or (3) Expand the available functionality of certified EHR technology capable of meeting meaningful use requirements at the practice site, including staffing, maintenance, and training, or upgrade from existing EHR technology to certified EHR technology per the ONC EHR certification criteria.</td>
</tr>
<tr>
<td>Allscripts</td>
<td>Vendor providing ePrescribing via the eScript solution with support for drug interactions and contraindications</td>
</tr>
<tr>
<td>Authentication</td>
<td>Authentication is a method or methods employed to prove that the person or entity accessing information has the proper authorization. Generally used to protect confidential information and network or application access.</td>
</tr>
<tr>
<td>Authorization</td>
<td>Authorization is a system established to grant access to information. Authorization also establishes the level of access an individual or entity has to a data set and includes a management component—an individual or individuals must be designated to authorize access and manage access once access is approved.</td>
</tr>
<tr>
<td>Broadband</td>
<td>A medium that can carry multiple signals, or channels of information, at the same time without interference. Broadband Internet connections enable high-resolution videoconferencing and other applications that require rapid, synchronous exchange of data.</td>
</tr>
<tr>
<td>Centers for Disease Control and Prevention (CDC)</td>
<td><a href="http://www.cdc.gov/">Centers for Disease Control and Prevention - http://www.cdc.gov/</a></td>
</tr>
<tr>
<td>Certification Commission for Health Information Technology (CCHIT)</td>
<td>A private not-for-profit organization functioning as an ONC-Authorized Testing and Certification Body of electronic health records.</td>
</tr>
<tr>
<td>Term</td>
<td>Definition</td>
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<tr>
<td>Computerized Physician Order Entry (CPOE)</td>
<td>Computer-based systems that automate and standardize the clinical ordering process in order to eliminate illegible, incomplete, and confusing orders. CPOE systems typically require physicians to enter information into predefined fields by typing or making selections from on-screen menus. CPOE systems often incorporate, or integrate with, decision support systems.</td>
</tr>
<tr>
<td>Conduent</td>
<td>Vendor providing the Medicaid Management Information System and services (MMIS) to provide core administrative capabilities for DOM. Conduent also provides the MS SLR for tracking provider attestations to the MPIP. Previously known as Xerox.</td>
</tr>
<tr>
<td>Continuity of Care Document (CCD); Consolidated-Clinical Document Architecture (C-CDA)</td>
<td>An electronic document exchange standard for sharing patient summary information, including the most commonly needed pertinent information about current and past health status in a form that can be shared by all computer applications, such as Web browsers and EMR/EHR software systems.</td>
</tr>
<tr>
<td>Critical Access Hospital (CAH)</td>
<td>A hospital that is certified to receive cost-based reimbursement from Medicare. The reimbursement that CAHs receive is intended to improve their financial performance and thereby reduce hospital closures.</td>
</tr>
<tr>
<td>Data Warehouse (DW)</td>
<td>A large database that stores information like a data repository but goes a step further, allowing users to access data to perform research-oriented analysis.</td>
</tr>
<tr>
<td>Decision Support System (DSS)</td>
<td>A computer-based information system that supports business or organizational decision-making activities intended to help decision makers compile useful information from a combination of raw data, documents, personal knowledge, or business models to identify and solve problems and make decisions.</td>
</tr>
<tr>
<td>De-identified health information</td>
<td>De-identified health information consists of individual health records with data redacted or edited to prevent it from being associated with a specific individual. See the HIPAA Privacy Rule for de-identification guidelines. The term is defined at 45 C.F.R. § 160.103.</td>
</tr>
<tr>
<td>Direct Project</td>
<td>Provides point-to-point messaging between providers and other healthcare related organizations – <a href="http://directproject.org">http://directproject.org</a></td>
</tr>
<tr>
<td>EA Server</td>
<td>Server enabling existing applications to leverage SOA architectures, J2EE, and CORBA.</td>
</tr>
<tr>
<td>EDIFECIS Certified</td>
<td>EDIFECIS Certified <a href="http://www.edifecs.com/">http://www.edifecs.com/</a></td>
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</tr>
<tr>
<td>Electronic Data Interchange (EDI)</td>
<td>Electronic Data Interchange – The electronic transmission of structured data between organizations.</td>
</tr>
<tr>
<td>Enterprise Master Patient Index (eMPI)</td>
<td>Master Patient Indices link smaller organizational level MPIs together to identify, match, merge, de-duplicate, and clean patient records to create a clear view of a patient’s medical record.</td>
</tr>
<tr>
<td>Electronic Health Record (EHR)</td>
<td>An electronic record of health-related information on an individual that conforms to nationally recognized interoperability standards that can be created, managed, and consulted by authorized clinicians and staff across more than one health care organization.</td>
</tr>
<tr>
<td>Electronic Medical Record (EMR)</td>
<td>An electronic record of health-related information for an individual that can be created, gathered, managed, and consulted by authorized clinicians and staff within one health care organization.</td>
</tr>
<tr>
<td>Envision</td>
<td>Mississippi’s HIPAA compliant Medicaid Management Information System (MMIS) developed by Affiliated Computer Systems (ACS).</td>
</tr>
<tr>
<td>e-prescribing</td>
<td>Practice in which drug prescriptions are entered into an automated data entry system (handheld, PC, or other), rather than handwriting them on paper. The prescriptions can then be printed for the patient or sent to a pharmacy via the Internet or other electronic means. <a href="https://www.cms.gov/eprescribing/">https://www.cms.gov/eprescribing/</a></td>
</tr>
<tr>
<td>Federal Health Architecture (FHA)</td>
<td>A collaborative body composed of several federal departments and agencies, including the Department of Health and Human Services (HHS), the Department of Homeland Security (DHS), the Department of Veterans Affairs (VA), the Environmental Protection Agency (EPA), the United States Department of Agriculture (USDA), the Department of Defense (DOD), and the Department of Energy (DOE). FHA provides a framework for linking health business processes to technology solutions and standards, and for demonstrating how these solutions achieve improved health performance outcomes.</td>
</tr>
<tr>
<td>Federally Qualified Health Center (FQHC)</td>
<td>A health center that receives cost-based reimbursement for Medicare and Medicaid patients as a mechanism to increase primary care services to high risk populations in underserved areas.</td>
</tr>
<tr>
<td>Formulary</td>
<td>A list of medications (both generic and brand names) that are covered by a specific health insurance plan or pharmacy benefit manager (PBM), used to encourage utilization of more cost-effective drugs. Hospitals sometimes use formularies of their own, for the same reason.</td>
</tr>
<tr>
<td>Geocoded Interoperable Population Summary Exchange (GIPSE)</td>
<td>GIPSE is a data format created by the U.S. Centers for Disease Control and Prevention (CDC) to allow the electronic exchange of health condition/syndrome summary data that has been stratified by a number of variables, including geography. GIPSE data will be utilized by public health agencies in the U.S. to conduct situational awareness, including early event detection and monitoring, for potential public health events.</td>
</tr>
<tr>
<td>Grabit</td>
<td>A tool provided by ACS that is able to search, read and download binary files</td>
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<tr>
<td>Term</td>
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<tr>
<td>Health Information Technology (HIT)</td>
<td>The application of information processing involving both computer hardware and software that deals with the storage, retrieval, sharing, and use of health care information, data, and knowledge for communication and decision-making.</td>
</tr>
<tr>
<td>Health Information Technology for Economic and Clinical Health Act (HITECH)</td>
<td>Legislation enacted under Title XIII of the American Recovery and Reinvestment Act (ARRA) of 2009. The purpose of HITECH was to promote spending to expand adoption rates of HIT.</td>
</tr>
<tr>
<td>Health Information Exchange (HIE)</td>
<td>The electronic movement of health-related information among organizations according to nationally recognized standards. Health Information Exchange is a term commonly used to describe a Regional Health Information Organization (RHIO). The notion of HIE is the precursor to RHIO and is used interchangeably when discussing RHIO.</td>
</tr>
<tr>
<td>Health Insurance Exchange (HIX)</td>
<td>As part of the Affordable Care Act (ACA), states are to establish, implement and operate a Health Insurance Exchange by January 1, 2014 that acts as a marketplace for individuals seeking affordable insurance options.</td>
</tr>
<tr>
<td>Health Insurance Portability and Accountability Act of 1996 (HIPAA)</td>
<td>A federal law intended to improve the portability of health insurance and simplify health care administration. HIPAA sets standards for electronic transmission of claims-related information and for ensuring the security and privacy of all individually identifiable health information.</td>
</tr>
<tr>
<td>Health Level 7 (HL7)</td>
<td>HL7 is one of several American National Standards Institute (ANSI)-accredited standards-developing organizations operating in the health care arena. Health Level 7’s domain is clinical and administrative data.</td>
</tr>
<tr>
<td>Healthcare Information Technology Standards Panel (HITSP)</td>
<td>Sponsored by ANSI under a contract from ONC, HITSP is a public/private partnership dedicated to facilitating the harmonization of consensus-based standards necessary to enable the widespread interoperability of health care information in the United States.</td>
</tr>
<tr>
<td>Indian Health Service (HIS)</td>
<td>Indian Health Service - <a href="http://www.ihs.gov/">http://www.ihs.gov/</a></td>
</tr>
<tr>
<td>Integrating the Healthcare Enterprise (IHE)</td>
<td>An initiative by healthcare professionals and industry to improve the way computer systems in healthcare share information. IHE promotes the coordinated use of established standards such as DICOIM and HL7 to address specific clinical needs in support of optimal patient care.</td>
</tr>
<tr>
<td>Interoperability</td>
<td>HIMSS’ definition of interoperability is “ability of health information systems to work together within and across organizational boundaries in order to advance the effective delivery of healthcare for individuals and communities.” For further information, visit HIMSS Interoperability Definition and Background (PDF).</td>
</tr>
<tr>
<td>Java Surveillance Utilization Review System (J-SURS)</td>
<td>A suite of claims-based, data mining software applications designed to identify potentially fraudulent or abusive practices by both those who provide and receive healthcare service.</td>
</tr>
<tr>
<td>Meaningful Use (MU)</td>
<td>Meaningful Use - <a href="https://www.cms.gov/EHRIncentivePrograms/30_Meaningful_Use.asp">https://www.cms.gov/EHRIncentivePrograms/30_Meaningful_Use.asp</a></td>
</tr>
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<td>Term</td>
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<tr>
<td>Medicaid Information Technology Architecture (MITA)</td>
<td>A federal, business-driven initiative that affects the Medicaid enterprise in all states by improving Medicaid program administration, via the establishment of national guidelines for processes and technologies. MITA is a common business and technology vision for state Medicaid organizations that supports the unique needs of each state. <a href="https://www.cms.gov/MedicaidInfoTechArch/">https://www.cms.gov/MedicaidInfoTechArch/</a></td>
</tr>
<tr>
<td>Mississippi Coastal Health Information Exchange (MSCHIE)</td>
<td>The predecessor HIE to MS-HIN.</td>
</tr>
<tr>
<td>Mississippi Coordinated Access Network (MississippiCAN)</td>
<td>A Coordinated Care Program for Mississippi Medicaid beneficiaries to improve access to needed medical services, improve quality care, and improve efficiencies and cost effectiveness.</td>
</tr>
<tr>
<td>Mississippi Department of Human Services (MDHS)</td>
<td>Mississippi Department of Human Service - <a href="http://www.MDHS.state.ms.us/">http://www.MDHS.state.ms.us/</a></td>
</tr>
<tr>
<td>Mississippi Department of Mental Health (DMH)</td>
<td>Mississippi Department of Mental Health - <a href="http://www.dmh.state.ms.us/">http://www.dmh.state.ms.us/</a></td>
</tr>
<tr>
<td>Mississippi Department of Rehabilitation Services (MDRS)</td>
<td>Mississippi Department of Rehabilitation Services - <a href="http://www.mdrs.state.ms.us/">http://www.mdrs.state.ms.us/</a></td>
</tr>
<tr>
<td>Mississippi Division of Medicaid</td>
<td>Mississippi Division of Medicaid - <a href="http://www.medicaid.ms.gov/">http://www.medicaid.ms.gov/</a></td>
</tr>
<tr>
<td>Mississippi EHR Provider Incentive Program</td>
<td>MS EHR PIP - <a href="https://msehrpip.wordpress.com">https://msehrpip.wordpress.com</a></td>
</tr>
<tr>
<td>Mississippi Health Information Network (MS-HIN)</td>
<td>The Mississippi Health Information Exchange.</td>
</tr>
<tr>
<td>Mississippi Information Technology Services (ITS)</td>
<td>Mississippi Information Technology Services - <a href="http://www.its.ms.gov/">http://www.its.ms.gov/</a></td>
</tr>
<tr>
<td>Mississippi Insurance Department (MID)</td>
<td>Mississippi Insurance Department - <a href="http://www.mid.state.ms.us/">http://www.mid.state.ms.us/</a></td>
</tr>
<tr>
<td>Mississippi State Department of Health (MSDH)</td>
<td>Mississippi State Department of Health - <a href="http://www.mdsh.state.ms.us/">http://www.mdsh.state.ms.us/</a></td>
</tr>
<tr>
<td>Nationwide Health Information Network (NwHIN)</td>
<td>The federal government’s program to implement a national interoperable system for sharing electronic medical records or EMRs (a.k.a. electronic health records or EHR). NwHIN (HealtheWay) describes the technologies, standards, laws, policies, programs and practices that enable health information to be shared among health decision makers, including consumers and patients, to promote improvements in health and healthcare. The development of a vision for the NwHIN began more than a decade ago with publication of an Institute of Medicine report, “The Computer-Based Patient Record”. <a href="http://healthit.hhs.gov/portal/server.pt/community/healthit_hhs_gov__nationwide_health_information_network/1142">http://healthit.hhs.gov/portal/server.pt/community/healthit_hhs_gov__nationwide_health_information_network/1142</a></td>
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<tr>
<td>National Coordinator for Health Information Technology (ONC)</td>
<td>Previously referred to as ONCHIT, ONC provides leadership for the development and nationwide implementation of an interoperable health information technology infrastructure to improve the quality and efficiency of health care and the ability of consumers to manage their care and safety. <a href="http://healthit.hhs.gov/portal/server.pt/community/healthit_hhs_gov__home/1204">http://healthit.hhs.gov/portal/server.pt/community/healthit_hhs_gov__home/1204</a></td>
</tr>
<tr>
<td>Personal Health Record (PHR)</td>
<td>An electronic record of health-related information on an individual that conforms to nationally recognized interoperability standards and that can be drawn from multiple sources while being managed, shared, and controlled by the individual.</td>
</tr>
<tr>
<td>Pharmacy Benefit Management (PBM)</td>
<td>A third party administrator of prescription drug programs primarily responsible for processing and paying prescription drug claims. They also are responsible for developing and maintaining the formulary, contracting with pharmacies, and negotiating discounts and rebates with drug manufacturers.</td>
</tr>
<tr>
<td>Physician Quality Reporting Initiative (PQRI)</td>
<td>A voluntary program that provides a financial incentive to physicians and other eligible professionals who successfully report quality data related to services provided under the Medicare Physician Fee Schedule (MPFS).</td>
</tr>
<tr>
<td>Portal</td>
<td>A Web site that offers a range of resources, such as e-mail, chat boards, search engines, and content.</td>
</tr>
<tr>
<td>Prospective Payment System</td>
<td>A payment mechanism for reimbursing hospitals for inpatient health care services in which a predetermined rate is set for treatment of specific illnesses. The system was originally developed by the U.S. federal government for use in treatment of Medicare recipients.</td>
</tr>
<tr>
<td>Provider</td>
<td>A provider is an individual or group of individuals who directly (primary care physicians, psychiatrists, nurses, surgeons, etc) or indirectly (laboratories, radiology clinics, etc) provide health care to patients. In the case of this SMHP and the MPIP, provider refers to both Eligible Professionals (EPs) and Eligible Hospitals (EHs).</td>
</tr>
<tr>
<td>Public Health</td>
<td>Public health is the art and science of safeguarding and improving community health through organized community effort involving prevention of disease, control of communicable disease, application of sanitary measures, health education, and monitoring of environmental hazards.</td>
</tr>
<tr>
<td>Quality Reporting Document Architecture (QRDA)</td>
<td>The emerging quality reporting architecture, based upon the HL7 CDA document.</td>
</tr>
<tr>
<td>Real-Time Innovations (RTI)</td>
<td>A company that develops a middleware solution.</td>
</tr>
<tr>
<td>Regional Extension Center (REC)</td>
<td>An organization that has received funding under the Health Information Technology for Economic and Clinical Health Act to assist health care providers with the selection and implementation of electronic health record technology.</td>
</tr>
</tbody>
</table>

Appendix B: Glossary
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regional Health Information Organization (RHIO)</td>
<td>A health information organization that brings together health care stakeholders within a defined geographic area and governs health information exchange among them for the purpose of improving health and care in that community.</td>
</tr>
<tr>
<td>Rural Health Clinic (RHC)</td>
<td>A clinic certified to receive special Medicare and Medicaid reimbursement, intended to increase primary care services for Medicaid and Medicare patients in rural communities.</td>
</tr>
<tr>
<td>Secure Sockets Layer (SSL)</td>
<td>A cryptographic protocol that enables secure communication over the internet.</td>
</tr>
<tr>
<td>Software as a Service (SaaS)</td>
<td>A business model for software delivery in which software is hosted in the cloud and accessed by users through a client.</td>
</tr>
<tr>
<td>Stakeholder</td>
<td>A stakeholder is any organization or individual that has a stake in the exchange of health information, including health care providers, health plans, health care clearinghouses, regulatory agencies, associations, consumers, and technology vendors.</td>
</tr>
<tr>
<td>Telehealth</td>
<td>The use of telecommunications and information technology to deliver health services and transmit health information over distance. Sometimes called telemedicine.</td>
</tr>
<tr>
<td>Telemedicine</td>
<td>The use of telecommunications and information technology to deliver health services and transmit health information over distance. Sometimes called telehealth.</td>
</tr>
</tbody>
</table>
| Transaction Types (EDI)                        | 270/271 – EDI Healthcare Eligibility/Benefit Inquiry (270) and EDI Healthcare Eligibility/Benefits Response (271)  
276/277/277U – EDI Healthcare Claim Status Request (276) and EDI Healthcare Claim Status Notification (277)  
278 – EDI Healthcare Service Review Information (278)  
820 – EDI Payroll Deducted and other group Premium Payment for Insurance Products (820)  
834 – EDI Benefit Enrollment and Maintenance Set (834)  
835 – EDI Healthcare Claim Payment/Advice Transaction Set  
837P/D/I – EDI Healthcare Claim Transaction Set (837), Professional (P), Dental (D), and Institutional (I) |
<p>| Vendors                                         | Vendors are organizations that provide services and supplies to other organizations. In the context of health information exchange, the term usually refers to technology vendors who provide hardware or software, such as electronic health records, e-prescribing technology, or security software. |
| Veteran’s Affairs                               | Veteran’s Affairs - <a href="http://www.va.gov/">http://www.va.gov/</a>                                                                                                                                               |</p>
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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</thead>
<tbody>
<tr>
<td>Virtual Private Network</td>
<td>Provides secure and remote access to a private Local Area Network via the Internet or other networks.</td>
</tr>
<tr>
<td>Xerox</td>
<td>See Conduent</td>
</tr>
</tbody>
</table>
Appendix C: HIE Readiness Assessment Focus Group Results

The HIE Readiness Assessment was conducted in June 2010 for the Mississippi Department of Information Technology Services (ITS) for its Strategic and Operational Planning (SOP) effort. The assessment included interviews with representatives of 27 facilities across Mississippi that were conducted with a cross section of urban and rural facilities, including both clinics and hospitals. This assessment was aimed primarily at gathering information from hospitals but included certain other entities, such as hospital clinics, FQHCs, and the Indian Tribe.

Two provider focus group meetings were conducted in Mississippi on August 18th and 19th, 2010. The 18th meeting was held in Jackson and had 20 participants representing 12 different providers. The 19th meeting was conducted in Hattiesburg and had 21 participants representing 9 different providers.

Each group was asked the same basic question set. Based on the responses to the basic questions, additional follow up questions were asked for clarification and additional information. The results of each focus group were similar. Therefore, these results are combined and shown as a collective response.

**Question 1 – How many participants are using an Electronic Health Record application?**
- 11 out of 20 in Jackson.
- 12 out of 21 in Hattiesburg.

**Question 2 – What EHR application are you using?**
- Allscripts
- Relay Health
- Greenway
- Epic associated with tertiary hospital
- Practice Works

**Question 3 – How long have you been using the EHR application?**
Most were relatively recent acquisitions with two (2) years being the longest for three (3) providers.

**Question 4 – Describe your experience with EHR technology to date.**
- On All Scripts (3 different responders).
  - Older physicians not as happy as younger physicians as their work flow is altered
  - Of 25 total physicians, 9 are fully using it while the rest are adjusting to the new system
  - One group was dissatisfied and looking to convert to tertiary hospital system
- Greenway user is having a positive experience and sees definite cost savings. No lost charts.
- Billing has become easier.
- Recent move to EPIC, 240+ physicians in locations over southern part of state are using the EHR and the organization could not function without it.
  - Does not know how they would ever go back to paper record, but does not know how to show meaningful use
- Some are using Voice recognition for clinical notes.
- Some physicians are using a point and click system with customized templates
- Customization of templates by each physicians is important
Question 5 – Why did you or why are you considering making the change to an electronic health records system?

- Driven by the fear of lost reimbursement not the incentive dollars
- Doctors concerned about loss of volume which is pay criteria when convert to EMR
- Change for the doctor must be coordinated with hospital EMR so change is not done twice.
- Incentive is nice, not primary driver
- Most would do EMR adoption without incentive because:
  - Improved quality of care
  - Difficult to manage volume of data with paper, they are running out of storage space
  - Federal requirement
  - Access information anywhere
  - Patient safety, easier to read notes and comments, prescription built in, automatic data feeds to different applications
  - Ease of use
  - Needed to recruit new doctors

Question 6 – For those participants without an EHR application, what are your plans?

- Have been looking for a year and hope to make a decision later this year
- Tried one system but it did not integrate with existing practice management system so they are continuing to look
- Five participants indicated they were unfamiliar with EHR applications in general and were looking for assistance (They were introduced to the Regional Extension Center staff at the end of the focus group meeting)

Question 7 – What features are you seeking in an electronic health record application?

- Ease of use
- Product suited to specialty
- Customization to fit the needs of individual doctor or specialty
- Integration with key services like labs
- Legibility leading for improved patient safety
- Customized templates to allow for additional detail information
- Assistance meeting quality metrics
- Improved access to data
- Improved coding features for better billing and collection

Question 8 – What are the primary resistance points for adoption of an EHR application?

- Takes time to learn a new process
- Physicians don’t like information they are getting. It seems template driven with a lot of irrelevant data to wade through to get to the data physicians really need
- Don’t like the templates, no time to customize
- Don’t like the workflow structure
- Medicine by check box, don’t like the built in intelligence
- Change
- Spending too much time looking at a computer and not enough face to face time with the patient
Question 9 – Are you aware of the Medicaid provider incentive program?
Most participants had heard of the incentive program but less than half had any real knowledge of how it worked and what they needed to do to apply. Of those familiar with the program (about 30 percent), they indicated they would apply for Medicaid because it paid more than the Medicare program.

Question 10 – Does the incentive influence affect your decision making about acquiring an EHR application?
Most of the respondents were moving forward without the incentives and a majority was skeptical the incentive program would actually pay them as promised.

Question 11 – When do you think you will apply for stimulus funds?
About half indicated they would apply in 2011. The remainder were unsure when they might apply because they did not know when they would convert to an EHR.

Question 12 – If you apply for Medicaid stimulus funds, Medicaid will be required to verify your eligibility. What would make verification easiest on your practice?
- Know the requirements and expectations from the beginning
- Keep it simple with minimal impact on administrative staff which adds expense
- Educate people on the process and how to meet meaningful use
  - PQRI example of what not to do, took too much time to get results and understand if submission was successful
  - Target audience to include public health
- Use random sampling for checking compliance and audits
- Do not want to do have to complete special data extractions. Follow the normal work flow practices that can be done as part of everyday business
- It should be as electronic as possible

Question 13 – Are you aware of Meaningful Use and what it may require?
- Most participants reported a limited understanding of Meaningful Use
- Most participants reported they were aware Meaningful Use was coming
- Most participants were aware there were quality measures in their future but lacked specifics on them

Question 14 – What is the value of an improved electronic claims submission process?
- Ability to bill every day with shorter turnaround times on reimbursement
- Will improve the throughput success
- Get money faster from Medicaid
- Medicare not impacted due to having set schedule and cutoffs
- Easier to address billing audits
- Billing success based on type of service performed, primary OK, specialty may cause issues
- Coding level is enhanced and good EHR’s can suggest code based on various components
- Documentation is there to help patients
- Helps with correct diagnosis coding
- From HIPPA standpoint, it helps track who is looking at records so there is better privacy and security
Question 15 – What is your experience with Medicaid in Mississippi?

- Do not like time it takes to approve claims. Denial two months after the treatment causes financial problems for clinics
- Process OK, reimbursement rate is too low
- Provider enrollment takes too long, some clinics not aware they can back bill new enrollments
- Deal with CHIPS and Medicaid, you do not ever know what to expect out of them. They are unpredictable
- Call center at Medicaid does not have the intelligence to deal with issues on phone. Frustrates the clinic
- Must ask for extended visits for kids and prior authorizations. Creates a lot of extra work for physicians

Question 16 – How many have heard about the Share Point EHR being offered by the Division of Medicaid in Mississippi

- 2 of 21 in Hattiesburg
- 5 of 20 in Jackson

Participant questions for the Moderator

Participants were provided an opportunity to ask questions of the moderator. The questions included:

- What is the Medicaid six year span for incentive payments and what is the relationship to relation to Meaningful Use?
- How do submit claims in the future without being ICD10 compliant? Does it require providers to have a certified EMR?
- Can you explain the Medicaid and Medicare incentive and disincentive programs?
- Are private payers incenting EMR adoption as well as Medicaid?
- Incentives not helpful if providers do not have the money to invest in EHR up front. How can Medicaid help financially strapped doctors get the money to get the technology
- Need to provide doctors a system to help doctors understand process and options
- States could tack on additional requirements for meaningful use. Is Mississippi planning on doing that?
- How would I find out what program I should choose and how do I apply for the incentives?
# Appendix D: Mississippi Hospital Association – IT Survey

<table>
<thead>
<tr>
<th>Hospital Name</th>
<th>Health Information System (HIS)</th>
<th>Electronic Health Record</th>
<th>Computerized Physician Order Entry</th>
<th>Lab Information System</th>
<th>Radiology Information System</th>
<th>Picture Archiving and Comm. System</th>
<th>Emergency Department</th>
<th>Pharmacy</th>
<th>Document Imaging</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baptist Memorial Hospital Booneville</td>
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<tr>
<td>Baptist Memorial Hospital Golden Triangle</td>
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<td></td>
<td></td>
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<tr>
<td>Baptist Memorial Hospital Union County</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
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<td>yes</td>
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### Mississippi Hospital Association – IT Survey

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Appendix E: DOM Medicaid Provider Survey Results

Mississippi Division of Medicaid

Provider Survey Results

The Medicaid Eligible Provider survey was launched in July of 2010 and consisted of a multi-part questionnaire that was made available online through the Division of Medicaid website and the MMIS website through September 2010. The questionnaire consisted of 22 questions, both in multiple choice and text entry format, concerning the present and planned use of health information technology among Eligible Professionals in the State. Following are the results of the survey:

1. In which county is your primary practice located? (Select County from drop-down list)

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<th>Alcorn</th>
<th>Amite</th>
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<th>Benton</th>
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<th>Holmes</th>
<th>Humphreys</th>
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<th>Itawamba</th>
<th>Jackson</th>
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| County          | Lamar | Lauderdale | Lawrence | Leake | Lee | Leffore | Lincoln | Lowndes | Madison | Marion | Marshall | Monroe | Montgomery | Neshobe | Newton | Noxubee | Oktibbeha | Panola | Pearl River | Perry | Pike | Pontotoc | Prentiss | Quitman | Rankin | Scott | Sharkey | }
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| County          | Simpson | Smith | Stone | Sunflower | Tallahatchie | Tate | Tippah | Trishomingo | Tunica | Union | Wallahall | Warren | Washington | Wayne | Webster | Wilkinson | Winston | Yalobusha | Yazoo | Out of State | Response Count |
|-----------------|---------|-------|-------|-----------|-------------|------|-------|-------------|--------|-------|-----------|--------|-------------|-------|---------|------------|---------|----------|-------|--------------|----------------|---|
| Number of Providers | 0      | 1     | 0     | 1         | 0           | 2    | 0     | 1           | 1      | 2     | 1         | 0      | 0           | 0     | 1       | 0          | 0       | 0        | 0     | 1            | 94              | 0  |

2. Please enter your contact information or that of your designee.

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4. Which of the following software products or services are you currently using?

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5. Which of your current software products or services are certified?

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### 6. Which of the following software products or services do you plan to add or upgrade to meet the EHR certification requirements?

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**answered question: 94**  
**skipped question: 8**

### 7. Your software or services are provided by:

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<td>On line Commercial Service</td>
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<td>Custom developed software</td>
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</table>

**answered question: 94**  
**skipped question: 8**

### 8. Please provide your software vendor/product information: (If outsourced, please include service bureau name)

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**answered question: 84**
### 9. What is the cost range for your planned software upgrades? (Select amounts from drop-down list)

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**Question Totals**

- Answered question: 81
- Skipped question: 21

### 10. Does your practice exchange or plan to exchange health information with the following?

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<td>Pharmacies</td>
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<td>Other Physicians</td>
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**Question Totals**

- Answered question: 94
- Skipped question: 8

### 11. Does your practice use or plan to use Telemedicine?

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<tr>
<th>Answer Options</th>
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<tbody>
<tr>
<td>Providing care</td>
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<tr>
<td>Consultation with other physicians or hospitals</td>
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<td>60</td>
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**Question Totals**

- Answered question: 88
- Skipped question: 8
### View patient information at home

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<tbody>
<tr>
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**Answered question:** 87  
**Skipped question:** 15

### Does your practice use computers in the exam room?

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**Answered question:** 94  
**Skipped question:** 8

### What are your practice specialties?

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**Answered question:** 90  
**Skipped question:** 12
14. Please estimate the percentage of services by payer type: (Total should equal 100%) (Select percentage from drop-down list)

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Appendix E: DOM Medicaid Provider Survey Results

Page 149
15. What is your level of interest in the following:

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<tr>
<th>Answer Options</th>
<th>High</th>
<th>Medium</th>
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<tbody>
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<td>Receiving updates on EHR information</td>
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<tr>
<td>Training on EHR Implementation</td>
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Answered question 94
Skipped question 8

16. Do you plan to apply for the Medicaid Provider Incentive Payments for Implementing EHR technology?

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<thead>
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<tr>
<td>No</td>
<td>17.0%</td>
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</table>

If yes, in what year do you plan to apply (2011 - 2016):

Answered question 94
Skipped question 8

17. If you plan to apply for the Medicaid Provider Incentive Payments, in which state do you plan to apply? (Select state from drop-down list)

<table>
<thead>
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<th>State</th>
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<tbody>
<tr>
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Answered question 83
Skipped question 19
**18.** Do you plan to apply for the Medicare Provider Incentive Payments for implementing EHR technology?

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<tr>
<td>No</td>
<td>26.6%</td>
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</table>

If yes, in what year do you plan to apply (2011 - 2014)

**answered question** 94
**skipped question** 8

**19.** If this practice is part of a group practice, how are your locations interconnected? (Bandwidths are ranges with the maximum bandwidth shown) (Check all that apply - multiple choices per row are allowed)

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<td>7</td>
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</tbody>
</table>

Please identify the hospital(s)

**answered question** 46
**skipped question** 56

**20.** If your practice electronically exchanges information with a hospital, what type of connection does your practice use? (Bandwidths are ranges with the maximum bandwidth shown) (Check all that apply - multiple choices per row are allowed)

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Please identify the hospital(s)

**answered question** 46
**skipped question** 56

**21.** What types of Internet services and bandwidths does your practice currently use? (Bandwidths are ranges with the maximum bandwidth shown) (Check all that apply - multiple choices per row are allowed)

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Please identify the hospital(s)

**answered question** 94
**skipped question** 8

**22.** What types of Internet services and maximum bandwidths are available to your practice location? (One choice per row for all rows)

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Please identify the hospital(s)

**answered question** 94
**skipped question** 8
Appendix F: Health Information Technology Act

Miss. Code Ann. § 41-119-1
This chapter shall be known and may be cited as the "Health Information Technology Act."

Miss. Code Ann. § 41-119-3
The Mississippi Health Information Network is a public-private partnership for the benefit of all of the citizens of this state.

Miss. Code Ann. § 41-119-5
(1) The Mississippi Health Information Network is established, and is referred to in this chapter as the "MS-HIN."

(2) The MS-HIN shall be governed by a board of directors (MS-HIN board) consisting of eleven (11) members. The membership of the MS-HIN board shall reasonably reflect the public-private and diverse nature of the MS-HIN.

(3) The membership of the MS-HIN board of directors shall consist of the following:

   (a) The Governor shall appoint one (1) member of the MS-HIN board of directors, who shall be a representative of a health insurance carrier in Mississippi with knowledge of information technology, to serve an initial term of three (3) years;

   (b) The State Board of Health shall appoint one (1) member of the MS-HIN board of directors, who shall be a representative of a Mississippi hospital with knowledge of information technology, to serve an initial term of three (3) years;

   (c) The Mississippi State Medical Association shall appoint a member of the MS-HIN board of directors, who shall be a licensed physician, to serve an initial term of three (3) years;

   (d) The Primary Health Care Association shall appoint a member of the MS-HIN board of directors to serve an initial term of one (1) year;

   (e) The Delta Health Alliance shall appoint a member of the MS-HIN board of directors to serve an initial term of four (4) years;

   (f) The Information and Quality Health Care-Mississippi Coastal Health Information Exchange (MCHIE) shall appoint a member of the MS-HIN board of directors to serve an initial term of one (1) year;

   (g) The State Board of Health shall appoint a member of the MS-HIN board of directors who shall be an employee of the State Department of Health to serve an initial term of one (1) year;

   (h) The Mississippi Board of Information Technology Services shall appoint a member of the MS-HIN board of directors to serve an initial term of two (2) years;

   (i) The Mississippi Board of Mental Health shall appoint a member of the MS-HIN board of directors who shall be an employee of the Department of Mental Health to serve an initial term of four (4) years;

   (j) The University of Mississippi Medical Center shall appoint a member of the MS-HIN board of directors to serve an initial term of two (2) years; and

   (k) The Division of Medicaid shall appoint a member of the MS-HIN board of directors who shall be an employee of the Division of Medicaid to serve an initial term of two (2) years.

Initial terms shall expire on June 30 of the appropriate year, and subsequent appointments shall be made by the appointing entity for terms of four (4) years. Members may be reappointed.

(4) No state officer or employee appointed to the MS-HIN board or serving in any other capacity for the MS-
(5) The chairperson of the MS-HIN board shall be elected by a majority of the members appointed to the MS-HIN board.

(6) The MS-HIN board is authorized to conduct its business by a majority of a quorum. A quorum is six (6) members of the MS-HIN board.

(7) The MS-HIN board may adopt bylaws for its operations, including, but not limited to, the election of other officers, the terms of officers, and the creation of standing and ad hoc committees.

**Miss. Code Ann. § 41-119-7**

(1) In furtherance of the purposes of this chapter, the MS-HIN shall have the following duties:

   (a) Initiate a statewide health information network to:

      (i) Facilitate communication of patient clinical and financial information;

      (ii) Promote more efficient and effective communication among multiple health care providers and payers, including, but not limited to, hospitals, physicians, nonphysician providers, third-party payers, self-insured employers, pharmacies, laboratories and other health care entities;

      (iii) Create efficiencies by eliminating redundancy in data capture and storage and reducing administrative, billing and data collection costs;

      (iv) Create the ability to monitor community health status;

      (v) Provide reliable information to health care consumers and purchasers regarding the quality and cost-effectiveness of health care, health plans and health care providers; and

      (vi) Promote the use of certified electronic health records technology in a manner that improves quality, safety, and efficiency of health care delivery, reduces health care disparities, engages patients and families, improves health care coordination, improves population and public health, and ensures adequate privacy and security protections for personal health information;

   (b) Develop or design other initiatives in furtherance of its purpose; and

   (c) Perform any and all other activities in furtherance of its purpose.

(2) The MS-HIN board is granted all incidental powers to carry out its purposes and duties, including the following:

   (a) To appoint an executive director, who will serve at the will and pleasure of the MS-HIN board. The qualifications and employment terms for the executive director shall be determined by the MS-HIN board;

   (b) To adopt, modify, repeal, promulgate, and enforce rules and regulations to carry out the purposes of the MS-HIN;

   (c) To establish a process for hearing and determining case decisions to resolve disputes under this chapter or the rules and regulations promulgated under this chapter among participants, subscribers or the public;

   (d) To enter into, and to authorize the executive director to execute contracts or other agreements with any federal or state agency, any public or private institution, or any individual in carrying out the provisions of this chapter; and

   (e) To discharge other duties, responsibilities, and powers as are necessary to implement the provisions of this chapter.
(3) The executive director shall have the following powers and duties:

(a) To employ qualified professional personnel as required for the operation of the MS-HIN and as authorized by the MS-HIN board;

(b) To administer the policies of the MS-HIN board; and

(c) To supervise and direct all administrative and technical activities of the MS-HIN.

(4) The MS-HIN shall have the power and authority to accept appropriations, grants and donations from public or private entities and to charge reasonable fees for its services. The revenue derived from grants, donations, fees and other sources of income shall be deposited into a special fund that is created in the State Treasury and earmarked for use by the MS-HIN in carrying out its duties under this chapter.

Miss. Code Ann. § 41-119-9
(1) All members of the MS-HIN board shall not be subject to and are immune from claim, suit, liability, damages or any other recourse, civil or criminal, arising from any act or proceeding, decision or determination undertaken, performed or reached in good faith and without malice by any such member or members acting individually or jointly in carrying out the responsibilities, authority, duties, powers and privileges of the offices conferred by law upon them under this chapter, or any other state law, or duly adopted rules and regulations of the aforementioned committees, good faith being presumed until proven otherwise, with malice required to be shown by a complainant. All employees and staff of the MS-HIN, whether temporary or permanent, shall enjoy the same rights and privileges concerning immunity from suit otherwise enjoyed by state employees under the Mississippi Constitution of 1890 and Section 11-46-1 et seq.

(2) The MS-HIN is not a health care provider and is not subject to claims under Sections 11-1-58 through 11-1-62. No person who participates in or subscribes to the services or information provided by the MS-HIN shall be liable in any action for damages or costs of any nature, in law or equity, that result solely from that person’s use or failure to use MS-HIN information or data that were imputed or retrieved in accordance with the rules or regulations of the MS-HIN. In addition, no person will be subject to antitrust or unfair competition liability based on membership or participation in the MS-HIN, which provides an essential governmental function for the public health and safety.

Miss. Code Ann. § 41-119-11
(1) All persons providing information and data to the MS-HIN shall retain a property right in that information or data, but grant to the other participants or subscribers a nonexclusive license to retrieve and use that information or data in accordance with the rules or regulations promulgated by the MS-HIN board and in compliance with the provisions of the Health Insurance Portability and Accountability Act of 1996, Public Law 104-191.

(2) Patients desiring to obtain a copy of their personal medical record or information are to request the copy from the health care provider who is the primary source of the information, and the MS-HIN shall not be required to provide this information directly to the patient.

(3) All processes or software developed, designed or purchased by the MS-HIN shall remain its property subject to use by participants or subscribers in accordance with the rules and regulations promulgated by the MS-HIN board.

Miss. Code Ann. § 41-119-13
(1) The MS-HIN board shall by rule or regulation ensure that patient specific health information be disclosed only in accordance with the provisions of the Health Insurance Portability and Accountability Act of 1996, Public Law 104-191, which governs the electronic transmission of that information.

(2) Patient specific health information and data of the MS-HIN shall not be subject to the Federal Freedom of Information Act, Mississippi Open Records Act (Section 25-61-1 et seq.) nor to subpoena by any court. That information may only be disclosed by consent of the patient or in accordance with the MS-HIN board’s rules, regulations or orders.

(3) Notwithstanding any conflicting statute, court rule or other law, the data in the network shall be
confidential and shall not be subject to discovery or introduction into evidence in any civil action. However, information and data otherwise discoverable or admissible from original sources are not to be construed as immune from discovery or use in any civil action merely because they were provided to the MS-HIN.

(4) Submission of information to and use of information by the State Department of Health shall be considered a permitted disclosure for uses and disclosures required by law and for public health activities under the Health Insurance Portability and Accountability Act and the privacy rules promulgated under that act.

(5) Any violation of the rules or regulations regarding access or misuse of the MS-HIN health information or data shall be reported to the Office of the Attorney General, and shall be subject to prosecution and penalties under state or federal law.

Miss. Code Ann. § 41-119-15
For the purposes of this chapter, the following terms shall be defined as provided in this section:

(a) "Electronic health records" or "EHR" means electronically maintained clinical and demographic information, used by a meaningful EHR user.

(b) "Health information technology" or "HIT" means the equipment, software and networks to be used by a meaningful EHR user.

(c) "Acquisition" of HIT systems or other computer or telecommunications equipment or services means the purchase, lease, rental or acquisition in any other manner of HIT systems or any other computer or telecommunications equipment or services used exclusively for HIT.

(d) "Meaningful EHR user" means an eligible professional or eligible hospital that, during the specified reporting period, demonstrates meaningful use of certified EHR technology in a form and manner consistent with certain objectives and measures presented in applicable federal regulations as amended or adopted. These objectives and measures shall include the use of certified EHR.

(e) "Entity" means and includes all the various state agencies, officers, departments, boards, commissions, offices and institutions of the state, but does not include any agency financed entirely by federal funds.

Miss. Code Ann. § 41-119-17
(1) Before the acquisition of any HIT system, an entity shall provide MS-HIN, at a minimum, description, purpose and intent of the proposed service or system, including a description and specifications of the ability to connect to MS-HIN.

(2) Where existing entities can be used to provide the proposed HIT system, in whole or in part, the submission shall include letters of commitment, memoranda of agreements, or other supporting documentation.

(3) The MS-HIN shall review proposals for acquisition of HIT systems for the purposes contained in Section 41-119-7, and provide guidance to entities including collaborative opportunities with MS-HIN members.

(4) Any acquisition of an HIT system that was approved by the Mississippi Department of Technology Services before April 28, 2010, is exempt from the requirements of Section 41-119-15 and this section.

Miss. Code Ann. § 41-119-19
The Legislative Audit Committee (PEER) shall develop and make a report to the Chairmen of the Senate and House Public Health and Welfare/Medicaid Committees regarding the following electronic health records (EHR) system items:

(a) Evaluate the Request for Proposals (RFP) for the implementation and operations services for the Division of Medicaid and the University Medical Center electronic health records system and e-prescribing system for providers;

(b) Evaluate the proposed expenditures of the Mississippi Division of Medicaid (DOM) and the University Medical Center (UMC) regarding electronic health information;
(c) Evaluate the use of American Recovery and Reinvestment Act (ARRA) funds for electronic health records system implementation in the State of Mississippi; and

(d) Evaluate the progress in implementing the electronic health records system in the State of Mississippi.

The PEER Committee shall make its report on or before December 1, 2014, including any recommendations for legislation.

Miss. Code Ann. § 41-119-21
Sections 41-119-1 through 41-119-21 shall stand repealed on July 1, 2019.
### Appendix G: Calculators

#### G1. Hospital EHR Patient Volume Calculator (Revised 2013) – Form 2552-96

**Mississippi Division of Medicaid**  
**Mississippi Provider Incentive Payment Program**

*White Areas are for data input*

<table>
<thead>
<tr>
<th>Hospital:</th>
<th>NPI:</th>
</tr>
</thead>
</table>

*Grey Areas are calculated results*

#### Average Length of Stay - 2552-96 Cost Report

<table>
<thead>
<tr>
<th>Measure</th>
<th>Cost Report Data Source</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Hospital Days</td>
<td>w/s S-3 part I, col. 6, lines 1,2,6,7,8,9,10</td>
<td>0</td>
</tr>
<tr>
<td>Total Hospital Discharges</td>
<td>w/s S-3 part I, col. 15, lines 1,2,6,7,8,9,10</td>
<td>0</td>
</tr>
</tbody>
</table>

**Average Length of Stay - 2552-96 Cost Report**

0.0

#### Patient Volume Calculation

**Inpatients - POS Code 21 - Discharges**

<table>
<thead>
<tr>
<th>Medicaid Primary Payer</th>
<th>Data Source - 2552-96 Cost Report</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>w/s S-3 part I, col. 15, lines 1,2,6,7,8,9,10</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Discharges</td>
<td>w/s S-3 part I, col. 14, lines 1,2,6,7,8,9,10</td>
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<td>0</td>
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</tbody>
</table>

**Medicaid Secondary Payer**

<table>
<thead>
<tr>
<th>Medicaid Primary Payer</th>
<th>Data Source - 2552-96 Cost Report</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>w/s S-3 part I, col. 14, lines 1,2,6,7,8,9,10</td>
<td></td>
<td>0</td>
</tr>
</tbody>
</table>
### Primary Payer - Discharges

<table>
<thead>
<tr>
<th></th>
<th>Data Source</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medicare</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Third Party</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total POS 21 Discharges</strong></td>
<td></td>
<td><strong>0</strong></td>
<td><strong>0</strong></td>
</tr>
</tbody>
</table>

### Emergency Room - POS Code 23 - Discharges

#### Medicaid Primary Payer

<table>
<thead>
<tr>
<th>All Patients</th>
<th>Data Source</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Payers</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Medicaid Primary Payer</td>
<td></td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

#### Medicaid Secondary Payer

<table>
<thead>
<tr>
<th>Primary Payer</th>
<th>Data Source</th>
<th>Medicaid</th>
<th>Total</th>
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<tbody>
<tr>
<td>Medicare</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Third Party</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total POS 23 Discharges</strong></td>
<td></td>
<td><strong>0</strong></td>
<td><strong>0</strong></td>
</tr>
</tbody>
</table>

**Total Discharges and Encounters for SLR Application**: 0

**Medicaid Percentage**: 0.0%

**Notes:**

- Hospital Patient Encounters are based on discharge data from both the Inpatient (POS Code 21) and Emergency Room (POS Code 23).
- Hospital must have a minimum of 10 percent Medicaid Patient Volume to qualify for the Medicaid Incentive Payment.
- Hospital Patient Volumes are from the prior federal fiscal year.

1. Medicaid Primary Payer Encounters for both the inpatient and emergency room are required. Medicaid primary payers include Medicaid and Mississippi CAN.
Medicaid Secondary Payer Encounters are optional (if Medicaid Secondary Payer encounters are included, then both inpatient and emergency room discharges must be used). Medicaid Secondary Payer Encounters include Medicare and third party payers when Medicaid is responsible for the copayment.

2 Supporting Documentation: (Must be attached to the application)
   a. Inpatient (POS 21) Discharges - Cost Reports from identified data locations.
   b. Emergency Room (POS 23) Discharges - Billing management reports

3 Inclusions in Medicaid Encounter (Discharges) Counts:
   a. Encounters include a Medicaid Eligible patient (regardless of payment Liability) New in 2013
   b. Encounters paid through the Mississippi CAN program

4 Exclusions from Medicaid Encounter (Discharges) Counts:
   a. Encounters not resulting in a payment by Medicaid
   b. All CHIP Encounters
   c. Emergency Room encounters that result in admission to the hospital

5 Each Emergency room visit will count as one encounter. (See 4.c. - Patients discharges into the hospital can’t be included in the patient discharges.)

G2. Hospital EHR Patient Volume Calculator (Revised 2013) – Form 2552-10

| Mississippi Division of Medicaid                        |
| Mississippi Provider Incentive Payment Program           |
| White Areas are for data input                          |
| Hospital:  | NPI:               |
| Grey Areas are calculated results                       |
| Average Length of Stay Calculation - 2552-10 Cost Report |
### Patient Volume Calculation

#### Inpatients - POS Code 21 - Discharges

<table>
<thead>
<tr>
<th>Measure</th>
<th>Cost Report Data Source</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Hospital Days</td>
<td>w/s S-3 part I, col. 8, lines 1,2,8,9,10,11,12</td>
<td>0</td>
</tr>
<tr>
<td>Total Hospital Discharges</td>
<td>w/s S-3 part I, col. 15, lines 1,2,8,9,10,11,12</td>
<td>0</td>
</tr>
</tbody>
</table>

**Average Length of Stay - 2010 Cost Report Year**

0.0

#### Patient Volume Calculation

**Medicaid Primary Payer (Required)(1)**

<table>
<thead>
<tr>
<th>Data Source - 2552-10 Cost Report</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discharges</td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/s S-3 part I, col. 15, lines 1,2,8,9,10,11,12</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Medicaid Primary Payer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/s S-3 part I, col. 14, lines 1,2,8,9,10,11,12</td>
<td></td>
<td>0</td>
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</table>

**Medicaid Secondary Payer - (Optional)(1)**

<table>
<thead>
<tr>
<th>Primary Payer - Discharges</th>
<th>Data Source</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medicare</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Third Party</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Total POS 21 Discharges: 0

**Emergency Room - POS Code 23 - Discharges**

<table>
<thead>
<tr>
<th>Medicaid Primary Payer - (Required)(1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Patients</td>
</tr>
<tr>
<td>Discharges</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Medicaid Primary Payer</td>
</tr>
</tbody>
</table>
## Medicaid Secondary Payer - (Optional)(1)

<table>
<thead>
<tr>
<th>Primary Payer</th>
<th>Data Source</th>
<th>Medicaid</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medicare</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Third Party</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

### Notes:

*Hospital Patient Encounters are based on discharge data from both the Inpatient (POS Code 21) and Emergency Room (POS Code 23).*

*Hospital must have a minimum of 10 percent Medicaid Patient Volume to qualify for the Medicaid Incentive Payment.*

*Hospital Patient Volumes are from the prior federal fiscal year.*

1. Medicaid Primary Payer Encounters for both the inpatient and emergency room are required. Medicaid primary payers include Medicaid and Mississippi CAN.

Medicaid Secondary Payer Encounters are optional (if Medicaid Secondary Payer encounters are included, then both inpatient and emergency room discharges must be used) Medicaid Secondary Payer Encounters include Medicare and third party payers when Medicaid is responsible for the copayment.

2. Supporting Documentation: (Must be attached to the application)
   a. **Inpatient (POS 21) Discharges** - Cost Reports from identified data locations
   b. **Emergency Room (POS 23) Discharges** - Billing management reports

3. Inclusions in Medicaid Encounter (Discharges) Counts:
   a. Encounters include a Medicaid Eligible patient (regardless of payment Liability) **New in 2013**
   b. Encounters paid through the Mississippi CAN program

4. Exclusions from Medicaid Encounter (Discharges) Counts:
   a. Encounters not resulting in a payment by Medicaid
b. All CHIP Encounters

5. Emergency Room encounters that result in admission to the hospital

Each Emergency room visit will count as one encounter. (See 4.c. - Patients discharges into the hospital can’t be included in the patient discharges.)


<table>
<thead>
<tr>
<th>Eligible Professional - Medicaid Percentage Calculation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Wholly Areas require provider input</strong></td>
</tr>
<tr>
<td><strong>Provider Name:</strong> Dr. Ben Dover</td>
</tr>
<tr>
<td><strong>NPI:</strong> 1234567890</td>
</tr>
<tr>
<td><strong>Payer Group Name:</strong></td>
</tr>
<tr>
<td><strong>NPI:</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medicaid Qualifying Period</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Period Start Date (3):</strong> 7/1/2014</td>
</tr>
<tr>
<td><strong>Period End Date (3):</strong> 8/31/2014</td>
</tr>
<tr>
<td><strong>Must begin on the first day of a month</strong></td>
</tr>
<tr>
<td><strong>90-day period from previous calendar year (cy 2014)</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Name of Patient Management System</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Appointment and billing management</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Encounters</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Medicaid Encounters / All Payers</strong></td>
</tr>
<tr>
<td><strong>Medicaid</strong></td>
</tr>
<tr>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>All Payer Encounters</td>
</tr>
<tr>
<td>Medicaid Encounters</td>
</tr>
<tr>
<td><em>(Medicaid FFS, MS CAN, Magnolia, Medicare Part B, United Health Care (non-commercial))</em></td>
</tr>
<tr>
<td><strong>Total Encounters used in Application</strong></td>
</tr>
<tr>
<td><strong>Medicaid Percentage</strong></td>
</tr>
</tbody>
</table>
### Hospital One Time Payment Calculation

**Calculation of Medicaid Electronic Health Records (EHR) Incentive Payment using 2552-96 Cost Report**

This Payment Calculation was approved by CMS on 06/13/2011

<table>
<thead>
<tr>
<th>Hospital</th>
<th>NPI</th>
</tr>
</thead>
</table>

*White Areas are for data input from your Cost Reports*

*Grey Areas are calculated by the MS SLR application - Do not change*

The overall "EHR" amount is the sum over 4 years of (a) the base amount of $2,000,000 plus (b) the discharge related amount defined as $200 for the 1,150 through the 23,000 discharge for the first payment year then a pro-rated amount of 75% in yr 2, 50% in yr 3, and 25% in yr 4

For years 2-4 the rate of growth is assumed to be the previous 3 years' average.
### Step 1: Compute the average annual growth rate over 3 years using previous Medicare cost reports.

Per the Medicare cost report, worksheet S-3, part I, line 12, column 15 - Total discharges

<table>
<thead>
<tr>
<th>Cost Report years used for one time calculations</th>
<th>PY</th>
<th>CY</th>
<th>Increase</th>
<th>Growth</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fiscal Year</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>0.00%</td>
</tr>
<tr>
<td>Fiscal Year</td>
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<td>0</td>
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<td>0.00%</td>
</tr>
<tr>
<td>Fiscal Year</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.00%</td>
</tr>
</tbody>
</table>

Enter most current Cost Report year used for Steps 2 - 6.

**Total Percent - Increase/(Decrease):** 0.0%

**Divided by 3 years:** 3

**The average annual growth rate over 3 years:** 0.00%

### Step 2: Compute total discharge related amount using proper transition factors

> discharges are capped at 23,000 each year

**INPUT FY total Discharges from worksheet S-3, part I, line 12, column 15:** 0 | 0

<table>
<thead>
<tr>
<th>Discharges</th>
<th>Total</th>
<th>Allowable</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
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<td>0</td>
<td>$0</td>
</tr>
<tr>
<td>Year 2</td>
<td>0</td>
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<td>$0</td>
</tr>
<tr>
<td>Year 3</td>
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<td>$0</td>
</tr>
<tr>
<td>Year 4</td>
<td>0</td>
<td>0</td>
<td>$0</td>
</tr>
</tbody>
</table>

**Total 4 year discharge related amount:** $0

### Step 3: Compute the initial amount for 4 years

<table>
<thead>
<tr>
<th>Years 1 - 4 base amount of $2,000,000 per year</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td>$2,000,000</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Years 1-4 discharge related amount (step 2)</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
</tr>
</tbody>
</table>

**Aggregate EHR amount for 4 years:** $2,000,000 | $2,000,000 | $2,000,000 | $2,000,000

### Step 4: Apply Transition Factor

| $2,000,000 | $1,500,000 | $1,000,000 | $500,000 |

### Step 5: Compute the overall EHR amount for 4 years

**$5,000,000**
### Step 6: Computation of Medicaid Share from the Medicare cost report (2552-96 Cost Report)

\[
\text{Medicaid Share} = \frac{\text{estimated Medicaid inpatient-bed-days} + \text{estimated Medicaid HMO inpatient-bed-days}}{\text{est. Medicaid IP-bed-days} \times \left(\frac{\text{est. total charges} - \text{est. charity care charges}}{\text{est. total charges}}\right)}
\]

<table>
<thead>
<tr>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Medicaid Days</td>
<td>0</td>
</tr>
<tr>
<td>Total Medicaid HMO days</td>
<td>0</td>
</tr>
<tr>
<td>Total Medicaid and HMO Medicaid days</td>
<td>0</td>
</tr>
<tr>
<td>Total Hospital Charges</td>
<td>0</td>
</tr>
<tr>
<td>Uncompensated care charges (negative amount)</td>
<td>0</td>
</tr>
<tr>
<td>Total Hospital Charges - charity chgs</td>
<td>0</td>
</tr>
<tr>
<td>Divided by Total Hospital Charges</td>
<td>0</td>
</tr>
<tr>
<td>Non-charity percentage</td>
<td>0.00%</td>
</tr>
<tr>
<td>Total Hospital Days</td>
<td>0</td>
</tr>
<tr>
<td>Non-charity total Hospital Days</td>
<td>0</td>
</tr>
</tbody>
</table>

### Step 7: Computation of Medicaid aggregate EHR incentive amount

\[
\text{Medicaid Aggregate EHR Incentive Amount} = \frac{\text{Aggregate EHR amount for 4 years}}{\text{Non-charity percentage}}
\]

<table>
<thead>
<tr>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aggregate EHR amount for 4 years</td>
<td>$5,000,000</td>
</tr>
<tr>
<td>(Total Medicaid and HMO Medicaid days) divide non-charity hospital days</td>
<td>0.00%</td>
</tr>
<tr>
<td>Medicaid Aggregate EHR Incentive Amount</td>
<td>$0.00</td>
</tr>
</tbody>
</table>

### Step 8: Computation of Medicaid annual EHR incentive payout

<table>
<thead>
<tr>
<th>Year</th>
<th>Percentage</th>
<th>Payment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td>50.0%</td>
<td>$0</td>
</tr>
<tr>
<td>Year 2</td>
<td>40.0%</td>
<td>$0</td>
</tr>
<tr>
<td>Year 3</td>
<td>10.0%</td>
<td>$0</td>
</tr>
</tbody>
</table>

### CMS Reference - Authorized Data Sources for One Time Payment Calculation

If the State chooses to use the cost report in the Medicaid EHR incentive hospital payment calculation, what data elements should be used in the Medicare cost report, Form CMS 2552-96 and the Form CMS 2552-10?

Based on the Medicare cost report guidance, Form CMS 2552-96 will be used until the implementation of the new Medicare cost report, Form CMS 2552-10. Although the State may choose to use the following data elements, it is the States’ and hospitals’ responsibility to ensure the integrity and regulatory compliance of the data.
For information about the cost report data elements that are used in the Medicare hospital incentive calculation, please see FAQ #10717.

<table>
<thead>
<tr>
<th>The CMS 2552-96 data elements are as follows:</th>
</tr>
</thead>
<tbody>
<tr>
<td>-Total Discharges - Worksheet S-3 Part 1, Column 15, Line 12</td>
</tr>
<tr>
<td>-Medicaid Days - Worksheet S-3, Part I, Column 5, Line 1 + Lines 6-10</td>
</tr>
<tr>
<td>-Medicaid HMO Days - Worksheet S-3, Part I, Column 5, Line 2</td>
</tr>
<tr>
<td>-Total Inpatient Days - Worksheet S-3 Part 1, Column 6, Line 1, 2 + Lines 6 -10</td>
</tr>
<tr>
<td>-Total Hospital Charges - Worksheet C Part 1, Column 8, Line 101</td>
</tr>
<tr>
<td>-Charity Care Charges - Worksheet S-10, Column 1, Line 30</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>The CMS 2552-10 data elements are as follows:</th>
</tr>
</thead>
<tbody>
<tr>
<td>-Total Discharges - Worksheet S-3 Part 1, Column 15, Line 14</td>
</tr>
<tr>
<td>-Medicaid Days - Worksheet S-3, Part I, Column 7, Line 1 + Lines 8-12</td>
</tr>
<tr>
<td>-Medicaid HMO Days - Worksheet S-3, Part I, Column 7, Line 2</td>
</tr>
<tr>
<td>-Total Inpatient Days - Worksheet S-3 Part 1, Column 8, Line 1, 2 + Lines 8 -12</td>
</tr>
<tr>
<td>-Total Hospital Charges - Worksheet C Part 1, Column 8, Line 200</td>
</tr>
<tr>
<td>-Charity Care Charges - Worksheet S-10, Column 3, Line 20</td>
</tr>
</tbody>
</table>

Appendix G: Calculators
### Hospital One Time Payment Calculation

Calculation of Medicaid Electronic Health Records (EHR) Incentive Payment using 2552-10 Cost Report

This Payment Calculation was approved by CMS on 06/13/2011

<table>
<thead>
<tr>
<th>Hospital:</th>
<th>NPI:</th>
</tr>
</thead>
</table>

**White Areas require provider input**

**Grey Areas are calculated by the MS SLR application - Do not change**

The overall "EHR" amount is the sum over 4 years of (a) the base amount of $2,000,000 plus (b) the discharge related amount defined as $200 for the 1,150 through the 23,000 discharge for the first payment year then a pro-rated amount of 75% in yr 2, 50% in yr 3, and 25% in yr 4

For years 2-4 the rate of growth is assumed to be the previous 3 years' average.
**Step 1:**  **Compute the average annual growth rate over 3 years using previous Medicare cost reports.**

Per the Medicare cost report 2552-10, worksheet S-3, part I, line 14, column 15 - Total discharges

<table>
<thead>
<tr>
<th>Fiscal Yr</th>
<th>PY 2552-96</th>
<th>CY 2552-96</th>
<th>Increase</th>
<th>Growth</th>
</tr>
</thead>
<tbody>
<tr>
<td>2009</td>
<td></td>
<td></td>
<td>0</td>
<td>0.00%</td>
</tr>
<tr>
<td>2010</td>
<td></td>
<td></td>
<td>0</td>
<td>0.00%</td>
</tr>
<tr>
<td>2011</td>
<td></td>
<td></td>
<td>0</td>
<td>0.00%</td>
</tr>
<tr>
<td>2012</td>
<td></td>
<td></td>
<td>0</td>
<td>0.00%</td>
</tr>
</tbody>
</table>

Total Percent - Increase/(Decrease) 0.0%

Divided by 3 years 3

The average annual growth rate over 3 years 0.0%

**Step 2:**  **Compute total discharge related amount using proper transition factors**

> discharges are capped at 23,000 each year

<table>
<thead>
<tr>
<th>Discharges</th>
<th>Total</th>
<th>Allowable</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td>0</td>
<td>0</td>
<td>$0</td>
</tr>
<tr>
<td>Year 2</td>
<td>0</td>
<td>0</td>
<td>$0</td>
</tr>
<tr>
<td>Year 3</td>
<td>0</td>
<td>0</td>
<td>$0</td>
</tr>
<tr>
<td>Year 4</td>
<td>0</td>
<td>0</td>
<td>$0</td>
</tr>
</tbody>
</table>

Total 4 year discharge related amount $0

**Step 3:**  **Compute the initial amount for 4 years**

<table>
<thead>
<tr>
<th>Years 1 - 4 base amount of $2,000,000 per year</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td>$2,000,000</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Years 1-4 discharge related amount (step 2)</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
</tr>
</tbody>
</table>

Aggregate EHR amount for 4 years $2,000,000

**Step 4:**  **Apply Transition Factor**

| $2,000,000 | $1,500,000 | $1,000,000 | $500,000 |

**Step 5:**  **Compute the overall EHR amount for 4 years**

| $5,000,000 |

*Appendix G: Calculators*
### Step 6: Computation of Medicaid Share from the Medicare cost report (Revised 2552-10 Cost Report)

\[
\text{(estimated Medicaid inpatient-bed-days + estimated Medicaid HMO inpatient-bed-days) / (est. Medicaid IP-bed-days x ((est. total charges - est. charity care charges) / est. total charges))}
\]

| w/s S-3 part I, col. 7, lines 1,8,9,10,11,12 | Total Medicaid Days | 0 |
| w/s S-3 part I, col. 7, line 2 | Total Medicaid HMO days | 0 |
| | Total Medicaid and HMO Medicaid days | 0 |
| w/s C part I, col. 8, line 200 | Total Hospital Charges | $0 |
| w/s S-10, line 20 | Uncompensated care charges (negative amount) | $0 |
| | Total Hospital Charges - charity chgs | $0 |
| | divided by Total Hospital Charges | $0 |
| | Non-charity percentage | 0.00% |
| w/s S-3 part I, col. 8, lines 1,2,8,9,10,11,12 | Total Hospital Days | 0 |
| | Non-charity total Hospital Days | 0 |

\[
\text{(Total Medicaid and HMO Medicaid days) divide non-charity hospital days} \quad 0.00%
\]

### Step 7: Computation of Medicaid aggregate EHR incentive amount

Aggregate EHR amount for 4 years $5,000,000

\[
\frac{(\text{Total Medicaid and HMO Medicaid days}) \text{ divide non-charity hospital days}}{0.00%}
\]

**Medicaid Aggregate EHR Incentive Amount** $0.00

### Step 8: Computation of Medicaid annual EHR incentive payout

<table>
<thead>
<tr>
<th>Annual Percentage</th>
<th>Payment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1 payment</td>
<td>50.0%</td>
</tr>
<tr>
<td>Year 2 payment</td>
<td>40.0%</td>
</tr>
<tr>
<td>Year 3 payment</td>
<td>10.0%</td>
</tr>
</tbody>
</table>

---

**CMS Reference - Authorized Data Sources for One Time Payment Calculation**

Published 08/09/2011 09:32 AM | Updated 12/05/2011 01:45 PM | Answer ID 10771

If the State chooses to use the cost report in the Medicaid EHR incentive hospital payment calculation, what data elements should be used in the Medicare cost report, Form CMS 2552-96 and the Form CMS 2552-10?

Based on the Medicare cost report guidance, Form CMS 2552-96 will be used until the implementation of the new Medicare cost report, Form CMS 2552-10. Although the State may choose to use the following data elements, it is the States’ and hospitals’ responsibility to ensure the integrity and regulatory compliance of the data.
## Appendix G: Calculators

### The CMS 2552-96 data elements are as follows:

- **Total Discharges** - Worksheet S-3 Part 1, Column 15, Line 12
- **Medicaid Days** - Worksheet S-3, Part I, Column 5, Line 1 + Lines 6-10
- **Medicaid HMO Days** - Worksheet S-3, Part I, Column 5, Line 2
- **Total Inpatient Days** - Worksheet S-3 Part 1, Column 6, Line 1, 2 + Lines 6-10
- **Total Hospital Charges** - Worksheet C Part 1, Column 8, Line 101
- **Charity Care Charges** - Worksheet S-10, Column 1, Line 30

### The CMS 2552-10 data elements are as follows:

- **Total Discharges** - Worksheet S-3 Part 1, Column 15, Line 14
- **Medicaid Days** - Worksheet S-3, Part I, Column 7, Line 1 + Lines 8-12
- **Medicaid HMO Days** - Worksheet S-3, Part I, Column 7, Line 2
- **Total Inpatient Days** - Worksheet S-3 Part 1, Column 8, Line 1, 2 + Lines 8-12
- **Total Hospital Charges** - Worksheet C Part 1, Column 8, Line 200
- **Charity Care Charges** - Worksheet S-10, Column 3, Line 20

For information about the cost report data elements that are used in the Medicare hospital incentive calculation, please see FAQ #10717.
Appendix H: Impact of Incentive Payments

Based on the results of the survey, at least 90% of the Providers who planned to attest to A/I/U indicated that incentive payments were a major factor in their decision. These results were consistent regardless of location or Provider type.

<table>
<thead>
<tr>
<th>Provider Type</th>
<th>High</th>
<th>Medium</th>
<th>Low</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dentist</td>
<td>4</td>
<td></td>
<td></td>
<td>4</td>
</tr>
<tr>
<td>FQHC</td>
<td>1</td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Hospital</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Optometry</td>
<td>8</td>
<td>1</td>
<td>9</td>
<td>18</td>
</tr>
<tr>
<td>Pediatrics</td>
<td>4</td>
<td>4</td>
<td></td>
<td>8</td>
</tr>
<tr>
<td>Physician</td>
<td>24</td>
<td>1</td>
<td>1</td>
<td>26</td>
</tr>
<tr>
<td>Grand Total</td>
<td>42</td>
<td>2</td>
<td>2</td>
<td>46</td>
</tr>
</tbody>
</table>

Percentages
- Overall Percentage: 91% 4% 4% 100%
- Non Physician Percentage: 90% 5% 5% 100%
- Physician Percentage: 92.3% 3.8% 3.8% 100%

<table>
<thead>
<tr>
<th>Location</th>
<th>High</th>
<th>Medium</th>
<th>Low</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coast Metro</td>
<td>5</td>
<td></td>
<td></td>
<td>5</td>
</tr>
<tr>
<td>Columbus Metro</td>
<td>2</td>
<td></td>
<td></td>
<td>2</td>
</tr>
<tr>
<td>JXN Metro</td>
<td>10</td>
<td>2</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>McComb</td>
<td>1</td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Memphis Metro</td>
<td>5</td>
<td></td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Meridian Metro</td>
<td>5</td>
<td></td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Picayune</td>
<td>1</td>
<td></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Tupelo Metro</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Under 50,000</td>
<td>11</td>
<td>1</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Grand Total</td>
<td>42</td>
<td>2</td>
<td>2</td>
<td>46</td>
</tr>
</tbody>
</table>

Percentages
- Overall Percentage: 91% 3% 6% 100%
- Metro Area Percentage: 91% 3% 6% 100%
- Rural Area Percentage: 91.7% 8.3% 0.0% 100%
Appendix I: MU Requirements (Updated 2017)

The Medicare and Medicaid EHR Incentive Programs provide financial incentives for the meaningful use of certified EHR technology to improve patient care. To receive an EHR incentive payment, providers have to show that they are meaningfully using their EHRs by meeting thresholds for a number of objectives. The EHR Incentive Programs are phased in three stages with increasing requirement complexity.

Eligible professionals participate in the program on the calendar year, while eligible hospitals and CAHs participate according to the federal fiscal year. Providers must attest to demonstrating meaningful use every year to receive an incentive and avoid a Medicare payment adjustment.

Requirements for 2014 Definition Stage 1

In May 2014, CMS released an NPRM that would grant flexibility to providers who are experiencing difficulties fully implementing 2014 Edition certified EHR technology (CEHRT) to attest this year. Providers scheduled to demonstrate Stage 1 in 2014 who have successfully implemented 2014 CEHRT would use 2014 Definition Stage 1 core and menu objectives. Providers who are still using 2011 Edition CEHRT or a combination of 2011 and 2014 Editions and choose to report 2013 Definition Stage 1 core and menu objectives should visit the 2013 Definition Stage 1 of Meaningful Use webpage.

Criteria for providers demonstrating the 2014 Definition of Stage 1 is listed below.

Eligible professionals must meet:
- 13 required core objectives
- 5 menu objectives from a list of 9
- Total of 18 objectives

Eligible hospitals and CAHs must meet:
- 11 required core objectives
- 5 menu objectives from a list of 10
- Total of 16 objectives
Requirements for Stage 2 of MU
The CMS Stage 2 Final Rule from 2012 specifies the criteria that eligible professionals, eligible hospitals, and critical access hospitals (CAHs) must meet in order to participate in Stage 2 of the Medicare and Medicaid EHR Incentive Programs. All providers must demonstrate Stage 1 of meaningful use before Stage 2.

To help providers better understand Stage 2 meaningful use requirements, CMS developed specification sheets for eligible professionals and eligible hospitals that provide detailed information on each objective, including:

- Numerator and denominator thresholds
- Exclusion criteria
- Definitions of important terms
- Requirements for achieving the objectives
- Certification information that corresponds with each objective

Stage 2 Timeline

The earliest providers will demonstrate Stage 2 of meaningful use is 2014. Eligible hospitals and CAHs participate on the fiscal year and eligible professionals participate on the calendar year.

Providers who began participation in the EHR Incentive Programs in 2011 will meet three consecutive years of meaningful use under the Stage 1 criteria before advancing to the Stage 2 criteria in 2014. All other providers would meet two years of meaningful use under the Stage 1 criteria before advancing to the Stage 2 criteria in their third year.

For 2014 Only

2014 CEHRT Flexibility
In May 2014, CMS released an NPRM that would grant flexibility to providers who are experiencing difficulties fully implementing 2014 Edition CEHRT to attest this year.

Providers scheduled to demonstrate Stage 2 of meaningful use in 2014 can:

- Demonstrate 2013 Definition of Stage 1 of meaningful use with 2011 Edition CEHRT or a combination of 2011 and 2014 Edition CEHRT
- Demonstrate 2014 Definition of Stage 1 of meaningful use with 2014 Edition CEHRT
- Demonstrate Stage 2 of meaningful use with 2014 Edition CEHRT

2014 Reporting Periods

All providers, regardless of their stage, are only required to demonstrate meaningful use for a 3-month EHR reporting period. For Medicare providers, this 3-month reporting period is fixed to the quarter of either the fiscal (for eligible hospitals and CAHs) or calendar (for eligible professionals). The 3-month reporting period is not fixed for Medicaid eligible professionals and hospitals that are only eligible to receive Medicaid EHR incentives.
Stage 2 Core and Menu Objectives

Stage 2 uses a core and menu structure for objectives that providers must achieve in order to demonstrate meaningful use. Core objectives are objectives that all providers must meet. There are also a predetermined number of menu objectives that providers must select from a list and meet in order to demonstrate meaningful use.

To demonstrate meaningful use under Stage 2 criteria—

Eligible professionals must meet:

- 17 core objectives
- 3 menu objectives that they select from a total list of 6
- Total of 20 objectives

Eligible hospitals and CAHs must meet:

- 16 core objectives
- 3 menu objectives that they select from a total list of 6
- Total of 19 objectives
Definition of Modified Stage 2

Eligible Professionals (EPs) Requirements

CMS published a final rule on October 16, 2015 that specifies criteria that eligible professionals (EPs), eligible hospitals, and critical access hospitals (CAHs) must meet in order to participate in the Medicare and Medicaid Electronic Health Record (EHR) Incentive Programs. The final rule’s provisions encompass the definition of meaningful use for 2015 through 2017.

Here’s what you need to know about meeting the requirements of the EHR Incentive Programs in 2016.

Objectives and Measures

All providers are required to attest to a single set of objectives and measures. This replaces the core and menu objectives structure of previous stages.

For EPs, there are 10 objectives.

In 2016, all providers must attest to objectives and measures using EHR technology certified to the 2014 Edition or the 2015 Edition, or a combination of the two.

Alternate Exclusions and Specifications

Many of the alternate exclusions that were available in 2015 are not applicable in 2016.

The Definition of Modified Stage 2 Meaningful Use Objectives for Eligible Professionals EPs

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Objective 1: Protect Patient Health Information</strong></td>
<td><strong>Measure</strong>: Conduct or review a security risk analysis in accordance with the requirements in 45 CFR 164.308(a)(1), including addressing the security (to include encryption) of ePHI created or maintained in CEHRT in accordance with requirements under 45 CFR 164.312(a)(2)(iv) and 45 CFR 164.306(d)(3), and implement security updates as necessary and correct identified security deficiencies as part of the EP’s risk management process.</td>
</tr>
<tr>
<td><strong>Objective 2: Clinical Decision Support</strong></td>
<td><strong>Measure 1</strong>: Implement five clinical decision support interventions related to four or more clinical quality measures at a relevant point in patient care for the entire EHR reporting period. Absent four clinical quality measures related to an EP’s scope of practice or patient population, the clinical decision support</td>
</tr>
</tbody>
</table>

**Appendix I: MU Requirements**

Page 175
interventions must be related to high priority health conditions.

**Measure 2:** The EP has enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period.

**Exclusion:** For the second measure, any EP who writes fewer than 100 medication orders during the EHR reporting period.

<table>
<thead>
<tr>
<th>Objective 3: Computerized Provider Order Entry</th>
</tr>
</thead>
<tbody>
<tr>
<td>An EP, through a combination of meeting the thresholds and exclusions (or both), must satisfy all three measures for this objective.</td>
</tr>
</tbody>
</table>

**Measure 1:** More than 60 percent of medication orders created by the EP during the EHR reporting period are recorded using computerized provider order entry.

**Exclusion for Measure 1:** Any EP who writes fewer than 100 medication orders during the EHR reporting period.

**Measure 2:** More than 30 percent of laboratory orders created by the EP during the EHR reporting period are recorded using computerized provider order entry.

**Exclusion for Measure 2:** Any EP who writes fewer than 100 laboratory orders during the EHR reporting period.

**Alternate Exclusion for Measure 2:** Providers scheduled to be in Stage 1 in 2016 may claim an exclusion for measure 2 (laboratory orders) of the Stage 2 CPOE objective for an EHR reporting period in 2016.

**Measure 3:** More than 30 percent of radiology orders created by the EP during the EHR reporting period are recorded using computerized provider order entry.

**Exclusion for Measure 3:** Any EP who writes fewer than 100 radiology orders during the EHR reporting period.

**Alternate Exclusion for Measure 3:** Providers scheduled to be in Stage 1 in 2016 may claim an exclusion for measure 3 (radiology orders) of the Stage 2 CPOE objective for an EHR reporting period in 2016.

<table>
<thead>
<tr>
<th>Objective 4: Electronic Prescribing</th>
</tr>
</thead>
<tbody>
<tr>
<td>EP Measure: More than 50 percent of permissible prescriptions written by the EP are queried for a drug formulary and transmitted electronically using CEHRT.</td>
</tr>
</tbody>
</table>

**Exclusions:** Any EP who (1) Writes fewer than 100 permissible prescriptions during the EHR reporting period; or (2) Does not have a pharmacy within his or her organization and there are no pharmacies
that accept electronic prescriptions within 10 miles of the EP's practice location at the start of his or her EHR reporting period.

| **Objective 5: Health Information Exchange** | **Measure:** The EP that transitions or refers their patient to another setting of care or provider of care must (1) use CEHRT to create a summary of care record; and (2) electronically transmit such summary to a receiving provider for more than 10 percent of transitions of care and referrals.  

**Exclusion:** Any EP who transfers a patient to another setting or refers a patient to another provider less than 100 times during the EHR reporting period. |

| **Objective 6: Patient Specific Education** | **EP Measure:** Patient specific education resources identified by CEHRT are provided to patients for more than 10 percent of all unique patients with office visits seen by the EP during the EHR reporting period.  

**Exclusion:** Any EP who has no office visits during the EHR reporting period. |

| **Objective 7: Medication Reconciliation** | **Measure:** The EP performs medication reconciliation for more than 50 percent of transitions of care in which the patient is transitioned into the care of the EP.  

**Exclusion:** Any EP who was not the recipient of any transitions of care during the EHR reporting period. |

| **Objective 8: Patient Electronic Access (VDT)** | **EP Measure 1:** More than 50 percent of all unique patients seen by the EP during the EHR reporting period are provided timely access to view online, download, and transmit to a third party their health information subject to the EP's discretion to withhold certain information.  

**Exclusion for Measure 1:** Any EP who neither orders nor creates any of the information listed for inclusion as part of the measures except for “Patient Name” and “Provider’s name and office contact information.”

**EP Measure 2:** For an EHR reporting period in 2016, at least one patient seen by the EP during the EHR reporting period (or patient-authorized representative) views, downloads or transmits his or her health information to a third party during the EHR reporting period.  

**Exclusion for Measure 2:** Any EP who (1) Neither orders nor creates any of the information listed for inclusion as part of the measures except for “Patient Name” and “Provider’s name and office contact information”; or (2) Conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period.
### Objective 9: Secure Messaging

**Measure:** For an EHR reporting period in 2016, for at least 1 patient seen by the EP during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient-authorized representative), or in response to a secure message sent by the patient (or the patient-authorized representative) during the EHR reporting period.

**Exclusion:** Any EP who has no office visits during the EHR reporting period, or any EP who conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period.

### Objective 10: Public Health Reporting

EPs in 2016 must meet 2 of the 3 measures.

**Measure Option 1 – Immunization Registry Reporting:** The EP is in active engagement with a public health agency to submit immunization data.

**Exclusions for Measure 1:** Any EP meeting one or more of the following criteria may be excluded from the immunization registry reporting measure if the EP:

- Does not administer any immunizations to any of the populations for which data is collected by its jurisdiction’s immunization registry or immunization information system during the EHR reporting period;

- Operates in a jurisdiction for which no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or

- Operates in a jurisdiction where no immunization registry or immunization information system has declared readiness to receive immunization data from the EP at the start of the EHR reporting period.

**Measure Option 2 – Syndromic Surveillance Reporting:** The EP is in active engagement with a public health agency to submit syndromic surveillance data.

**Exclusions for Measure 2:** Any EP meeting one or more of the following criteria may be excluded from the syndromic surveillance reporting measure if the EP:

- Is not in a category of providers from which ambulatory syndromic surveillance data is collected by their jurisdiction's syndromic...
surveillance system;

Operates in a jurisdiction for which no public health agency is capable of receiving electronic syndromic surveillance data from EPs in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or

Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from EPs at the start of the EHR reporting period.

**Measure Option 3 – Specialized Registry Reporting:** The EP is in active engagement to submit data to a specialized registry.

**Exclusions for Measure 3:** Any EP meeting at least one of the following criteria may be excluded from the specialized registry reporting measure if the EP:

Does not diagnose or treat any disease or condition associated with, or collect relevant data that is collected by, a specialized registry in their jurisdiction during the EHR reporting period;

Operates in a jurisdiction for which no specialized registry is capable of accepting electronic registry transactions in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or

Operates in a jurisdiction where no specialized registry for which the EP is eligible has declared readiness to receive electronic registry transactions at the beginning of the EHR reporting period.

**Alternate Exclusions for 2016:**

EPs scheduled to be in Stage 1 and Stage 2 in 2016: Must attest to at least 2 measures from the Public Health Reporting Objective Measures 1-3.

May claim an Alternate Exclusion for Measure 2 and Measure 3 (Syndromic Surveillance and Specialized Registry Reporting).

An Alternate Exclusion may only be claimed for up to two measures, then the provider must either attest to or meet the exclusion requirements for the remaining measure described in 495.22 (e)(10)(i)(C).
Eligible Hospitals and CAHs Requirements

CMS published a final rule on October 16, 2015 that specifies criteria that eligible professionals (EPs), eligible hospitals, and critical access hospitals (CAHs) must meet in order to participate in the Medicare and Medicaid Electronic Health Record (EHR) Incentive Programs. The final rule’s provisions encompass the definition of meaningful use for 2015 through 2017.

Here’s what you need to know about meeting the requirements of the EHR Incentive Programs in 2016.

Objectives and Measures

All providers are required to attest to a single set of objectives and measures. This replaces the core and menu objectives structure of previous stages.

There are 9 objectives for eligible hospitals and CAHs.

In 2016, all providers must attest to objectives and measures using EHR technology certified to the 2014 Edition or the 2015 Edition, or a combination of the two.

Alternate Exclusions and Specifications

Many of the alternate exclusions that were available in 2015 are not available in 2016.

The Definition of Modified Stage 2 Meaningful Use Objectives for Eligible Hospitals and CAHs

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td><strong>Objective 1: Protect Patient Health Information</strong></td>
<td><strong>Measure:</strong> Conduct or review a security risk analysis in accordance with the requirements in 45 CFR 164.308(a)(1), including addressing the security (to include encryption) of ePHI created or maintained in CEHRT in accordance with requirements under 45 CFR 164.312(a)(2)(iv) and 45 CFR 164.306(d)(3), and implement security updates as necessary and correct identified security deficiencies as part of the eligible hospital or CAH's risk management process.</td>
</tr>
<tr>
<td><strong>Objective 2: Clinical Decision Support</strong></td>
<td><strong>Measure 1:</strong> Implement five clinical decision support interventions related to four or more clinical quality measures at a relevant point in patient care for the entire EHR reporting period. Absent four clinical quality measures related to an eligible hospital or CAH's scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions.</td>
</tr>
<tr>
<td>Measure 2:</td>
<td>The eligible hospital or CAH has enabled and implemented the functionality for drug-drug and drug allergy interaction checks for the entire EHR reporting period.</td>
</tr>
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<td>-----------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| Objective 3: Computerized Provider Order Entry | *Eligible hospitals and CAHs must meet the thresholds of all three measures.*  
**Measure 1:** More than 60 percent of medication orders created by authorized providers of the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.  
**Measure 2:** More than 30 percent of laboratory orders created by authorized providers of the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.  
**Alternate Exclusion for Measure 2:** Providers scheduled to be in Stage 1 in 2016 may claim an exclusion for measure 2 (laboratory orders) of the Stage 2 CPOE objective for an EHR reporting period in 2016.  
**Measure 3:** More than 30 percent of radiology orders created by authorized providers of the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.  
**Alternative Exclusion for Measure 3:** Providers scheduled to be in Stage 1 in 2016 may claim an exclusion for measure 3 (radiology orders) of the Stage 2 CPOE objective for an EHR reporting period in 2016. |
| Objective 4: Electronic Prescribing | *Eligible Hospital/CAH Measure:* More than 10 percent of hospital discharge medication orders for permissible prescriptions (for new and changed prescriptions) are queried for a drug formulary and transmitted electronically using CEHRT.  
**Exclusion:** Any eligible hospital or CAH that does not have an internal pharmacy that can accept electronic prescriptions and is not located within 10 miles of any pharmacy that accepts electronic prescriptions at the start of their EHR reporting period.  
**Alternate Exclusion:** An eligible hospital or CAH may claim an exclusion for the eRx objective and measure for an EHR reporting period in 2016 if they were either scheduled to demonstrate Stage 1 in 2016, or if they are scheduled to demonstrate Stage 2 but did not intend to select the Stage 2 eRx objective for an EHR reporting period in 2016. |
<table>
<thead>
<tr>
<th>Objective</th>
<th>Measure/Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Objective 5: Health Information Exchange</strong></td>
<td>The eligible hospital or CAH that transitions or refers their patient to another information setting of care or provider of care must (1) use CEHRT to create a summary of care record; Exchange and (2) electronically transmit such summary to a receiving provider for more than 10 percent of transitions of care and referrals.</td>
</tr>
<tr>
<td><strong>Objective 6: Patient Specific Education</strong></td>
<td><strong>Eligible Hospital/CAH Measure:</strong> More than 10 percent of all unique patients admitted to the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) are provided patient specific education resources identified by CEHRT.</td>
</tr>
<tr>
<td><strong>Objective 7: Medication Reconciliation</strong></td>
<td><strong>Measure:</strong> The eligible hospital or CAH performs medication reconciliation for more than 50 percent of transitions of care in which the patient is admitted to the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23).</td>
</tr>
</tbody>
</table>
| **Objective 8: Patient Electronic Access (VDT)** | **Measure 1:** More than 50 percent of all unique patients who are discharged from the inpatient or emergency department (POS 21 or 23) of an eligible hospital or CAH are provided timely access to view online, download and transmit to a third party their health information.  
**Measure 2:** For an EHR reporting period in 2016, at least 1 patient who is discharged from the inpatient or emergency department (POS 21 or 23) of an eligible hospital or CAH (or patient-authorized representative) views, downloads or transmits his or her health information to a third party during the EHR reporting period.  
**Exclusion for Measure 2:** Any eligible hospital or CAH that is located in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period. |
| **Objective 9: Public Health Reporting** | In 2016, all eligible hospitals and CAHs must meet three measures.  
**Measure Option 1 – Immunization Registry Reporting:** The eligible hospital or CAH is in active engagement with a public health agency to submit immunization data.  
**Exclusions for Measure 1:** Any eligible hospital or CAH meeting one or more of the following criteria may be excluded from the immunization registry reporting measure if the eligible hospital or CAH:  
- Does not administer any immunizations to any of the populations for which data is collected by its jurisdiction’s immunization registry or immunization information system |
during the EHR reporting period;

- Operates in a jurisdiction for which no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no immunization registry or immunization information system has declared readiness to receive immunization data from the eligible hospital or CAHs at the start of the EHR reporting period.

**Measure Option 2 – Syndromic Surveillance Reporting:** The eligible hospital or CAH is in active engagement with a public health agency to submit syndromic surveillance data.

**Exclusions for Measure 2:** Any eligible hospital or CAH meeting one or more of the following criteria may be excluded from the syndromic surveillance reporting measure if the eligible hospital or CAH:

- Does not have an emergency or urgent care department;
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic syndromic surveillance data from eligible hospitals or CAHs in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from eligible hospitals or CAHs at the start of the EHR reporting period.

**Measure Option 3 – Specialized Registry Reporting:** The eligible hospital or CAH is in active engagement to submit data to a specialized registry.

**Exclusions for Measure 3:** Any eligible hospital or CAH meeting at least one of the following criteria may be excluded from the specialized registry reporting measure if the EP, eligible hospital, or CAH:

- Does not diagnose or treat any disease or condition associated with, or collect relevant data that is collected by, a specialized registry in their jurisdiction during the EHR reporting period;
- Operates in a jurisdiction for which no specialized registry is capable of accepting electronic registry transactions in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no specialized registry for which the eligible hospital or CAH is eligible has declared readiness to receive electronic registry transactions at the beginning of the EHR reporting period.
Measure Option 4—Electronic Reportable Laboratory Result Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit electronic reportable laboratory (ELR) results.

Exclusions for Measure 4: Any eligible hospital or CAH meeting one or more of the following criteria may be excluded from the electronic reportable laboratory result reporting measure if the eligible hospital or CAH:

- Does not perform or order laboratory tests that are reportable in their jurisdiction during the EHR reporting period;
- Operates in a jurisdiction for which no public health agency is capable of accepting the specific ELR standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no public health agency has declared readiness to receive electronic reportable laboratory results from eligible hospitals or CAHs at the start of the EHR reporting period.

Alternate Exclusion for 2016

Eligible hospitals/CAHs scheduled to be in Stage 1 and Stage 2 in 2016: Must attest to at least 3 measures from the Public Health Reporting Objective Measures 1-4.

- May claim an Alternate Exclusion for Measure 3 (Specialized Registry Reporting).
- If an Alternate Exclusion is claimed, then the provider must either attest to or meet the exclusion requirements for the remaining measures described in 495.22 (e)(10)(ii)(C).

Immunization Registry will periodically send DOM a list of providers that are no longer compliant, meaning no response when invited to participate in testing interfaces.
**Definition of Stage 3 Meaningful Use:**

**Stage 3 Program Requirements for Providers Attesting to their State’s Medicaid EHR Incentive Program**

In October 2015, CMS released a final rule that modified the requirements for participation in the Electronic Health Record (EHR) Incentive Programs for years 2015 through 2017 as well as in 2018 and beyond. This page provides information on requirements for Stage 3.

In 2018, all providers will be required to participate in Stage 3 regardless of their prior participation. Moving all participants to a single stage of meaningful use aims to reduce the program’s complexity and simplify reporting requirements.

Medicaid providers who are only eligible to participate in the Medicaid EHR Incentive Program are not subject to the Medicare payment adjustments.

Mississippi will continue manual attestation or reporting of Clinical Quality Measures (CQMs)

**NOTE:** All providers who have not successfully demonstrated meaningful use in a prior year and are seeking to demonstrate meaningful use for the first time in 2017 to avoid the 2018 payment adjustment must attest to Modified Stage 2 objectives and measures.

**Objectives and Measures**

- All providers are required to attest to a single set of objectives and measures.
- For eligible professionals (EPs) and eligible hospitals there are 8 objectives.
- To meet Stage 3 requirements, all providers must use technology certified to the 2015 Edition. A provider who has technology certified to a combination of the 2015 Edition and 2014 Edition may potentially attest to the Stage 3 requirements, if the mix of certified technologies would not prohibit them from meeting the Stage 3 measures. However, a provider who has technology certified to the 2014 Edition only may not attest to Stage 3.
- Please note there are no alternate exclusions or specifications available.
- There are changes to the measure calculations policy, which specifies that actions included the numerator must occur within the EHR reporting period if that period is a full calendar year, or if it is less than a full calendar year, within the calendar year in which the EHR reporting period occurs. Specific measures affected are identified in the Additional Information section of the specification sheets.

**Flexibility within Objectives and Measures**

- Stage 3 includes flexibility within certain objectives to allow providers to choose the measures most relevant to their patient population or practice. The Stage 3 objectives with flexible measure options include:
  - Coordination of Care through Patient Engagement – Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective.
  - Health Information Exchange – Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective.
  - Public Health Reporting – Eligible professionals must report on two measures and eligible hospitals must report on four measures.

**EHR Reporting Period**
Starting in 2018, all providers are required to use an EHR reporting period of a full calendar year, with the exception of providers attesting to meaningful use for the first time; these providers will have a minimum of any continuous 90-days EHR reporting period.

NOTE: In 2017, for all new and returning participants, the EHR reporting period is a minimum of any continuous 90 days between January 1 and December 31, 2017.

Here is what Eligible Professionals (EPs) should know about Stage 3 Meaningful Use:

<table>
<thead>
<tr>
<th>Stage Three Meaningful Use Objectives (beginning January 1, 2018)</th>
<th>Stage 3 Meaningful Use Measures for EPs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protect electronic protected health information (ePHI)</td>
<td>Protect electronic protected health information (ePHI) created or maintained by the CEHRT through the implementation of appropriate technical, administrative, and physical safeguards.</td>
</tr>
</tbody>
</table>
| Generate and transmit permissible prescriptions electronically (eRx) | Measure: More than 60 percent of all permissible prescriptions written by the EP are queried for a drug formulary and transmitted electronically using CEHRT. Exclusions: Any EP who: 
  - Writes fewer than 100 permissible prescriptions during the EHR reporting period; or
  - Does not have a pharmacy within their organization and there are no pharmacies that accept electronic prescriptions within 10 miles of the EP’s practice location at the start of his or her EHR reporting period. |
<p>| Clinical Decision Support                                      | Measure 1: Implement five clinical decision support interventions related to four or more CQMs at a relevant point in patient care for the entire EHR reporting period. Absent four CQMs related to an EP’s scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions. Measure 2: The EP has enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period. Exclusion: Any EP who writes fewer than 100 medication orders during the EHR reporting period. |</p>
<table>
<thead>
<tr>
<th>Computerized Provider Order Entry (CPOE)</th>
<th>An EP, through a combination of meeting the thresholds and exclusions (or both), must satisfy all three measures for this objective:</th>
</tr>
</thead>
</table>
|                                         | **Measure 1:**  
|                                         | More than 60 percent of medication orders created by the EP during the EHR reporting period are recorded using computerized provider order entry. |
|                                         | **Measure 2:**  
|                                         | More than 60 percent of laboratory orders created by the EP during the EHR reporting period are recorded using computerized provider order entry. |
|                                         | **Measure 3:**  
|                                         | More than 60 percent of diagnostic imaging orders created by the EP during the EHR reporting period are recorded using computerized provider order entry. |
| Exclusions:                             | **Measure 1:**  
|                                         | Any EP who writes fewer than 100 medication orders during the EHR reporting period. |
|                                         | **Measure 2:**  
|                                         | Any EP who writes fewer than 100 laboratory orders during the EHR reporting period. |
|                                         | **Measure 3:**  
|                                         | Any EP who writes fewer than 100 diagnostic imaging orders during the EHR reporting period. |

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<thead>
<tr>
<th>Patient Electronic Access</th>
<th>EPs must satisfy both measures in order to meet this objective:</th>
</tr>
</thead>
</table>
|                           | **Measure 1:**  
|                           | For more than 80 percent of all unique patients seen by the EP:  
|                           | 1) The patient (or the patient-authorized representative) is provided timely access to view online, download, and transmit his or her health information; and  
|                           | 2) The provider ensures the patient’s health information is available for the patient (or patient-authorized representative) to access using any application of their choice that is configured to meet the technical specifications of the Application Programming Interface (API) in the provider’s CEHRT. |
|                           | **Measure 2:**  
|                           | |
The EP must use clinically relevant information from CEHRT to identify patient-specific educational resources and provide electronic access to those materials to more than 35 percent of unique patients seen by the EP during the EHR reporting period.

Exclusions:

**Measure 1 and Measure 2:**
A provider may exclude the measures if one of the following applies:
- An EP may exclude from the measure if they have no office visits during the EHR reporting period.
- Any EP that conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period may exclude the measure.

<table>
<thead>
<tr>
<th>Coordination of Care</th>
<th>Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective:</th>
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<tbody>
<tr>
<td><strong>Measure 1:</strong></td>
<td>For an EHR reporting period in 2017, more than 5 percent of all unique patients (or their authorized representatives) seen by the EP actively engage with the electronic health record made accessible by the provider and either—</td>
</tr>
<tr>
<td></td>
<td>1. View, download or transmit to a third party their health information; or</td>
</tr>
<tr>
<td></td>
<td>2. Access their health information through the use of an API that can be used by applications chosen by the patient and configured to the API in the provider's CEHRT; or</td>
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<tr>
<td></td>
<td>3. A combination of (1) and (2)</td>
</tr>
<tr>
<td><strong>Threshold for 2018 and Subsequent Years:</strong></td>
<td>The resulting percentage must be more than 10 percent.</td>
</tr>
</tbody>
</table>

**Measure 2:**
For an EHR reporting period in 2017, more than 5 percent of all unique patients seen by the EP during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient authorized representative), or in response to a secure message sent by the patient or their authorized representative.

**Threshold in 2018 and Subsequent Years:** The resulting percentage must be more than 25 percent in order for an EP to meet this measure.
<table>
<thead>
<tr>
<th>Measure 3:</th>
<th>Patient generated health data or data from a nonclinical setting is incorporated into the CEHRT for more than 5 percent of all unique patients seen by the EP during the EHR reporting period.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exclusions:</td>
<td>Measure 1, 2 and 3 Exclusion: A provider may exclude the measures if one of the following apply: An EP may exclude from the measure if they have no office visits during the EHR reporting period, or; Any EP that conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period may exclude the measure.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Health Information Exchange</th>
<th>Providers must attest to all three measures and must meet the threshold for at least two measures to meet the objective.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure 1:</td>
<td>For more than 50 percent of transitions of care and referrals, the EP that transitions or refers their patient to another setting of care or provider of care: 1) Creates a summary of care record using CEHRT; and 2) Electronically exchanges the summary of care record</td>
</tr>
<tr>
<td>Measure 2:</td>
<td>For more than 40 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the EP incorporates into the patient’s EHR an electronic summary of care document.</td>
</tr>
<tr>
<td>Measure 3:</td>
<td>For more than 80 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the EP performs a clinical information reconciliation. The provider must implement clinical information reconciliation for the following three clinical information sets: 1) Medication. Review of the patient’s medication, including the name, dosage, frequency, and route of each medication. 2) Medication allergy. Review of the patient’s known medication allergies. 3) Current Problem list. Review of the patient’s current and active...</td>
</tr>
</tbody>
</table>
diagnoses.

Exclusions:

**Measure 1:**
A provider may exclude from the measure if any of the following apply:
- Any EP who transfers a patient to another setting or refers a patient to another provider less than 100 times during the EHR reporting period.
- Any EP that conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period may exclude the measures.

**Measure 2:**
A provider may exclude from the measure if any of the following apply:
- Any EP for whom the total of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, is fewer than 100 during the EHR reporting period is excluded from this measure.
- Any EP that conducts 50 percent or more of his or her patient encounters in a county that does not have 50 percent or more of its housing units with 4Mbps

<table>
<thead>
<tr>
<th>Public Health Reporting</th>
<th>Measure 1: Immunization Registry Reporting: The EP is in active engagement with a public health agency to submit immunization data and receive immunization forecasts and histories from the public health immunization registry/immunization information system (IIS).</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Measure 2: Syndromic Surveillance Reporting: The EP is in active engagement with a public health agency to submit syndromic surveillance data from an urgent care setting.</td>
</tr>
<tr>
<td></td>
<td>Measure 3: Electronic Case Reporting: The EP is in active engagement with a public health agency to submit case reporting of reportable conditions.</td>
</tr>
<tr>
<td></td>
<td>Measure 4: Public Health Registry Reporting: The EP is in active engagement with a</td>
</tr>
</tbody>
</table>
public health agency to submit data to public health registries.

**Measure 5:**
Clinical Data Registry Reporting: The EP is in active engagement to submit data to a clinical data registry.

**Exclusions:**

**Measure 1:**
Any EP meeting one or more of the following criteria may be excluded from the immunization registry reporting measure if the EP—

- Does not administer any immunizations to any of the populations for which data is collected by their jurisdiction’s immunization registry or immunization information system during the EHR reporting period;
- Operates in a jurisdiction for which no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no immunization registry or immunization information system has declared readiness to receive immunization data as of 6 months prior to the start of the EHR reporting period.

**Measure 2:**
Any EP meeting one or more of the following criteria may be excluded from the syndromic surveillance reporting measure if the EP—

- Is not in a category of providers from which ambulatory syndromic surveillance data is collected by their jurisdiction’s syndromic surveillance system;
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic syndromic surveillance data from EPs in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period; or
- Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from EPs as of 6 months prior to the start of the EHR reporting period.
Clinical Quality Measures (CQMs) Reporting:

EPs will select and report on 6 of 53 Clinical Quality Measures without the 6 domain reporting requirements.

Here is what Eligible Hospitals need to know about Stage 3 Meaningful Use:

<table>
<thead>
<tr>
<th>Stage 3 Meaningful Use Objectives (beginning January 1, 2018)</th>
<th>Stage 3 Meaningful Use Measures:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protect Electronic Protected Health Information (PHI)</td>
<td>Measure:</td>
</tr>
<tr>
<td></td>
<td>Conduct or review a security risk analysis in accordance with the requirements under 45 CFR 164.308(a)(1), including addressing the security (including encryption) of data created or maintained by CEHRT in accordance with requirements under 45 CFR 164.312(a)(2)(iv) and 45 CFR 164.306(d)(3), implement security updates as necessary, and correct identified security deficiencies as part of the provider's risk management process.</td>
</tr>
<tr>
<td></td>
<td>Exclusion:</td>
</tr>
<tr>
<td></td>
<td>There is no exclusion for this Stage 3 Meaningful Use Objective</td>
</tr>
<tr>
<td>Transmitting Electronic Prescriptions</td>
<td>Measure:</td>
</tr>
<tr>
<td></td>
<td>More than 25 percent of hospital discharge medication orders for permissible prescriptions (for new and changed prescriptions) are queried for a drug formulary and transmitted electronically using CEHRT.</td>
</tr>
<tr>
<td></td>
<td>Exclusion:</td>
</tr>
<tr>
<td></td>
<td>Any eligible hospital or CAH that does not have an internal pharmacy that can accept electronic prescriptions and there are no pharmacies that accept electronic prescriptions within 10 miles at the start of their EHR reporting period.</td>
</tr>
<tr>
<td>Clinical Decision Support</td>
<td>In order for eligible hospitals and CAHs to meet the objective they must satisfy both of the following measures:</td>
</tr>
<tr>
<td></td>
<td>Measure 1:</td>
</tr>
<tr>
<td></td>
<td>Implement five clinical decision support interventions related to four or more CQMs at a relevant point in patient care for the entire EHR reporting period. Absent four CQMs related to an eligible hospital or CAH's scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions.</td>
</tr>
<tr>
<td></td>
<td>Measure 2:</td>
</tr>
<tr>
<td></td>
<td>The eligible hospital or CAH has enabled and implemented the</td>
</tr>
<tr>
<td><strong>Computerized Provider Order Entry (CPOE)</strong></td>
<td><strong>Patient Electronic Access</strong></td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>An eligible hospital/CAH must meet the thresholds for all three measures:</td>
<td>Eligible Hospitals and CAHs must satisfy both measures in order to meet the objective:</td>
</tr>
<tr>
<td>Measure 1: More than 60 percent of medication orders created by the authorized providers of the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.</td>
<td>Measure 1: For more than 80 percent of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23):</td>
</tr>
<tr>
<td>Measure 2: More than 60 percent of laboratory orders created by the authorized providers of the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.</td>
<td></td>
</tr>
<tr>
<td>Measure 3: More than 60 percent of diagnostic imaging orders created by the authorized providers of the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.</td>
<td></td>
</tr>
<tr>
<td>There are no exclusions for the Stage 3 Meaningful Use Objective.</td>
<td>The patient (or the patient authorized representative) is provided timely access to view online, download, and transmit his or her health information; and</td>
</tr>
<tr>
<td></td>
<td>The provider ensures the patient’s health information is available for the patient (or patient authorized representative) to access using any application of their choice that is configured to meet the technical specifications of the API in the provider’s CEHRT.</td>
</tr>
<tr>
<td>Measure 2:</td>
<td>The eligible hospital or CAH must use clinically relevant information from CEHRT to identify patient-specific educational resources and provide electronic access to those materials to more than 35 percent of unique patients seen by the EP or discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period.</td>
</tr>
</tbody>
</table>

**Exclusion:**

**Measures 1 and 2:**
Any eligible hospital or CAH will be excluded from the measure if it is located in a county that does not have 50 percent or more of their housing units with 4Mbps broadband availability according to the latest information available from the FCC at the start of the EHR reporting period.

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| Coordination of Care | Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective: |

**Measure 1:**
For an EHR reporting period in 2017, more than 5 percent of all unique patients (or their authorized representatives) discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) actively engage with the electronic health record made accessible by the provider and either—
1. View, download or transmit to a third party their health information; or
2. Access their health information through the use of an API that can be used by applications chosen by the patient and configured to the API in the provider's CEHRT; or
3. A combination of (1) and (2)

**Threshold for 2018 and Subsequent Years:** The resulting percentage must be more than 10 percent.

**Measure 2:**
For an EHR reporting period in 2017, more than 5 percent of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient authorized representative), or in response to a secure message sent by the patient or their authorized representative.
<table>
<thead>
<tr>
<th>Threshold in 2018 and Subsequent Years:</th>
<th>The resulting percentage must be more than 25 percent in order for an EP, eligible hospital, or CAH to meet this measure.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure 3:</td>
<td>Patient generated health data or data from a nonclinical setting is incorporated into the CEHRT for more than 5 percent of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period.</td>
</tr>
<tr>
<td>Exclusion:</td>
<td>Any eligible hospital or CAH will be excluded from the measure if it is located in a county that does not have 50 percent or more of their housing units with 4Mbps broadband availability according to the latest information available from the FCC at the start of the EHR reporting period.</td>
</tr>
<tr>
<td>Health Information Exchange</td>
<td>Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective.</td>
</tr>
<tr>
<td>Measure 1:</td>
<td>For more than 50 percent of transitions of care and referrals, the eligible hospital or CAH that transitions or refers their patient to another setting of care or provider of care: 1) Creates a summary of care record using CEHRT; and 2) Electronically exchanges the summary of care record.</td>
</tr>
<tr>
<td>Measure 2:</td>
<td>For more than 40 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the eligible hospital or CAH incorporates into the patient’s EHR an electronic summary of care document.</td>
</tr>
<tr>
<td>Measure 3:</td>
<td>For more than 80 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the eligible hospital or CAH performs a clinical information reconciliation. The provider must implement clinical information reconciliation for the following three clinical information sets: 1) Medication. Review of the patient’s medication, including the name, dosage, frequency, and route of each medication. 2) Medication allergy. Review of the patient’s known medication allergies. 3) Current Problem list. Review of the patient’s current and active diagnoses.</td>
</tr>
</tbody>
</table>
Exclusions:

**Measure 1:**
Any eligible hospital or CAH will be excluded from the measure if it is located in a county that does not have 50 percent or more of their housing units with 4Mbps broadband availability according to the latest information available from the FCC at the start of the EHR reporting period.

**Measure 2:**
A provider may exclude from the measure if any of the following apply:
- Any eligible hospital or CAH for whom the total of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, is fewer than 100 during the EHR reporting period is excluded from this measure.
- Any eligible hospital or CAH that is located in a county that does not have 50 percent or more of their housing units with 4Mbps broadband availability according to the latest information available from the FCC at the start of the EHR reporting period.

**Measure 3:**
Any eligible hospital or CAH for whom the total of transitions or referrals received and patient encounters in which the provider has never before

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**Public Health Reporting**

**Measure 1:**
Immunization Registry Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit immunization data and receive immunization forecasts and histories from the public health immunization registry/immunization information system (IIS).

**Measure 2:**
Syndromic Surveillance Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit syndromic surveillance data from an urgent care setting.

**Measure 3:**
Electronic Case Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit case reporting of reportable conditions.

NOTE: Electronic Case Reporting is not required until 2018.
| Measure 4 – Public Health Registry Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit data to public health registries. |
| Measure 5 – Clinical Data Registry Reporting: The eligible hospital or CAH is in active engagement to submit data to a clinical data registry. |
| Measure 6 – Electronic Reportable Laboratory Result Reporting: The eligible hospital or CAH is in active engagement with a public health agency to submit electronic reportable laboratory results. |
Appendix J: Post-Payment Audit Strategy for Meaningful Use

Appendix J will be submitted to CMS separate from this SMHP update to maintain confidentiality.
Appendix K: Meaningful Use Screenshots

DOM submitted an SMHP Addendum for the 2015-2017 Modifications for the CMS Final Rule in January 2016, including screen shots of the:

- Proposed SLR Attestation Portal Screen Shots for Eligible Professionals, and
- Proposed SLR Attestation Portal Screen Shots for Eligible Hospitals.

CMS approved the SMHP Addendum on February 5, 2016. Following are the screenshots that were included in the Addendum.
Protect Patient Health Information

**Objective:** Protect electronic health information created or maintained by the CEHRT through the implementation of appropriate technical capabilities.

**Measure:** Conduct or review a security risk analysis in accordance with the requirements in 45 CFR 164.308(a)(1), including addressing the security (to include encryption) of PHI created or maintained by CEHRT in accordance with requirements under 45 CFR 164.312 (a)(2)(iv) and 45 CFR 164.306(d)(3), and implement security updates as necessary and correct identified security deficiencies as part of the eligible hospital or CAH risk management process.

Replace text regarding the help text. Help text is configurable for each client and is hidden by default.

Complete the following information:

* Have you conducted or reviewed a security risk analysis in accordance with the requirements?

  - No
  - Yes

* Date security risk analysis was completed:

  Will display with calendar selector

Clinical Decision Support

**Objective:** Please select which objective you will report:

- Stage 1 Objective: Implement one clinical decision support rule relevant to specialty or high priority hospital condition, along with the ability to track compliance with that rule.

- Stage 2 Objective: Use clinical decision support to improve performance on high-priority health conditions.

**Measure:** Implement one clinical decision support rule.

Complete the following information:

* Have you implemented one clinical decision support rule relevant to specialty or high clinical priority along with the ability to track compliance to that rule?

  - Yes
  - No

* Please enter the name of one Clinical Decision Support Rule you have implemented:
Computerized Provider Order Entry (CPOE)

Objective: Use computerized provider order entry (CPOE) for medication, laboratory, and radiology orders directly entered by any licensed healthcare professional who can enter orders into the medical record per state, local, and professional guidelines.

Please select which measures you will report:
- Stage 1 Alternative Exclusions and Specifications
- Stage 2 Measures

Exclusion Criteria: Meeting the following criteria qualifies for the exclusion of the Associated measure.

- Do you wish to exclude Measure #2 as there is no equivalent Stage 1 measure? (Measure #2)
  - No
  - Yes

- Do you wish to exclude Measure #3 as there is no equivalent Stage 1 measure? (Measure #3)
  - No
  - Yes

Measure #1: Please select which measure you will report:

- More than 30% of all unique patients with at least one medication in their medication list admitted to eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reporting period have at least one medication order entered using CPOE.
- More than 30% of medication orders created by the authorized providers of the eligible hospital’s or CAH’s for patients admitted to their inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using CPOE.

Complete the following information:

Numerator = The number of patients in the denominator that have at least one medication order entered using CPOE

Denominator = Number of unique patients with at least one medication in their medication list seen by the eligible hospital or CAH during the EHR reporting period.

Measure #2: More than 30% of laboratory orders created by authorized providers of the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using CPOE.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.

Measure #3: More than 30% of radiology orders created by authorized providers of the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using CPOE.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.

Appendix K: Meaningful Use Screenshots
Electronic Prescribing

Objective: Generate and transmit permissible discharge prescriptions electronically (eRx).

Exclusion Criteria: Meeting the following criteria qualifies for the exclusion for this measure.

- Do you have an internal pharmacy that can accept electronic prescriptions or any pharmacy that accepts electronic prescriptions within 10 miles of the hospital? No Yes

Stage 1 Exclusion: Do you wish to exclude the measure as there is no equivalent Stage 1 measure? No Yes

Measure: More than 10% of hospital discharge medication orders for permissible prescriptions (for new, and changed prescriptions) are queried for a drug formulary and transmitted electronically using CEHRT.

Complete the following information:

- Numerator = The number of prescriptions in the denominator generated, queried for a drug formulary and transmitted electronically.
- Denominator = Number of new or changed permissible prescriptions written for drugs requiring a prescription in order to be dispensed other than controlled substances for patients discharged during the EHR reporting period.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.
Health Information Exchange

Objective: The eligible hospital or CAH who transitions their patient to another setting of care or provider of care or refers their patient to another provider of care provides a summary of care record for each transition of care or referral.

Exclusion Criteria: Meeting the following criteria qualifies for the exclusion for this measure.

Stage 1 Exclusion: Do you wish to exclude the measure as there is no equivalent Stage 1 measure? ☐ No ☑ Yes

Measure: The eligible hospital or CAH that transitions or refers their patient to another setting of care or provider of care must - (1) use CEHRT to create a summary of care record; and (2) electronically transmits such summary to a receiving provider for more than 10% of transitions of care and referrals.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.

Patient-Specific Education

Objective: Use clinically relevant information from CEHRT to identify patient-specific education resources and provide those resources to the patient.

Exclusion Criteria: Meeting the following criteria qualifies for the exclusion for this measure.

Stage 1 Exclusion: Do you wish to exclude the measure as you were not intending to attest to the Stage 1 menu objective? ☐ No ☑ Yes

Measure: More than 10% of all unique patients admitted to the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reportin period are provided patient specific education resources identified by CEHRT.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.
**Medication Reconciliation**

**Objective:** The eligible hospital or CAH that receives a patient from another setting of care or provider of care or believes an encounter is relevant performs medication reconciliation.

**Exclusion Criteria:** Meeting the following criteria qualifies for the exclusion for this measure.

**Stage 1 Exclusion:** Do you wish to exclude the measure as you were not intending to attest to the Stage 1 menu objective?

- ☐ No
- ☑ Yes

**Measure:** The eligible hospital or CAH performs medication reconciliation for more than 50% of transitions of care in which the patient is admitted to the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23).

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.

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**Patient Electronic Access**

**Objective:** Provide patients the ability to view online, download, and transmit information within 36 hours of hospital discharge.

**Exclusion Criteria:** Meeting either of the following criteria qualifies for the exclusion for the associated measure.

- Is the hospital or CAH located in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #2)
  - ☐ No
  - ☑ Yes

  **Stage 1 Exclusion:** Do you wish to exclude Measure #2 as there is no equivalent Stage 1 measure?
  - ☐ No
  - ☑ Yes

**Measure #1:** More than 50% of all unique patients who are discharged from the inpatient or emergency department (POS 21 or 23) of an eligible hospital or CAH are provided timely access to view online, download and transmit to a third party their health information.

Complete the following information:

1. **Numerator:** The number of patients in the denominator who are able to access their health information within 36 hours after the information is available to the eligible hospital or CAH.
2. **Denominator:** Number of unique patients discharged from an eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reporting period.

**Measure #2:** For an EHR reporting period in 2015 and 2016, at least 1 patient who is discharged from the inpatient or emergency department (POS 21 or 23) of an eligible hospital or CAH (or patient-authorized representative) views, downloads, transmits to a third party his or her health information during the EHR reporting period.

You meet the requirement for exclusion from this measure. Exclusion from the requirement does not prevent an eligible hospital or CAH from achieving Meaningful Use.
Public Health Reporting

Objective: The eligible hospital or CAH is in active engagement with a Public Health Agency (PHA) or clinical data registry (CDR) to submit electronic public health data in a meaningful way using certified EHR technology, except where prohibited and in accordance with applicable law and practice.

In order to meet this objective, eligible hospitals and CAHs must meet two of the total number of measures available to them. Reporting an exclusion for a measure does not qualify towards meeting the objective unless the eligible hospital or CAH can report on fewer than two measures. If an eligible hospital or CAH can report on fewer than 2 measures, the eligible hospital or CAH must report on any possible measures and claim the exclusion for the remaining measures.

Eligible hospitals and CAHs may choose to report more than one public health registry or clinical data registry to meet the number of measures required to meet the objective. Up to 3 public health registries of clinical data registries may be reported.

Select “I will report on this measure” to report for the specific measure. Select “I will claim exclusion for this measure” to claim exclusion for the specific measure.

Active Engagement: If your state has one public health agency that manages registration for all the public health measures, you may use this option to select the level of active engagement to apply to all measures with reporting to a Public Health Agency. Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Complete Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days, failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- Option 3 - Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

<table>
<thead>
<tr>
<th>Measure</th>
<th>I will report on this measure</th>
<th>I will claim exclusion for this measure</th>
</tr>
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<tbody>
<tr>
<td>Measure 1 - Immunization Registry Reporting</td>
<td>☐</td>
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<tr>
<td>Measure 2 - Syndromic Surveillance Reporting</td>
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<td>Measure 3 - Specialized Registry Reporting</td>
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<td>Measure 3 - Specialized Registry Reporting</td>
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<tr>
<td>Measure 4 - Electronic Reportable Laboratory Results</td>
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Please select the ‘Previous Screen’ button to go back or the ‘Save & Continue’ button to proceed.

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Measure 1 - Immunization Registry Reporting

Measure: The eligible hospital or CAH is in active engagement with a Public Health Agency (PHA) to submit immunization data.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply:

- Does not administer any immunizations to any of the populations for which data is collected by their jurisdiction's immunization registry or immunization information system during the EHR reporting period.
- Operates in a jurisdiction where no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEHRT definition at the start of the EHR reporting period.
- Operates in a jurisdiction where no immunization registry or immunization information system has declared readiness to receive immunization data from the eligible hospital or CAH at the start of the EHR reporting period.
- Alternate Exclusion: Did not intend to select the menu objective for an EHR reporting period in 2015.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- Option 3 - Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

Attach Files

The following attachments are optional:
- Other Attachment

File Name | Subject | Remove
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No records to display.

Add Files | Remove Selected

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Previous Screen | Save & Continue
Measure 2 - Syndromic Surveillance Reporting

**Measure:** The eligible hospital or CAH is in active engagement with a Public Health Agency (PHA) to submit syndromic surveillance data.

**Exclusion Criteria:**
- Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.
- Does not have an emergency or urgent care department
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic syndromic surveillance data from eligible hospitals or CAHs in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period
- Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from eligible hospitals or CAHs at the start of the EHR reporting period.
- **Alternate Exclusion:** Did not intend to select the menu objective for an EHR reporting period in 2015.

**Active Engagement:** Select the level of active engagement you demonstrate for this measure. Only one level may be selected.
- **Option 1 - Completed Registration to Submit Data:** The eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted, registration was completed within 60 days after the start of the EHR reporting period, and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- **Option 2 - Testing and Validation:** The eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days: failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- **Option 3 - Production:** The eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

**Attach Files**
The following attachments are optional
- Other Attachment

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<thead>
<tr>
<th>File Name</th>
<th>Subject</th>
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Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.
Measure 3 - Specialized Registry Reporting

**Measure:** The eligible hospital or CAH is in active engagement with a Public Health Agency (PHA) to submit data to a specialized registry.

**Exclusion Criteria:** Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- [ ] Does not diagnose or treat any disease or condition associated with or collect relevant data that is required by a specialized registry in their jurisdiction during the EHR reporting period.
- [ ] Operates in a jurisdiction for which no public health agency is capable of accepting electronic registry transactions in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period.
- [ ] Operates in a jurisdiction where no public health agency for which the eligible hospital or CAH is eligible has declared readiness to receive electronic registry transactions at the start of the EHR reporting period.
- [ ] **Alternate Exclusion:** Did not intend to select the menu objective for an EHR reporting period in 2015.

**Active Engagement:** Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- [ ] **Option 1 - Completed Registration to Submit Data:** the eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted; registration was completed within 90 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- [ ] **Option 2 - Testing and Validation:** the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- [ ] **Option 3 - Production:** the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

**Registry Name:** (please enter name of registry)

**Attach Files**

The following attachments are optional:

- Other Attachment

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Previous Screen  Save & Continue
Measure 4 - Reportable Electronic Laboratory Results Reporting

Measure: The eligible hospital or CAH is in active engagement with a public health agency (PHA) to submit electronic reportable laboratory (ELR) results.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Does not perform or order laboratory tests that are reportable in their jurisdiction during the EHR reporting period.
- Operates in a jurisdiction for which no public health agency is capable of accepting the specific ELR standards required to meet the CEHRT definition at the start of the EHR reporting period.
- Operates in a jurisdiction where no public health agency has declared readiness to receive electronic reportable laboratory results from eligible hospitals or CAHs at the start of the EHR reporting period.

Alternate Exclusion: Did not intend to select the menu objective for an EHR reporting period in 2015.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- Option 3 - Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

Public Health Agency Name: (please enter name of Public Health Agency)

Attach Files

The following attachments are optional:

- Other Attachment

File Name Subject Remove

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Please select the ‘Previous screen’ button to go back or the ‘Save & Continue’ button to proceed.

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1 OVERVIEW

1.1 Background

In October 2015, Centers for Medicare & Medicaid Services (CMS) released a final rule that specifies the meaningful use (MU) objectives that eligible professionals (EPs), eligible hospitals (EHs), and critical access hospitals (CAHs) must meet in order to continue to participate in the Medicare and Medicaid Electronic Health Record (EHR) Incentive programs.

The provisions of this rule specify Modified Meaningful Use requirements that will apply for providers attesting to Stage 2 in 2017, including text or threshold changes to the measures for Patient Electronic Access and Secure electronic Messaging. In addition, this rule requires that providers have the option to attest to Stage 3 in 2017.

In November 2016, CMS released a final rule that updated payment rates and policy changes in the Hospital Outpatient Prospective Payment System (OPPS) and Ambulatory Surgical Center (ACS) Payment System for calendar year 2017.

One of the provisions of this rule extends the 90-day EHR reporting period for program year 2017, allowing providers to select any continuous 90-day period between January 1 and December 31 in calendar year 2017. Another provision requires that all actions included in the numerator of Meaningful Use measures must occur within the HER reporting period if that period is a full calendar year, or if less than a full calendar year, within the calendar year in which the HER reporting period occurs.

Also in November 2016, CMS finalized a rule establishing the Merit Based Incentive Payment System (MIPS), which consolidates components of three existing programs into a single, cohesive program called the Quality Payment Program. The Quality Payment Program focuses on quality, cost, and use of certified EHR technology to support interoperability and advanced quality objectives.

The purpose of this addendum is to describe Mississippi’s plan for implementing the provisions in
each of these new rules that affect Program Year 2017 for Mississippi’s Medicaid EHR Incentive Program. No changes are required in Mississippi’s IAPD to implement this new rule.

### 1.2 Policy Considerations and Audit Strategy Adjustments

Supporting documentation for meaningful use measures will be required in Program Year 2017 for eligible professionals. The State Level Registry accepts the Meaningful Use data from CMS for its dually eligible hospitals. These include the summary MU reports generated by certified EHR systems, a summary document for the Security Risk Assessment, and written confirmation from the State’s Public Health Agency for active engagement and/or qualification for exclusions. Additionally, providers will have the option to attach supporting documentation for exclusions for any measure. The internal system configuration document reflects these requirements, which will be tested during UAT prior to implementing the system changes for Program Year 2017.

Mississippi will update its post payment audit procedures to incorporate requirements for rule changes that apply to Program Year 2017 in a future update of the Audit Appendix.

## 2 SYSTEM CHANGES

The Conduent State Level Registry (SLR) User Group Community has reviewed all CMS requirements with the Xerox team to assure changes to the portal will accommodate the requirements of the new rule.

### 2.1 2015 – 2017 Modifications Rule

#### Eligible Professional (EP) Option to Attest to Stage 3 in 2017

On the initial Meaningful Use (MU) page in the State Level Registry, EPs attesting for 2017 MU are presented with a selection option to report on either Stage 3 Objectives or to report on Stage 2 Objectives. (Please see Screen shots in Appendices)

This option will be used as selection screen for PY 2017 for both EHs and EPs:

---

**Meaningful Use**

Please select which Stage of MU you will report for 2017. You must attest to Stage 2 objectives if you are a new participant in 2017. You must have 2016 edition certified EHR technology if you attest to Stage 3 Objectives. You may not change your MU Stage selection option on individual MU objectives pages.

- [ ] I will report Stage 2 objectives in 2017
- [ ] I will report Stage 3 objectives in 2017 (you must have 2015 edition CEHRT to attest to Stage 3 in 2017)

---

When 2017 Stage 3 MU Objective option is selected, the EP is presented with a summary of 2017 Stage 3 Objectives. From the Summary page, the EP can navigate to any specific Stage 3 Objectives. The table below shows Objective name and sequence, which differs from Stage 2 objectives in 2017:
Upon advancing from the selection page to the detailed MU pages, each Stage 3 Objective displays with 2017 Objective, measure text, and relevant exclusion criteria per the final rule. Five Public Health measures options display for EP in 2017 under the Stage 3 selection option. EPs must report on two measures. The State Level Registry also allows Coordination of Care and Health Information Exchange objectives to pass validation when the thresholds are met for at least two of the three measures.

Validations that prevent Stage 3 section options from displaying are as follows:

- Stage 3 MU option is disabled for providers that do not have a 2015 edition CEHRT validation on EHR Certification Page in Conduent SLR. In this scenario, providers can only attest to Stage 2 objectives and measures.
- Stage 3 MU option is disabled for providers that have not successfully demonstrated MU in a prior year. Validation is based on B6 data and/or prior year(s) attestation data in Conduent SLR.

When 2017 Stage 2 MU Objective option is selected, the EP is presented with a summary of 2017 Stage 2 Objectives. From the Summary page, the EP can navigate to any specific Stage 2 Objectives. There are two objectives with measure text changes or threshold changes per the final rule:

- Objective 8: Patient Electronic Access
- Objective 9: Secure Electronic Messaging

**Eligible Hospital (EH) Option to Attest to Stage 3 in 2017**

On the initial Meaningful Use page in the State Level Registry, EHs attesting for 2017 MU are presented with a selection option to report on either Stage 3 Objectives or to report on Stage 2 Objectives.

When 2017 Stage 3 MU Objective option is selected, the EH is presented with a summary of 2017 Stage 3 Objectives. From the Summary page, the EH can navigate to any specific Stage 3 Objectives. The table below shows Objective name and sequence, which differs from Stage 2 objectives in 2017:

<table>
<thead>
<tr>
<th>Stage 3 Objective</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protect Patient Health Information</td>
</tr>
</tbody>
</table>

---

**Appendix K: Meaningful Use Screenshots**

---

**Updated**

State Medicaid Health Information Technology Planning Document

November 3, 2017
Updated
State Medicaid Health Information Technology
Planning Document
November 3, 2017

<table>
<thead>
<tr>
<th>Electronic Prescribing (eRx)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Decision Support</td>
</tr>
<tr>
<td>Computerized Provider Order Entry (CPOE)</td>
</tr>
<tr>
<td>Patient Electronic Access</td>
</tr>
<tr>
<td>Coordination of Care</td>
</tr>
<tr>
<td>Health Information Exchange</td>
</tr>
<tr>
<td>Public Health and Clinical Data Registry Reporting</td>
</tr>
</tbody>
</table>

Upon advancing from the selection page to the detailed MU pages, each Stage 3 Objective displays with 2017 objective, measure text, and relevant exclusion criteria per the final rule. Six Public Health measures options display for EH in 2017 under the Stage 3 selection option. EHs must report on four measures. The State Level Registry also allows Coordination of Care and Health Information Exchange objectives to pass validation when the thresholds are met for at least two of the three measures.

The Conduent SLR will populate MU objective data from the C5 received in 2017, which will not contain data for Clinical Decision Support and CPOE objectives – the Conduent SLR will pass validation for these objectives and allow the EH to advance in the attestation process.

Validations occur that prevent Stage 3 section options for the following reasons:

- Stage 3 MU option is disabled for providers that do not have a 2015 edition CEHRT validation on EHR Certification Page in Conduent SLR. In this scenario, providers can only attest to Stage 2 objectives and measures.
- Stage 3 MU option is disabled for providers that have not successfully demonstrated MU in a prior year. Validation is based on B6 data and/or prior year(s) attestation data in Conduent SLR.

When 2017 Stage 2 MU Objective option is selected, the EH is presented with a summary of 2017 Stage 2 Objectives. From the Summary page, the EH can navigate to any specific Stage 2 Objectives. There is one EH objective with measure text changes or threshold change per the final rule:

- Objective 8: Patient Electronic Access

**Program Year 2017 MU Requirements**

As indicated previously, all providers are presented with the option to report on either Stage 3 or Stage 2 MU.

For Eligible Professionals (EPs), when 2017 Stage 2 MU Objective option is selected, the system displays a summary of 2017 Stage 2 Objectives. From the Summary page, the EP can navigate to any specific Stage 2 Objectives. The system displays the measure threshold changes for 2017, per the final rule, for the following:

- Objective 8: Patient Electronic Access
Objective 9: Secure Electronic Messaging

The Conduent SLR will not allow login for EPs who did not participate in EHR Incentive Program prior to the end of program year 2016. The system validates B6 data and prior year data in the Conduent SLR database.

For Eligible Hospitals (EHs), when 2017 Stage 2 MU Objective option is selected, the system displays a summary of 2017 Stage 2 Objectives. From the Summary page, the EH can navigate to any specific Stage 2 Objectives. The system displays the measure threshold changes for 2017, per the final rule, for the following:

- Objective 8: Patient Electronic Access

The Conduent SLR will not allow login for an EHs that was not paid in the prior year. The system validates B6 data and prior year data in the Conduent SLR database.

The Conduent SLR disabled AIU in 2017 for both EP and EH.

2.2 Outpatient Prospective Payment System (OPPS)

Rule

Eligible 90-day EHR Reporting Period

The workflow for Eligible Professionals and Eligible Hospitals allows the provider to report using an EHR reporting period of any continuous 90-day period in calendar year 2017, regardless of prior attestations, while retaining the standard reporting period requirements for subsequent years. As of the published date of this document, the system requires a full year CQM reporting period for returning MU providers.

Modification to Measure Calculation Timeframe

There is no system change to validate that actions included in the numerator data occur within the calendar year in which the EHR reporting period occurs. The Conduent SLR validates that EHR reporting period dates are in the calendar year 2017 and displays an attestation statement at the EHR Reporting Period page that the numerator and denominator data are for the reporting period.

On each MU objective page, users are required to enter data in numerator and denominator fields specific for applicable MU measures. The State Level Registry calculates the percentage based on the data entered by the user. The threshold is met when the calculated percentage meets or exceeds the requirement mandated by CMS. The system displays a confirmation that the user has meet the MU objective when all measure(s) meet the threshold(s).

2.3 Medicare Quality Payment Program
Updates to Definition of Meaningful EHR

User
For program year 2017, the Conduent SLR displays two attestation statements on the EHR Certification page. These statements are related to supporting providers with the performance of CEHRT (SPPC). Users are required to select a check box indicating their confirmation they engaged in SPPC activities as stated by the rule. A second check box is optional to select. Selection of this checkbox indicates their confirmation of engagement in surveillance of health information, as stated by the rule.

On the same EHR Certification page, the system displays an attestation statement related to the support for health information exchange and the prevention of information blocking. Users are also required to select a check box indicating their confirmation they engaged in prevention of health information blocking, as stated by the rule.

When the two required attestation statements are selected, the system allows the user to continue to the next step. The optional attestation statement checkbox is not required in order for the user to advance to the next step. If the user fails to select required checkboxes, the system displays an error message that the fields are required and the user cannot advance until the selections are made.

2.4 Screenshots and Program Year Extensions

Screen shots for both EPs and EHs are shown separately for Stage 2 and Stage 3 in Appendices of this document.

Screen Shot Appendices:
   2.4.1 EP - Modified Stage 2 changes - 2017
   2.4.2 EH - Modified Stage 2 changes - 2017
   2.4.3 EP - Stage 3 - 2017
   2.4.4 EH - Stage 3 – 2017

Mississippi plans to open Program Year 2017 in early January of 2018. This will allow providers to use a 90-day EHR reporting period and a 365-day CQM reporting period (under current CMS regulations) for Program Year 2017.

We will determine actual starting dates for PY 2017 once we have an estimate of the time to implement and test system changes. Requests for extensions to the grace periods will be submitted to our CMS program contact.

At present, Mississippi’s grace period for Program Year 2016 for EP attestations is in effect through the April 30, 2017 for Eligible Hospitals and Eligible Professionals.

Stage 2 EP 2017 – Objectives with measure text changes in 2017

2.4.1 EP - Modified Stage 2 Screen Shots
<table>
<thead>
<tr>
<th>Objective:</th>
<th>Provide patients the ability to view online, download, and transmit their health information within 4 business days of the information being available to the EP.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exclusion Criteria:</td>
<td>Meeting the following criteria qualifies for the exclusion for Measure #1 only: Did you order or create any of the information listed as part of this measure except for “Patient name” and “Provider's name and office contact information”? (Measure #1)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Meeting either of the following criteria qualifies for the exclusion for Measure #2 only. Did you order or create any of the information listed as part of this measure except for “Patient name” and “Provider's name and office contact information”? (Measure #2)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Did you conduct 50% or more of your encounters in a county/area that does not have more than 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #2)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>No</td>
</tr>
</tbody>
</table>

**Measure #1:** More than 50 percent of all unique patients seen by the EP during the EHR reporting period are provided timely access to view online, download, and transmit to a third party their health information subject to the EP’s discretion to withhold certain information.

Complete the following information:

- **Numerator** = The number of patients in the denominator who have access to view online, download, and transmit their health information within 4 business days after the information is available to the EP.

- **Denominator** = Number of unique patients seen by the EP during the EHR reporting period.

**Measure #2:** For an EHR reporting period in 2017, more than 5 percent of unique patients seen by the EP during the EHR reporting period (or his or her authorized representative) view, download, or transmit to a third party their health information during the EHR reporting period.

Complete the following information:

- **Numerator** = The number of patients in the denominator who view, download, or transmit to a third party their health information.

- **Denominator** = The number of patients seen by the EP during the EHR reporting period.

**Attach Files**

- The following attachments are optional.
- Other Attachment
Secure Electronic Messaging

Red asterisk indicates a required field.

Objective: Use secure electronic messaging to communicate with patients on relevant health information.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure.

- Did you have office visits during the EHR reporting period?  
  - ☐ No  ☐ Yes

- Did you conduct 50% or more of your encounters in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? 
  - ☐ No  ☐ Yes

Measure: For an EHR reporting period in 2017, for more than 5 percent of unique patients seen by the EP during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient-authorized representative), or in response to a secure message sent by the patient (or the patient-authorized representative) during the EHR reporting period.

Complete the following information:

- Numerator = The number of patients in the denominator for whom a secure electronic message is sent to the patient (or patient-authorized representative), or in response to a secure message sent by the patient (or patient-authorized representative).

- Denominator = Number of unique patients seen by the EP during the EHR reporting period.

Attach Files

The following attachments are optional:

- Other Attachment

2.4.2 EH - Modified Stage 2 Screen Shots
### Patient Electronic Access

- **Objective:** Provide patients the ability to view online, download, and transmit information within 36 hours of hospital discharge.

- **Exclusion Criteria:** Meeting the following criteria qualifies for Measure #2 only.
  - Is the hospital or CAH located in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #2)

#### Measure #1:

More than 50% of all unique patients who are discharged from the inpatient or emergency department (POSO 21 or 23) of an eligible hospital or CAH are provided timely access to view online, download and transmit to a third party their health information.

**Complete the following information:**

- **Numerator:** The number of patients in the denominator who are have access to view, download, and transmit their health information within 36 hours after the information is available to the eligible hospital or CAH.

- **Denominator:** Number of unique patients discharged from an eligible hospital's or CAH's inpatient or emergency department (POSO 21 or 23) during the EHR reporting period.

#### Measure #2:

For an EHR reporting period in 2017, more than 5 percent of unique patients discharged from the inpatient or emergency department (POSO 21 or 23) of an eligible hospital or CAH (or patient-authorized representative) view, downloads or transmits to a third party his or her health information during the EHR reporting period.

**Complete the following information:**

- **Numerator:** The number of patients (or patient-authorized representative) who view, download, or transmit to a third party their health information.

- **Denominator:** Number of unique patients discharged from the inpatient or emergency department (POSO 21 or 23) of the eligible hospital or CAH during the EHR reporting period.

### Attach Files

- The following attachments are optional:
  - Other Attachment

---

**2.4.3 EP – Stage 3 Screen Shots**
3. Attestation of EHR

Important Note - Please Read!

As part of the American Recovery and Reinvestment Act of 2009 (ARRA) Congress mandated payment adjustments to be applied to Medicare eligible professionals (EPs) and eligible hospitals (EHs) who do not demonstrate meaningful use of Certified Electronic Health Record (EHR) Technology by 2014. Payment adjustments were mandated to begin on the first day of 2014. Medicaid only hospitals and Medicaid EPs who do not bill Medicare are not subject to these payment adjustments, however all other providers are subject to the penalties regardless of which program they choose to participate in. This means that EPs participating in the Medicaid EHR Incentive Program who bill Medicare are subject to the payment adjustment if they do not demonstrate Meaningful Use.

Providers who first attested to MU in 2011 or 2012 must have demonstrated meaningful use for a full year in 2013 to avoid payment adjustments in 2015, and must continue to demonstrate meaningful use every year to avoid payment adjustments in subsequent years.

Providers who first attested to MU in 2013 must have demonstrated meaningful use for a full year in 2014 to avoid payment adjustments in 2015, and must continue to demonstrate meaningful use every year to avoid payment adjustments in subsequent years.

Providers who first demonstrate meaningful use in 2014 must have demonstrated meaningful use for a full year in 2015 to avoid payment adjustments in 2016. The reporting period must have occurred in the first 9 months of the program year 7/1/14 for eligible hospitals and 10/1/14 for eligible professionals in order to avoid payment adjustments. Providers must continue to demonstrate meaningful use every year to avoid payment adjustments for providers who first demonstrate meaningful use in 2014.

Providers who first demonstrate meaningful use in 2016 must have demonstrated meaningful use for a full year in 2017 to avoid payment adjustments in 2018. This reporting period must occur in the first 10 months of calendar year 2017, in order to avoid the payment adjustments in calendar year 2018.

In certain circumstances, eligible hospitals and eligible professionals may apply for hardship exceptions to avoid the payment adjustments. Hardship exceptions will be granted only under certain circumstances and only if CMS determines that providers have demonstrated that those circumstances pose a significant barrier to their achieving meaningful use. CMS will publish information on how to apply for a hardship exception on the CMS EHR Incentive Programs website (www.CMS.gov/Regulations-and-Guidance/Legislation/EHRIncentivePrograms).

For more information on payment adjustments and hardship exceptions, please select one of the links below:

Payment Adjustments and Hardship Exceptions Tipsheet for Eligible Professionals

Click here for a printable version of this notice.

Attest to Adopt, Implement, Upgrade

Attest to Meaningful Use

AIU is not an option in 2017. 2016 was the last program year for eligible professionals and hospitals to start the EHR Incentive Program.

Select this option to attest to demonstrating Meaningful Use of certified EHR Technology by:

1. Use of certified EHR technology in a meaningful manner, such as e-prescribing;
2. That the certified EHR technology is connected in a manner that provides for the electronic exchange of health information to improve the quality of care; and
3. Using certified EHR technology to submit information on clinical quality measures and other such measures of Meaningful Use.
### EHR Certification

#### EHR Certification

Providers must provide information demonstrating that their EHR technology is certified through the Office of the National Coordinator (ONC). The ONC Certified HIT Product List (CHPL) contains the list of all certified EHR technology products and is used by the providers to generate the unique EHR Certification ID that contains the list of all certified EHR technology products and is used by the providers to generate the unique EHR Certification ID that represents the system or combination of modules that is capable of meeting Meaningful Use. The State is required to validate the verification of the Certified EHR information before making any payment to providers.

It is the provider’s responsibility to generate an EHR Certification ID that accurately reflects the complete EHR or combination of modules representing a complete EHR used by the provider before attesting to the State. Failure to do so could result in a false negative result that may disqualify the provider from receiving payment.

To proceed, please indicate your understanding of this responsibility by agreeing to the following statements. Note: the second statement is not required.

**Provider Understands Responsibility:**

- Eligible Professionals must attest that they engaged in SPEP activities by attesting that they: (1) acknowledge the option to cooperate in good faith with ONC’s direct review of their health information technology certified under the ONC Health IT Certification Program if a request to assist in ONC direct review is received; and (2) if a request is received, cooperate in good faith in ONC direct review of health information technology certified under the ONC Health IT Certification Program as authorized by 45 CFR part 170, subpart E, to the extent that such technology meets (or can be used to meet) the definition of CEHRT, including by permitting timely access to such technology and demonstrating its capabilities as implemented and used by the EP in the field.

- Optionally, EPs may also attest that they engaged in SPEP activities by attesting: (1) acknowledge the option to cooperate in good faith with ONC’s direct review of their health information technology certified under the ONC Health IT Certification Program if a request to assist in ONC’s direct review is received, and (2) if a request is received, cooperate in good faith in ONC direct review of health information technology certified under the ONC Health IT Certification Program, as authorized by 45 CFR part 170, subpart E, to the extent that such technology meets (or can be used to meet) the definition of CEHRT, including by permitting timely access to such technology and demonstrating its capabilities as implemented and used by the EP in the field.

---

### EHR Certification Number

2. Search for your product and select “Add to” to add the CMS EHR Certification ID widget on the right side of the page.
3. Once you have entered all of the desired products, click the “Get EHR Certification ID button”.
4. Your CMS EHR Certification ID will be displayed on the screen. This is the number you will need to enter above as part of your attestation.

**NOTE:** ONC does not allow you to mix inpatient and Ambulatory products together to represent a complete EHR solution. Additionally, if the product you add to your shopping cart did not represent a complete EHR solution (as per choosing meaning use criteria, you will not be able to click “Get CMS EHR Certification ID” in step 3.

You must enter an EHR Certification ID that meets the 2014 certification criteria or 2015 certification criteria.

Systems certified to the 2011 criteria no longer qualify toward meeting Meaningful Use.
EHR Reporting Period

CMS requires that providers meet the following regulations for attesting to Meaningful Use:

- 60% of patients must have records in the certified EHR technology

\[
\text{Numerator} = \text{number of patients with records in the certified EHR technology during this reporting period} \\
\text{Denominator} = \text{total number of patients during this reporting period}
\]

- Eligible Professionals who work at multiple locations but don’t have certified EHR technology available at all of their locations must:
  - Have 50% of their total patient encounters at locations where certified EHR technology is available
  - Base all meaningful use measures only on encounters that occurred at locations where certified EHR technology is available

---

EHR Reporting Period

<table>
<thead>
<tr>
<th>Street Address</th>
<th>City</th>
<th>State</th>
<th>Zip</th>
<th>Numerator</th>
<th>Denominator</th>
<th>EHR Technology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1303 SOUTH 28TH AVENUE</td>
<td>HATTIESBURG</td>
<td>MS</td>
<td>39401</td>
<td>1211</td>
<td>1211</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Percentage of total patient encounters at locations where certified EHR technology is available: 100.00%

Numerator = Number of patient encounters in the denominator at the specified location during this reporting period

Denominator = Number of patient encounters at the specified location during this reporting period
Meaningful Use

Please select which Stage of MU you will report for 2017. You must attest to Stage 2 objectives if you are a new participant in 2017. You must have 2015 edition certified EHR technology if you attest to Stage 3 Objectives. You may not change your MU Stage selection option on individual MU objectives pages.

- [ ] I will report Stage 2 objectives in 2017
- [ ] I will report Stage 3 objectives in 2017 (you must have 2015 edition CEHRT to attest to Stage 3 in 2017)

Objectives
Select the Save and Continue button to open each Objective Data page in turn to complete the information for Meaningful Use attestation. Alternatively, select any of the links below to complete that Objective’s Detail page. All objectives must be answered.

Import Meaningful Use Objective Data

**Stage 3 Objective**

<table>
<thead>
<tr>
<th>Objective</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protect Patient Health Information</td>
<td></td>
</tr>
<tr>
<td>Electronic Prescribing (eRx)</td>
<td></td>
</tr>
<tr>
<td>Clinical Decision Support</td>
<td></td>
</tr>
<tr>
<td>Computerized Provider Order Entry (CPOE)</td>
<td></td>
</tr>
<tr>
<td>Patient Electronic Access</td>
<td></td>
</tr>
<tr>
<td>Coordination of Care</td>
<td></td>
</tr>
<tr>
<td>Health Information Exchange</td>
<td></td>
</tr>
<tr>
<td>Public Health and Clinical Data Registry Reporting</td>
<td></td>
</tr>
</tbody>
</table>

To be added configurable in 4.1
**Protect Patient Health Information**

- **Objective:** Protect electronic protected health information (ePHI) created or maintained by the CEHRT through the implementation of appropriate technical, administrative, and physical safeguards.

- **Measure:** Conduct or review a security risk analysis in accordance with the requirements in 45 CFR 164.308(a)(1), including addressing the security (to include encryption) of data created or maintained by CEHRT in accordance with requirements under 45 CFR 164.312(a)(1)(iv) and 45 CFR 164.308(b)(5), and implement security updates as necessary and correct identified security deficiencies as part of the EHR’s risk management process.

**Complete the following information:**

1. Have you conducted or reviewed a security risk analysis in accordance with the requirements?
   - [ ] No
   - [ ] Yes

2. Date security risk analysis was completed:
   
   **Will display with calendar selector**

---

**Attach Files**

- The following attachment is required:
  - Security Risk Analysis – Summary Report

<table>
<thead>
<tr>
<th>File Name</th>
<th>Subject</th>
</tr>
</thead>
</table>

---

**Electronic Prescribing**

- **Objective:** Generate and transmit permissible prescriptions electronically (eRx).

- **Exclusion Criteria:**
  - Meeting either of the following criteria qualifies for the exclusion for this measure.
    - Did you write fewer than 100 prescriptions during the EHR Reporting Period?  
      - [ ] No
      - [ ] Yes
    - Do you have a pharmacy within your organization or one that accepts electronic prescriptions within 10 miles of your practice location at the start of his or her EHR reporting period?  
      - [ ] No
      - [ ] Yes

- **Measure:** More than 60% of all permissible prescriptions written by the EP are queried for a drug formulary and transmitted electronically using CEHRT.

**Complete the following information:**

<table>
<thead>
<tr>
<th>Numerator</th>
<th>Denominator</th>
</tr>
</thead>
</table>

- Numerator = The number of prescriptions in the denominator generated, queried for a drug formulary, and transmitted electronically using CEHRT.
- Denominator = Number of prescriptions written for drugs requiring a prescription in order to be dispensed other than controlled substances during the EHR reporting period, or number of prescriptions written for drugs requiring a prescription in order to be dispensed during the EHR reporting period.
Attach Files

The following attachments are optional:

- Other Attachment

File Name | Subject | Remove
---|---|---

No records to display.

[Add Files] + [Remove Selected]

Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.

[Previous Screen] [Save & Continue]
Clinical Decision Support

Objective: Implement clinical decision support (CDS) interventions focused on improving performance on high-priority health conditions.

Exclusion Criteria: Did you write fewer than 100 of the following orders during the EHR reporting period? Writing fewer than 100 orders qualifies for the exclusion for Measure #0 only.

Measure #1: Implement five clinical decision support interventions related to four or more clinical quality measures at a relevant point in patient care for the entire EHR reporting period. Absent four clinical quality measures related to the EP's scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions.

Complete the following information:

- Have you implemented five clinical decision support interventions related to four or more clinical quality measures or high-priority health conditions at a relevant point in patient care for the entire EHR reporting period?
  - No
  - Yes

List the five clinical decision support interventions you have implemented:

1. 
2. 
3. 
4. 
5. 

These clinical decision support interventions are related to:

- 4 or more clinical quality measures  
- 4 or more high priority health conditions

A cross reference listing of the Clinical Quality Measures is located in the User Guide to assist you with identifying the applicable CGM numbers, if needed.

Measure #2: The EP has enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period.

Complete the following information:

- Have you enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period?
  - No
  - Yes

Attachment Files

The following attachments are optional:

- Other Attachment

File Name  Subject  Remove

No records to display.

Add Files  Remove Selected
### Computerized Provider Order Entry (CPOE)

**Objective:** Use computerized provider order entry (CPOE) for medication, laboratory, and diagnostic imaging orders directly entered by any licensed healthcare professional, credentialed medical assistant, or a medical staff member credentialed to and performing the equivalent duties of a credentialed medical assistant, who can enter orders into the medical record per state, local, and professional guidelines.

**Exclusion Criteria:** Did you write fewer than 100 of the following orders during the EHR reporting period? Writing fewer than 100 orders qualifies for the exclusion for the associated measure.

- Medication Orders (Measure #1) - Yes/No
- Laboratory Orders (Measure #2) - Yes/No
- Diagnostic Imaging Orders (Measure #3) - Yes/No

**Measure #1:** More than 60% of medication orders created by the EP during the EHR reporting period are recorded using CPOE.

Complete the following information:
- Numerator = The number of orders in the denominator recorded using CPOE.
- Denominator = Number of medication orders created by the EP during the EHR reporting period.

**Measure #2:** More than 60% of laboratory orders created by the EP during the EHR reporting period are recorded using CPOE.

Complete the following information:
- Numerator = The number of orders in the denominator recorded using CPOE.
- Denominator = Number of laboratory orders created by the EP during the EHR reporting period.

**Measure #3:** More than 60% of diagnostic imaging orders created by the EP during the EHR reporting period are recorded using CPOE.

Complete the following information:
- Numerator = The number of orders in the denominator recorded using CPOE.
- Denominator = Number of diagnostic imaging orders created by the EP during the EHR reporting period.

### Attach Files

- EHR – Meaningful Use Summary Report (All Objectives)

No records to display.

- Add Files
- Remove Selected

Please select the ‘Previous Screen’ button to go back or the ‘Save & Continue’ button to proceed.
**Patient Electronic Access To Health Information**

**Objective:** Provide patients with timely access to their health information and patient-specific education

**Exclusion Criteria:**
- Did you have any office visits during the EHR reporting period? [ ] No  [ ] Yes
- Did you conduct 50% or more of your encounters in a county/area that does not have 50% or more of its housing units with 50Mbs broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? [ ] No  [ ] Yes

---

**Measure #1:**
More than 80% of all unique patients seen by the EP during the EHR reporting period are 1) provided timely access to view, online, download, and transmit his or her health information; and 2) the EP ensures the patient’s health information is available for the patient (or patient-authorized representative) to access using any application of their choice that is configured to meet the technical specifications of the API in the provider’s CEHR.

Complete the following information:

- **Numerator** = The number of patients in the denominator (or patient authorized representative) who are provided timely access to health information to view online, download, and transmit to a third party and to access using an application of their choice that is configured to meet the technical specifications of the API in the provider’s CEHR.

- **Denominator** = Number of unique patients seen by the EP during the EHR reporting period.

---

**Measure #2:**
The provider must use clinically relevant information from CEHR to identify patient-specific educational resources and provide electronic access to those materials to more than 35% of unique patients seen by the EP during the EHR reporting period.

Complete the following information:

- **Numerator** = The number of patients in the denominator who were provided electronic access to patient-specific educational resources using clinically relevant information identified from CEHR during the EHR reporting period.

- **Denominator** = The number of unique patients seen by the EP during the EHR reporting period.

---

**Attach Files**
The following attachments are optional

- [ ] Other Attachment

<table>
<thead>
<tr>
<th>File Name</th>
<th>Subject</th>
<th>Remove</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
</tbody>
</table>

No records to display.

[Add Files]  [Remove Selected]

Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.

[Previous Screen]  [Save & Continue]
Coordination of Care Through Patient Engagement

Objective: Use CEHRT to engage with patients or their authorized representatives about the patient’s care.

Exclusion Criteria: Meeting either of the following criteria qualifies for the exclusion for the measures:

- Did you have any office visits during the EHR reporting period? [ ] No [ ] Yes
- Did you conduct 50% or more of your encounters in a county/area that does not have 50% or more of its housing units with 4Gbps broadband availability, according to the latest information available from the FCC, on the first day of the EHR reporting period? [ ] No [ ] Yes

Measure #1: During the EHR reporting period, more than 5 percent of all unique patients (or their authorized representatives) seen by the EP actively engage with the electronic health record made accessible by the provider and either: (1) View, download or transmit to a third party their health information; or (2) access their health information through the use of an API that can be used by applications chosen by the patient and configured to the API in the provider’s CEHRT; or (3) a combination of (1) and (2)

Complete the following information:

**Numerator:** The number of patients (or patient authorized representative) in the denominator who have viewed online, downloaded, or transmitted to a third party the patient's health information during the EHR reporting period and the number of unique patients (or patient authorized representative) in the denominator who have accessed their health information through the use of an API during the EHR reporting period.

**Denominator:** Number of unique patients seen by the EP during the EHR reporting period.

Measure #2: For more than 5 percent of all unique patients seen by the EP during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient authorized representative), or in response to a secure message sent by the patient or their authorized representative.

Complete the following information:

**Numerator:** The number of patients in the denominator for whom a secure electronic message is sent to the patient (or patient-authorized representative) or in response to a secure message sent by the patient (or patient-authorized representative), during the EHR reporting period.

**Denominator:** The number of unique patients seen by the EP during the EHR reporting period.

Measure #3: Patient generated health data or data from non-clinical setting is incorporated into the CEHRT for more than 5 percent of all unique patients seen by the EP during the EHR reporting period.

Complete the following information:

**Numerator:** The number of patients in the denominator for whom data from non-clinical settings, which may include patient-generated data, is captured through the CEHRT into the patient record during the EHR reporting period.

**Denominator:** The number of unique patients seen by the EP during the EHR reporting period.
Health Information Exchange

1. About You
2. Confirm Medicaid Eligibility
   - Objective: The EP provides a summary of care or record when transitioning or referring their patient to another setting of care; receives or retrieves a summary of care record upon receipt of a transition or referral or upon the first patient encounter with a new patient, and incorporates summary of care information from other providers into their EHR using the functions of CHERT.
   - Exclusion Criteria: Meeting the following criteria qualifies for the exclusion for any of the measures.

   - Did you transfer a patient to another setting or refer a patient to another provider less than 100 times during the EHR reporting period? (Measure #1)
     - No  Yes

   - Did you conduct 50% or more of your encounters in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #2)
     - No  Yes

   - Were transitions or referrals received and patient encounters in which the provider has never before encountered the patient fewer than 100 during the EHR reporting period? (Measure #3)
     - No  Yes

Measure #1: For more than 50 percent of transitions of care and referrals, the EP that transitions or refers their patient to another setting of care or provider of care: (1) Creates a summary of care record using CHERT, and (2) electronically exchanges the summary of care record.

   Complete the following information:

   - Numerator = The number of transitions of care and referrals in the denominator where a summary of care record was created using Certified EHR technology and was exchanged electronically.

   - Denominator = Number of transitions of care and referrals during the EHR reporting period for which the EP was the transferring or referring provider.

Measure #2: For more than 40 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the EP incorporates the patient’s EHR an electronic summary of care document.

   Complete the following information:

   - Numerator = Number of patient encounters in the denominator where an electronic summary of care record received is incorporated by the provider into the certified EHR technology.

   - Denominator = Number of patient encounters during the EHR reporting period for which an EP was the receiving party of a transition or referral or has never before encountered the patient and for which an electronic summary of care record is available.

Measure #3: For more than 80 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the EP performs a clinical information reconciliation. The provider must implement clinical information reconciliation for the following three clinical information sets: (1) Medication. Review of the patient’s medication, including the name, dosage, frequency, and route of each medication. (2) Medication allergy. Review of the patient’s known medication allergies. (3) Current Problem list. Review of the patient’s current and active diagnoses.

   Complete the following information:

   - Numerator = The number of transitions of care or referrals in the denominator where the following three clinical information reconciliations were performed: Medication list, medication allergy list, and current problem list.

   - Denominator = Number of transitions of care or referrals during the EHR reporting period for which the EP was the recipient of a transition or referral or has never before encountered the patient.
Public Health Reporting

Objective: The EP is in active engagement with a Public Health Agency (PHA) or clinical data registry (CDR) to submit electronic public health data in a meaningful way using certified EHR technology, except where prohibited, and in accordance with applicable law and practice.

In order to meet this objective, EPs would need to meet two of the total number of measures available to them. Reporting an exclusion for a measure does not qualify towards meeting the objective unless the EP can report on fewer than 2 measures. If an EP can report on fewer than 2 measures, the EP must report on any possible measures and claim the applicable exclusions for the remaining measures. If no measures remain available, the EP can meet the objective by claiming applicable exclusions for all measures.

For Measure 4, EPs may choose to report to more than one public health registry to meet the number of measures required to meet the objective. For Measure 5, EPs may choose to report to more than one clinical data registry to meet the number of measures required to meet the objective.

Select "I will report on this measure" to report for the specific measure. Select "I will claim exclusion for this measure" to claim exclusion for the specific measure.

Active Engagement: If your state has one public health agency that manages registration for all the public health measures, you may use this option to select the level of active engagement to apply to all measures with reporting to a Public Health Agency. Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the EP registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days of the start of the EHR reporting period; and the EP is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- Option 2 - Testing and Validation: the EP is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA, or where applicable, the clinical data registry within 30 days. Failure to respond within the EHR reporting period would result in the provider not meeting the measure.

- Option 3 - Production: the EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

<table>
<thead>
<tr>
<th>Measure</th>
<th>I will report on this measure</th>
<th>I will claim exclusion for this measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure 1 - Immunization Registry Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 2 - Syndromic Surveillance Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 3 - Electronic Case Reporting (not required until 2018)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 4 - Public Health Registry Reporting (Registry #1)</td>
<td>☐</td>
<td>☐</td>
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<tr>
<td>Measure 4 - Public Health Registry Reporting (Registry #2)</td>
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<tr>
<td>Measure 4 - Public Health Registry Reporting (Registry #3)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 5 - Clinical Data Registry Reporting (Registry #1)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 5 - Clinical Data Registry Reporting (Registry #2)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 5 - Clinical Data Registry Reporting (Registry #3)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 8 - Electronic Reportable Lab Results Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Please select the 'Previous Screen' button to go back or the 'Save & Continue' button to proceed.
Measure 1 - Immunization Registry Reporting

Measure: The EP is in active engagement with a Public Health Agency (PHA) to submit immunization data and receive immunization forecasts and histories from the public health immunization registry/immunization information system.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply:

- Does not administer any immunizations to any of the populations for which data is collected by its jurisdiction's immunization registry or immunization information system during the EHR reporting period.
- Operates in a jurisdiction for which no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEMIRM definition at the start of the EHR reporting period.
- Operates in a jurisdiction where no immunization registry or immunization information has declared readiness to receive immunization data as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

Option 1 - Completed Registration to Submit Data: the EP registered to submit data with the PHAs to which the information is being submitted; registration was completed within 90 days after the start of the EHR reporting period; and the EP is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

Option 2 - Testing and Validation: the EP is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

Option 3 - Production: the EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

Attach Files

The following attachment is required:

- Evidence of Level of Engagement (Registration, Testing, Production)

File Name | Subject
--- | ---
No records to display.

Add Files | Remove Selected
Measure 2 - Syndromic Surveillance Reporting

■ Red asterisk indicates a required field.

Measure: The EP is in active engagement with a Public Health Agency (PHA) to submit syndromic surveillance data from an urgent care setting.

Exclusion Criteria: Meeting one or more of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Is not in a category of providers from which ambulatory syndromic surveillance data is collected by their jurisdiction’s syndromic surveillance system;
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic syndromic surveillance data from EPs in the specific standards required to meet the CCHIT definition at the start of the EHR reporting period;
- Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from EPs as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data- the EP registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted, registration was completed within 60 days after the start of the EHR reporting period, and the EP is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- Option 2 - Testing and Validation: the EP is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA, or where applicable, the clinical data registry within 30 days, failure to respond within an EHR reporting period would result in the provider not meeting the measure.

- Option 3 - Production: the EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

Attach Files

The following attachments are optional:

- Other Attachment

No records to display.

Add files + Remove Selected 

Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.

Previous Screen Save & Continue

Measure 4 - Public Health Registry Reporting

■ Red asterisk indicates a required field.

Measure: The EP is in active engagement with a Public Health Agency (PHA) to submit data to public health registries.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Does not diagnose or treat any disease or condition associated with a public health registry in their jurisdiction during the EHR reporting period;
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic registry transactions in the specific standards required to meet the CCHIT definition at the start of the EHR reporting period;
- Operates in a jurisdiction where no specialized registry for which the EP has declared readiness to receive electronic registry transactions as of 6 months prior to the start of the EHR reporting period.

Appendix K: Meaningful Use Screenshots
Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1: Completed Registration to Submit Data: The EP registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted. Registration was completed within 60 days after the start of the EHR reporting period, and the EP is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- Option 2: Testing and Validation: The EP is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or where applicable, the clinical data registry within 30 days. Failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

- Option 3: Production: The EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

Registry Name: (please enter name of registry)

Attach Files

The following attachments are optional:
- Other Attachment

File Name | Subject | Remove
---|---|---
No records to display.

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Measure 5 - Clinical Data Registry Reporting

- About You
- Confirm Medicaid Eligibility
- Alteration of EHR
  - EHR Certification
  - EHR Reporting Period
  - MU - Import
- MU Objectives
  - Protect Health
  - CDSS
  - CPOE
  - CQM - Import
  - CQM

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Does not diagnose or directly treat any disease or condition associated with a clinical data registry in their jurisdiction during the EHR reporting period.
- Operates in a jurisdiction for which no clinical data registry is capable of accepting electronic registry transactions in the specific standards required to meet the CERBT definition at the start of the EHR reporting period.
- Operates in a jurisdiction where no clinical data registry for which the EP is eligible has declared readiness to receive electronic registry transactions as of 6 months prior to the start of the EHR reporting period.
Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1. Completed Registration to Submit Data: the EP registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the EP is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- Option 2. Testing and Validation: the EP is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or where applicable, the clinical data registry within 30 days; failure to respond within an EHR reporting period would result in the provider not meeting the measure.

- Option 3. Production: the EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

Registry Name: [please enter name of registry]
Clinical Quality Measure Selection Screen: (no changes for PY 2015-2017 and for Stage 3)

Clinical Quality Measures
EPs must report on a total of nine (9) Clinical Quality Measures that cover at least three (3) of the National Quality Strategy domains. EPs should select the CQMs that best apply to their scope of practice and/or unique patient population. If the EP’s CEHRT does not contain patient data for at least 5 CQMs covering at least 3 domains, then the EP must report the CQMs for which there is patient data and report the remaining required CQMs as “zero denominators” as displayed by the EP’s CEHRT.

CMS has recommended core sets of clinical quality measures that focus on high priority clinical conditions for Eligible Professionals for both Adult and Pediatric measures. To select one of the recommended core measure sets, please select the appropriate option below. To select measures individually, check the Select check box for the measure. If you select measures individually, you must ensure you select 9 measures that cover 3 National Quality Strategy domains.

Placehoder for client configurable text

- I wish to report on the Adult Recommended Core Measures
- I wish to report on the Pediatric Recommended Core Measures
- I wish to select 9 Measures from the list

Import Clinical Quality Measure Data

Efficient Use of Healthcare Resources Domain

<table>
<thead>
<tr>
<th>CMS eMeasure ID</th>
<th>Title</th>
<th>Description</th>
<th>Domain</th>
<th>Select</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS146</td>
<td>Appropriate Testing for Children with Pharyngitis</td>
<td>Percentage of children 2-18 years of age who were diagnosed with pharyngitis, ordered an antibiotic, and received a group A Streptococcus (Strep) test for the episode.</td>
<td>Efficient Use of Healthcare Resources</td>
<td></td>
</tr>
<tr>
<td>CMS168</td>
<td>Use of Imaging Studies for Low Back Pain</td>
<td>Percentage of patients 18-50 years of age with a diagnosis of low back pain who did not have an imaging study (plain X-ray, MRI, CT scan) within 28 days of the diagnosis.</td>
<td>Efficient Use of Healthcare Resources</td>
<td></td>
</tr>
<tr>
<td>CMS154</td>
<td>Appropriate Treatment for Children with Upper Respiratory Infection (URI)</td>
<td>Percentage of children 3 months - 18 years of age who were diagnosed with upper respiratory infection (URI) and were not dispensed an antibiotic prescription on or three days after the episode.</td>
<td>Efficient Use of Healthcare Resources</td>
<td></td>
</tr>
<tr>
<td>CMS129</td>
<td>Prostate Cancer: Avoidance of Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients</td>
<td>Percentage of patients, regardless of age, with a diagnosis of prostate cancer at low risk of recurrence receiving interstitial prostate brachytherapy, OR external beam radiotherapy to the prostate, OR radiotherapy to the groin lymph nodes. OR cryotherapy who did not have a bone scan performed at any time since diagnosis of prostate cancer.</td>
<td>Efficient Use of Healthcare Resources</td>
<td></td>
</tr>
<tr>
<td>CMS Measure ID</td>
<td>Title</td>
<td>Description</td>
<td>Domain</td>
<td>Select</td>
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</tr>
<tr>
<td>CMS137</td>
<td>Initiation and Engagement of Alcohol and Other Drug Dependence Treatment</td>
<td>Percentage of patients 13 years of age and older with a new code of alcohol and other drug (AOD) dependence who received the following. Two rates are reported: a. Percentage of patients who initiated treatment within 14 days of the diagnosis. b. Percentage of patients who initiated treatment and who had two or more additional services with an AOD diagnosis within 30 days of the initiation visit.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS165</td>
<td>Controlling High Blood Pressure</td>
<td>Percentage of patients 18-65 years of age who had a diagnosis of hypertension and whose blood pressure was adequately controlled (≤140/90 mmHg) during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS125</td>
<td>Breast Cancer Screening</td>
<td>Percentage of women 40-69 years of age who had a mammogram to screen for breast cancer.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS124</td>
<td>Cervical Cancer Screening</td>
<td>Percentage of women 21-64 years of age, who received one or more Pap tests to screen for cervical cancer.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS130</td>
<td>Colorectal Cancer Screening</td>
<td>Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS126</td>
<td>Use of Appropriate Medications for Asthma</td>
<td>Percentage of patients 5-64 years of age who were identified as having persistent asthma and were appropriately prescribed medication during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS127</td>
<td>Pneumonia Vaccination Status for Older Adults</td>
<td>Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS131</td>
<td>Diabetes: Eye Exam</td>
<td>Percentage of patients 18-75 years of age with diabetes who had a retinal or dilated eye exam by an eye care professional during the measurement period or a negative retinal exam (no evidence of retinopathy) in the 12 months prior to the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS120</td>
<td>Diabetes: Foot Exam</td>
<td>Percentage of patient aged 18-75 years of diabetes who had a foot exam during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS122</td>
<td>Diabetes: Hemoglobin A1c Poor Control</td>
<td>Percentage of patients 18-75 years of age with diabetes who had hemoglobin A1c ≥ 9.0% during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS148</td>
<td>Hemoglobin A1c Test for Pediatric Patients</td>
<td>Percentage of patients 5-17 years of age with diabetes who had an HbA1c test during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS134</td>
<td>Diabetes: Urine Protein Screening</td>
<td>The percentage of patients 18-75 years of age with diabetes who had a nephropathy screening test or evidence of nephropathy during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS163</td>
<td>Diabetes, Low Density Lipoprotein (LDL) Management</td>
<td>Percentage of patients 18-75 years of age with diabetes whose LDL-C was adequately controlled (&lt;100 mg/dL) during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>CMS164</td>
<td>Ischemic Vascular Disease (IVD): Use of Aspirin or Another Antithrombotic</td>
<td>Percentage of patients 16 years of age and older who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) in the 12 months prior to the measurement period, or who had an active diagnosis of ischemic vascular disease (IVD) during the measurement period, who had documentation of use of aspirin or another antithrombotic during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS165</td>
<td>Coronary Artery Disease (CAD): Beta-Blocker Therapy Prior Myocardial Infarction (AMI) or Left Ventricular Systolic Dysfunction (LVSD &lt;40%)</td>
<td>Percentage of patients 18 years of age and older with a diagnosis of coronary artery disease seen within a 12 month period who also have a prior MI or a current or prior LVEF &lt;40% who were prescribed beta-blocker therapy.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS162</td>
<td>Ischemic Vascular Disease (IVD): Complete Lipid Panel and LDL Control</td>
<td>Percentage of patients 16 years of age and older who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) in the 12 months prior to the measurement period, or who had an active diagnosis of ischemic vascular disease (IVD) during the measurement period, and who had a complete lipid profile performed during the measurement period and whose LDL-C was adequately controlled (&lt;100 mg/dL).</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS135</td>
<td>Heart Failure (HF): Angiotensin-Converting Enzyme (ACE) Inhibitor or Angiotensin Receptor Blocker (ARB) Therapy for Left Ventricular Systolic Dysfunction (LVSD)</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of heart failure (HF) with a current or prior left ventricular ejection fraction (LVEF) &lt;40% who were prescribed ACE inhibitor or ARB therapy either within a 12 month period or within 30 days of the outpatient setting OR at each hospital discharge.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS144</td>
<td>Heart Failure (HF): Beta-Blocker Therapy for Left Ventricular Systolic Dysfunction (LVSD)</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of heart failure (HF) with a current or prior left ventricular ejection fraction (LVEF) &lt;40% who were prescribed beta-blocker therapy either within a 12 month period or within 30 days of the outpatient setting OR at each hospital discharge.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS143</td>
<td>Primary Open Angle Glaucoma (POAG): Optic Nerve Evaluation</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of POAG who have an optic nerve head evaluation during one or more office visits within 12 months.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
<tr>
<td>CMS167</td>
<td>Diabetic Retinopathy, Documentation of Presence or Absence of Macular Edema and Level of Severity of Retinopathy</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of diabetic retinopathy who had a dilated macular or fundus exam performed which included documentation of the level of severity of retinopathy and the presence or absence of macular edema during one or more office visits within 12 months.</td>
<td>Clinical Process/Effectiveness</td>
<td></td>
</tr>
</tbody>
</table>

Appendix K: Meaningful Use Screenshots  Page 237
<table>
<thead>
<tr>
<th>CMS</th>
<th>Measure Description</th>
<th>Percentage Criteria</th>
<th>Measure Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS142</td>
<td>Diabetic Retinopathy: Communication with the Physician Managing Ongoing Diabetes Care</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of diabetic retinopathy who had a dilated macular or fundus exam performed with documented communication to the physician who manages the ongoing care of the patient with diabetes mellitus regarding the findings of the macular or fundus exam at least once within 12 months.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS161</td>
<td>Adult Major Depressive Disorder (MDD): Suicide Risk Assessment</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of major depressive disorder (MDD) who had a suicide risk assessment completed during the visit in which a new diagnosis or treatment modality was identified</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS128</td>
<td>Anti-depressant Medication Management</td>
<td>Percentage of patients 18 years of age and older who were diagnosed with major depression and treated with antidepressant medication, and who remained on antidepressant medication treatment. Two rates are reported: a. Percentage of patients who remained on an antidepressant medication for at least 54 days (12 weeks) b. Percentage of patients who remained on an antidepressant medication for at least 180 days (6 months)</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS156</td>
<td>ADHD: Follow-Up Care for Children Prescribed Attention Deficit/Hyperactivity Disorder (ADHD) Medication</td>
<td>Percentage of children 6-12 years of age and newly diagnosed a medication for attention-deficit/hyperactivity disorder (ADHD) who had appropriate follow-up care. Two rates are reported: a. Percentage of children who had one follow-up visit within 210 days and who, in addition to the visit in the Initiation Phase, had at least two additional follow-up visits with a practitioner within 270 days (9 months) after the Initiation Phase ended.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS169</td>
<td>Bipolar Disorder and Major Depression: Appraisal for alcohol or chemical substance use</td>
<td>Percentage of patients with depression or bipolar disorder with evidence of an initial assessment that includes an appraisal for alcohol or chemical substance use.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS141</td>
<td>Colon Cancer: Chemotherapy for AJCC Stage III Colon Cancer Patients</td>
<td>Percentage of patients aged 18 through 80 years with AJCC Stage III colon cancer who are referred for adjuvant chemotherapy, prescribed adjuvant chemotherapy, or have previously received adjuvant chemotherapy during the 12-month reporting period.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS140</td>
<td>Breast Cancer: Hormonal Therapy for Stage I - IIIC Estrogen Receptor (ER)/Progesterone Receptor (PR)/Positive Breast Cancer</td>
<td>Percentage of female patients aged 18 years and older with Stage I or II breast cancer who were prescribed tamoxifen or aromatase inhibitor (AI) during the 12-month reporting period.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS52</td>
<td>HIV/AIDS: Medical Visit</td>
<td>Percentage of patients, regardless of age, with a diagnosis of HIV/AIDS, with at least two medical visits during the measurement year with a minimum of 90 days between each visit.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS55</td>
<td>HIV/AIDS: Pneumocystis jiroveci pneumonia (PCP) Prophylaxis</td>
<td>Percentage of patients aged 6 years and older with a diagnosis of HIV/AIDS who were prescribed pneumocystis jiroveci pneumonia (PCP) prophylaxis.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS Nbr</td>
<td>Measure Description</td>
<td>Details</td>
<td>Domain</td>
</tr>
<tr>
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</tr>
<tr>
<td>CMS77</td>
<td>HIV/AIDS: RNA control for Patients with HIV</td>
<td>Percentage of patients aged 13 years and older with a diagnosis of HIV/AIDS, with at least two visits during the measurement year, with at least 90 days between each visit, whose most recent HIV RNA level is &lt;200 copies/mL.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS133</td>
<td>Cataract: 20/40 or Better Visual Acuity within 90 Days Following Cataract Surgery</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of uncomplicated cataract who had cataract surgery and no significant ocular conditions impacting the visual outcome of surgery and had best-corrected visual acuity of 20/40 or better (distance or near) achieved within 90 days following the cataract surgery.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS159</td>
<td>Pregnant women that had HIV-1 testing</td>
<td>This measure identified pregnant women who had an HIV-1 test during their pregnancy.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS169</td>
<td>Depression Remission at Twelve Months</td>
<td>Adult patients age 18 and older with major depression or dysthymia and an initial PHQ-9 score ≥18 who demonstrate remission at twelve months defined as PHQ-9 score &lt;5. This measure applies to both patients with newly diagnosed and existing depression whose current PHQ-9 score indicates a need for treatment.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS170</td>
<td>Depression Utilization of the PHQ-9 Tool</td>
<td>Adult patients age 18 and older with the diagnosis of major depression or dysthymia who have a PHQ-9 tool administered at least once during a 4-month period in which there was a qualifying visit.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS75</td>
<td>Children who have dental decay or cavities</td>
<td>Percentage of children ages 0-20 who have had dental decay or cavities during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS74</td>
<td>Primary Care Preventive Intervention as Offered by Primary Care Providers, Including Dentists</td>
<td>Percentage of children, age 0-20 years, who received a fluoride varnish application during the measurement period.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS61</td>
<td>Preventive Care and Screening: Cholesterol - Fasting Low Density Lipoprotein (LDL-C) Test Performed</td>
<td>Percentage of patients aged 20 through 79 years whose risk factors have been assessed and a fasting LDL-C test has been performed.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS64</td>
<td>Preventive Care and Screening: Risk-Stratified Cholesterol - Fasting Low Density Lipoprotein (LDL-C)</td>
<td>Percentage of patients aged 20 through 79 years who had a fasting LDL-C test performed and whose risk-stratified fasting LDL-C is at or below the recommended LDL-C goal.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS146</td>
<td>Dementia, Cognitive Assessment</td>
<td>Percentage of patients, regardless of age, with a diagnosis of dementia for whom an assessment of cognition is performed and the results reviewed at least once within a 12 month period.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
<tr>
<td>CMS86</td>
<td>Hypertension: Improvement in blood pressure</td>
<td>Percentage of patients aged 18-65 years of age with a diagnosis of hypertension whose blood pressure improved during the measurement year.</td>
<td>Clinical Process/Effectiveness</td>
</tr>
</tbody>
</table>

**Patient Safety Domain**
### Patient Safety Domain

<table>
<thead>
<tr>
<th>CMS eMeasure ID</th>
<th>Title</th>
<th>Description</th>
<th>Domain</th>
<th>Select</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS156</td>
<td>Use of High-Risk Medications in the Elderly</td>
<td>Percentage of patients 65 years of age and older who were ordered high-risk medications. Two rates are reported: a. Percentage of patients who were ordered at least one high-risk medication. b. Percentage of patients who were ordered at least two different high-risk medications.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMS159</td>
<td>Falls: Screening for Future Fall Risk</td>
<td>Percentage of patients 65 years of age and older who were screened for future fall risk during the measurement period.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMS668</td>
<td>Documentation of Current Medications in the Medical Record</td>
<td>Percentage of visits for patients aged 18 years and older for which the eligible professional attests to documenting a list of current medications to the best of his/her knowledge and ability. This list must include all medications, over-the-counter, herbal and vitamin/herbal/alternative (nutritional) supplements AND must contain the medications’ name, dosage, frequency and route of administration.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMS132</td>
<td>Cataract: Complications within 30 Days Following Cataract Surgery Requiring Additional Surgical Procedures</td>
<td>Percentage of patients aged 18 years and older with a diagnosis of uncomplicated cataract who had cataract surgery and had any of a specified list of surgical procedures in the 30 days following the cataract surgery which would indicate the occurrence of any of the following major complications: retained nuclear fragments, endophthalmitis, dislocated or wrong power IOL, retinal detachment, or wound dehiscence.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMS177</td>
<td>Child and Adolescent Major Depressive Disorder: Suicide Risk Assessment</td>
<td>Percentage of patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder who are assessed for suicide risk.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMS178</td>
<td>ADE Prevention and Monitoring: Warfarin Time in Therapeutic Range</td>
<td>Average percentage of time in which patients aged 18 and older with atrial fibrillation who are on chronic warfarin therapy have International Normalized Ratio (INR) test results within the therapeutic range (1.6-1.9) during the measurement period.</td>
<td>Patient Safety</td>
<td></td>
</tr>
<tr>
<td>CMSMeasure ID</td>
<td>Title</td>
<td>Description</td>
<td>Domain</td>
<td>Select</td>
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</tr>
<tr>
<td>CM5156</td>
<td>Weight Assessment and Counseling for Nutrition and Physical Activity for Children and Adolescents</td>
<td>Percentage of patients 3-17 years of age who had an outpatient visit with a Primary Care Physician (PCP) or Chiropractic/Gynecologist (CM/GYN) and who had evidence of the following during the measurement period. Three rates are reported. * Percentage of patients with height, weight, and body mass index (BMI) percentile documentation * Percentage of patients with counseling for nutrition * Percentage of patients with counseling for physical activity</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM5138</td>
<td>Preventive Care and Screening; Tobacco Use: Screening and Cessation Intervention</td>
<td>Percentage of patients aged 15 years and older who were screened for tobacco use one or more times within 24 months AND who received cessation counseling intervention if identified as a tobacco user.</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM5153</td>
<td>Chlamydia Screening for Women</td>
<td>Percentage of women 16-24 years of age who were identified as sexually active and who had at least one test for chlamydia during the measurement period.</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM5117</td>
<td>Childhood Immunization Status</td>
<td>Percentage of children 2 years of age who had four diphtheria, tetanus and acellular pertussis (DTaP); three polo (IPV); one mumps, measles, and rubella (MMR); three H influenza type B (Hib); three hepatitis B (Hep B); one chicken pox (IZV); four pneumococcal conjugate (PCV); one Hepatitis A (Hep A); two or three rotavirus (RV); and two influenza (flu) vaccines by their second birthday</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM5147</td>
<td>Preventive Care and Screening: Influenza Immunization</td>
<td>Percentage of patients aged 6 months and older seen for a visit between October 1 and March 31 who received an influenza immunization OR who reported previous receipt of an influenza immunization.</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM52</td>
<td>Preventive Care and Screening: Screening for Clinical Depression and Follow-Up Plan</td>
<td>Percentage of patients aged 12 years and older screened for clinical depression on the date of the encounter using an age appropriate standardized depression screening tool AND if positive, a follow-up plan is documented on the date of the positive screen.</td>
<td>Population/Public Health</td>
<td></td>
</tr>
<tr>
<td>CM5689</td>
<td>Preventive Care and Screening: Body Mass Index (BMI) Screening and Follow-Up</td>
<td>Percentage of patients aged 10 and older with a documented BMI during the encounter or during the previous six months. AND when the BMI is outside of normal parameters, a follow-up plan is documented during the encounter or during the previous six months of the encounter. Normal parameters: Age 60 years and older BMI &gt;= 23 and &lt; 30 Age 10-64 years BMI &gt;= 18.5 and &lt;25</td>
<td>Population/Public Health</td>
<td></td>
</tr>
</tbody>
</table>
### Patient and Family Engagement Domain

<table>
<thead>
<tr>
<th>CMS eMeasure ID</th>
<th>Title</th>
<th>Description</th>
<th>Domain</th>
<th>Select</th>
</tr>
</thead>
<tbody>
<tr>
<td>CM5157</td>
<td>Oncology; Medical and Radiation - Pain Intensity Measured</td>
<td>Percentage of patient visits, regardless of patient age, with a diagnosis of cancer currently receiving chemotherapy or radiation therapy in which pain intensity is quantified.</td>
<td>Patient and Family Engagement</td>
<td></td>
</tr>
<tr>
<td>CM566</td>
<td>Functional status assessment for knee replacement</td>
<td>Percentage of patients aged 18 years and older with primary total knee arthroplasty (TKA) who completed baseline and follow-up (patient-reported) functional status assessments.</td>
<td>Patient and Family Engagement</td>
<td></td>
</tr>
<tr>
<td>CM566</td>
<td>Functional status assessment for hip replacement</td>
<td>Percentage of patients aged 18 years and older with primary total hip arthroplasty (THA) who completed baseline and follow-up (patient-reported) functional status assessments.</td>
<td>Patient and Family Engagement</td>
<td></td>
</tr>
<tr>
<td>CM590</td>
<td>Functional status assessment for complex chronic conditions</td>
<td>Percentage of patients aged 65 years and older with heart failure who completed initial and follow-up patient-reported functional status assessments.</td>
<td>Patient and Family Engagement</td>
<td></td>
</tr>
</tbody>
</table>

### Care Coordination Domain

<table>
<thead>
<tr>
<th>CMS eMeasure ID</th>
<th>Title</th>
<th>Description</th>
<th>Domain</th>
<th>Select</th>
</tr>
</thead>
<tbody>
<tr>
<td>CM590</td>
<td>Closing the referral loop: receipt of specialist report</td>
<td>Percentage of patients with referrals, regardless of age, for which the referring provider receives a report from the provider to whom the patient was referred.</td>
<td>Care Coordination</td>
<td></td>
</tr>
</tbody>
</table>

Please select the 'Previous Screen' button to go back or the 'Save & Continue' button to proceed.
## 2.4.3 EH – Stage 3 Screen Shots

### EHR Certification

**EHR Certification**

Providers must provide information demonstrating that their EHR technology is certified through the Office of the National Coordinator (ONC). The ONC Certified HIT Product List (CHPL) contains the list of all certified EHR technology products and is used by the providers to generate the unique EHR Certification ID that contains the list of all certified EHR technology products and is used by the providers to generate the unique EHR Certification ID that represents the system or combination of modules that is capable of meeting Meaningful Use. The State is required to validate the verification of the Certified EHR information before making any payment to providers.

It is the provider's responsibility to generate an EHR Certification ID that accurately reflects the complete EHR or combination of modules representing a complete EHR used by the provider before attesting to the State. Failure to do so could result in a false negative result that may disqualify the provider from receiving payment.

To proceed, please indicate your understanding of this responsibility by agreeing to the following statements. Note: the second statement is not required.

### Provider Understands Responsibility *

- Eligible Hospital or CAH must attest that they engaged in SPCC activities by attesting that they: (1) acknowledge the requirement to cooperate in good faith with ONC direct review of their health information technology certified under the ONC Health IT Certification Program if a request to assist in ONC direct review is received; and (2) if a request is received, cooperate in good faith with ONC direct review of health information technology certified under the ONC Health IT Certification Program as authorized by 45 CFR part 170, subpart E, to the extent that such technology meets (or can be used to meet) the definition of CEHRT, including by permitting timely access to such technology and demonstrating its capabilities as implemented and used by the eligible hospital, or CAH in the field.

- Optionally, the eligible hospital or CAH may also attest that they engaged in SPCC activities by attesting: (1) acknowledge the option to cooperate in good faith with ONC-ACB surveillance of their health information technology certified under the ONC Health IT Certification Program if a request to assist in ONC-ACB surveillance is received; and (2) if requested, cooperate in good faith with ONC-ACB surveillance of their health information technology certified under the ONC Health IT Certification Program, as authorized by 45 CFR part 170, subpart E, to the extent that such technology meets (or can be used to meet) the definition of CEHRT, including by permitting timely access to such technology and demonstrating its capabilities as implemented and used by the eligible hospital, or CAH in the field.

- Understand that it is my responsibility, as the provider, to ensure that my certified EHR technology code is listed on the ONC public web service before submitting my attestation to the State. I understand that failing to ensure my code is listed may result in a false negative result that may disqualify me from receiving payment.

### EHR Certification Number *

- [Box for EHR Certification Number]

  1. Go to the ONC website: https://healthit.gov. penalties/ and select “CertID” to add to the CMS EHR Certification ID widget on the right side of the page. 
  2. Search for your product(s) and select “CertID” to add to the CMS EHR Certification ID widget on the right side of the page.
  3. Once you have entered all of the desired products, click the “Get EHR Certification ID button”.
  4. Your CMS EHR Certification ID will be displayed on the screen. This is the number you will need to enter above as part of your attestation.

**NOTE:** ONC does not allow you to mix reppliant products and Ambulatory products together to represent a complete EHR solution. Additionally, if the products you used to your shopping cart do not represent a complete EHR solution capable of achieving meaningful use criteria, you will not be able to click “Get CMS EHR Certification ID” in step 3.

You must enter an EHR Certification ID that meets the 2014 certification criteria or 2015 certification criteria.

Systems certified to the 2014 criteria no longer qualify toward meeting Meaningful Use.
EHR Reporting Period

CMS requires that providers meet the following regulations for attesting to Meaningful Use:

* 85% of patients must have records in the certified EHR technology

Numerator *  [ ] Denominator *  [ ] Calculate Percentage 190.00%

Numerator = number of patients with records in the certified EHR technology during this reporting period
Denominator = total number of patients during this reporting period

- I agree that I meet the additional CMS regulations for attesting to Meaningful Use. I understand that the State may choose to audit my records to verify that I meet these regulations.
- I agree with the following statements:
  - The information submitted for clinical quality measures (CQMs) was generated as an output from an identified certified EHR technology
  - The information submitted is accurate to the knowledge and belief of the person submitting on behalf of the eligible hospital or CAH
  - The information submitted is accurate and complete for numerators, denominators, exclusions and measures applicable to the eligible hospital or CAH
  - The information submitted includes information on all patients to whom the measure applies

Recording Period: Start Date* Blank End Date* Blank

- I am reporting CQMs for a different reporting period than my meaningful use objectives

Attach Files

The following attachments are optional:
- Other Attachment

<table>
<thead>
<tr>
<th>File Name</th>
<th>Subject</th>
<th>Remove</th>
</tr>
</thead>
<tbody>
<tr>
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<td></td>
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</tr>
</tbody>
</table>

No records to display.

Add File Remove Selected

Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed:

Previous Screen Save & Continue
### Meaningful Use

Please select which Stage of MU you will report for 2017. You must attest to Stage 2 objectives if you are a new participant in 2017. You must have 2015 edition certified EHR technology if you attest to Stage 3 Objectives. You may not change your MU Stage selection option on individual MU objectives pages.

- [ ] I will report Stage 2 objectives in 2017
- [x] I will report Stage 3 objectives in 2017 (you must have 2015 edition CEHRT to attest to Stage 3 in 2017)

#### Objectives

Select the Save and Continue button to open each Objective Detail page in turn to complete the information for Meaningful Use attestation. Alternatively, select any of the links below to complete that Objective’s Detail page. All objectives must be answered.

Import Meaningful Use Objective Data

<table>
<thead>
<tr>
<th>Stage 3 Objective</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protect Patient Health Information</td>
<td></td>
</tr>
<tr>
<td>Electronic Prescribing (eRx)</td>
<td></td>
</tr>
<tr>
<td>Clinical Decision Support</td>
<td></td>
</tr>
<tr>
<td>Computerized Provider Order Entry (CPOE)</td>
<td></td>
</tr>
<tr>
<td>Patient Electronic Access</td>
<td></td>
</tr>
<tr>
<td>Coordination of Care</td>
<td></td>
</tr>
<tr>
<td>Health Information Exchange</td>
<td></td>
</tr>
<tr>
<td>Public Health and Clinical Data Registry Reporting</td>
<td></td>
</tr>
</tbody>
</table>

### Protect Patient Health Information

**Objective:** Protect electronic protected health information (ePHI) created or maintained by the CEHRT through the implementation of appropriate technical, administrative, and physical safeguards.

**Measure:** Conduct or review a security risk analysis in accordance with the requirements in 45 CFR 164.308(a)(1), including addressing the security (to include encryption) of data created or maintained by CEHRT in accordance with requirements under 45 CFR 164.312(a)(2)(iv) and 45 CFR 164.306(d)(3), and implement security updates as necessary and correct identified security deficiencies as part of the EP’s risk management process.

**Complete the following information:**

- [ ] Have you conducted or reviewed a security risk analysis in accordance with the requirements?
  - [ ] No
  - [x] Yes

- [ ] Date security risk analysis was completed:

Will display with calendar selector

### Attach Files

The following attachments are optional:

- [ ] Other Attachment

No records to display.

[Add Files] [Remove Selected]
### Electronic Prescribing

**Objective:** Generate and transmit permissible prescriptions electronically (eRx).

**Exclusion Criteria:** Meeting either of the following criteria qualifies for the exclusion for this measure:
- Does eligible hospital or CAH have an internal pharmacy that can accept electronic prescriptions or any pharmacy that accept electronic prescriptions within 10 miles at the start of their EHR reporting period?
- No
- Yes

**Measure:** More than 25 percent of hospital discharge medication orders for permissible prescriptions (for new and changed prescriptions) are queried for a drug formulary and transmitted electronically using CDS/CPOE.

Complete the following information:

- Numerator: The number of prescriptions in the denominator generated, queried for a drug formulary, and transmitted electronically.
- Denominator: The number of new or changed prescriptions written for drugs requiring a prescription in order to be dispensed other than controlled substances for patients discharged during the EHR reporting period.

### Attach Files

The following attachments are optional:
- Other Attachment

### Clinical Decision Support

**Objective:** Implement clinical decision support (CDS) interventions focused on improving performance on high-priority health conditions.

**Measure #1:** Implement five clinical decision support interventions related to four or more CQMs at a relevant point in patient care for the entire EHR reporting period. Absent four CQMs related to an eligible hospital or CAH’s scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions.

Complete the following information:
- Have you implemented five clinical decision support interventions related to four or more clinical quality measures or high-priority health conditions at a relevant point in patient care for the entire EHR reporting period?
- No
- Yes

List the five clinical decision support interventions you have implemented:

1. 
2. 
3. 
4. 
5. 

---

**Appendix K: Meaningful Use Screenshots**

Page 246
These clinical decision support interventions are related to:

- [ ] 4 or more clinical quality measures
- [ ] 4 or more high priority health conditions

Select CGM:

- [ ] CGM1
- [ ] CGM4

A cross reference listing of the Clinical Quality Measures is located in the User Guide to assist you with identifying the applicable CGM numbers, if needed.

**Measure #2:**

The EHR or CAH has enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period.

Complete the following information:

- [ ] Have you enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire EHR reporting period?
  - [ ] No
  - [ ] Yes

**Computerized Provider Order Entry (CPOE)**

- [ ] This table indicates a required field.

**Objective:**

Use computerized provider order entry (CPOE) for medication, laboratory, and diagnostic imaging orders directly entered by any licensed healthcare professional, credentialed medical assistant, or a medical staff member credentialed to and performing the equivalent duties of a credentialed medical assistant, who can enter orders into the medical record per state, local, and professional guidelines.

**Measure #1:**

More than 60 percent of medication orders created by the authorized providers of the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.

Complete the following information:

- Numerator = The number of orders in the denominator recorded using CPOE.
- Denominator = Number of medication orders created by the authorized providers in the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) during the EHR reporting period.

**Measure #2:**

More than 60 percent of laboratory orders created by the authorized providers of the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period are recorded using computerized provider order entry.

Complete the following information:

- Numerator = The number of orders in the denominator recorded using CPOE.
- Denominator = Number of laboratory orders created by the authorized providers in the eligible hospital's or CAH's inpatient or emergency department (POS 21 or 23) during the EHR reporting period.
Patient Electronic Access To Health Information

Objective: Provide patients with timely access to their health information and patient-specific education.

Exclusion Criteria: Meeting either of the following criteria qualifies for the exclusion for both measures:
- Is the EHR or CAH in a county/area that does not have 60% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period?
- Yes ☑️ No ☐

Measure #1: More than 60% of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) are, 1) provided timely access to view online, download, and transmit his or her health information, and 2) the provider ensures the patient’s health information is available for the patient (or patient-authorized representative) to access using any application of their choice that is configured to meet the technical specifications of the API in the provider’s CEHRT.

Complete the following information:

Numerator = The number of patients in the denominator (or patient authorized representative) who are provided timely access to health information to view online, download, and transmit to a third party and to access using an application of their choice that is configured to meet the technical specifications of the API in the provider’s CEHRT.

Denominator = The number of unique patients discharged from an eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) during the EHR reporting period.
**Measure #1:** The eligible hospital or CAH must use clinically relevant information from CEHRT to identify patient-specific educational resources and provide electronic access to those materials to more than 35% of unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period.

Complete the following information:

- **Numerator:** The number of patients in the denominator who were provided electronic access to patient-specific educational resources using clinically relevant information identified from CEHRT during the EHR reporting period.

- **Denominator:** The number of unique patients discharged from an eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period.

**Attach Files**

The following attachments are optional:

- **Other Attachment**

**File Name** | **Subject**
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No records to display.

- **Add Files**
- **Remove Selected**

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**Coordination of Care Through Patient Engagement**

1. About You
2. Confirm Medicaid Eligibility
3. Attestation of EHR
   - EHR Certification
   - EHR Reporting Period
   - MU - Import
   - MU Objectives
     - Protect Health
     - CDS
     - CPOE
     - CQM - Import
     - CQM

- **Objective:** Use CEHRT to engage with patients or their authorized representatives about the patient’s care.

- **Exclusion Criteria:**
  - Meeting either of the following criteria qualifies for the exclusion for the measures:
    - Is the EHR or CAH in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability according to the latest information available from the FCC on the first day of the EHR reporting period?
    - Yes
    - No

**Measure #1:** During the EHR reporting period, more than 5 percent of all unique patients (or their authorized representatives) discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) actively engage with the electronic health record made accessible by the provider and either: (1) View, download, or transmit to a third party their health information; or (2) access their health information through the use of an API that can be used by applications chosen by the patient and configured to the API in the provider’s CEHRT, or (3) a combination of (1) and (2).

CMS says providers need to attest with meaningful data but only pass threshold for 2 out of 3 measures.

**Measure #2:** For more than 25 percent of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period, a secure message was sent using the electronic messaging function of CEHRT to the patient (or the patient authorized representative), or in response to a secure message sent by the patient or their authorized representative.
Measure 83: Patient generated health data or data from a non-clinical setting is incorporated into the CEMRT for more than 5 percent of all unique patients discharged from the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) during the EHR reporting period.

**Attach Files**

The following attachments are optional:

- Other Attachment

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</thead>
<tbody>
<tr>
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</tbody>
</table>

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Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.

[Previous Screen] [Save & Continue]
### Health Information Exchange

<table>
<thead>
<tr>
<th>Objective: The eligible hospital or CAH provides a summary of care or record when transitioning or referring their patient to another setting of care, receives or retrieves a summary of care record upon receipt of a transition or referral or upon the first patient encounter with a new patient, and incorporates summary of care information from other providers into their EHR using the functions of CEHRT.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exclusion Criteria: Meeting the following criteria qualifies for the exclusion for relevant measures.</td>
</tr>
<tr>
<td>Is the EH or CAH in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability, according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #1)</td>
</tr>
<tr>
<td>Were transitions or referrals received and patient encounters in which the provider has never before encountered the patient fewer than 100 during the EHR reporting period? (Measure #2)</td>
</tr>
<tr>
<td>Is the EH or CAH in a county/area that does not have 50% or more of its housing units with 4Mbps broadband availability, according to the latest information available from the FCC on the first day of the EHR reporting period? (Measure #3)</td>
</tr>
</tbody>
</table>

#### Measure #1: For more than 50 percent of transitions of care and referrals, the eligible hospital or CAH that transitions or refers their patient to another setting of care or provider of care: (1) Creates a summary of care record using CEHRT, and (2) electronically exchanges the summary of care record.

**Complete the following information:**

- **Numerator:** The number of transitions of care and referrals in the denominator where a summary of care record was created using Certified EHR technology and is exchanged electronically.
- **Denominator:** Number of transitions of care and referrals during the EHR reporting period for which the eligible hospital’s or CAH’s inpatient or emergency department (POS 21 or 23) was the transferring or referring provider.

#### Measure #2: For more than 50 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the eligible hospital or CAH incorporates into the patient’s EHR an electronic summary of care document.

**Complete the following information:**

- **Numerator:** Number of patient encounters in the denominator where an electronic summary of care record received is incorporated by the provider into the certified EHR technology.
- **Denominator:** Number of patient encounters during the EHR reporting period for which an eligible hospital or CAH was the receiving party of a transition or referral or has never before encountered the patient and for which an electronic summary of care record is available.

#### Measure #3: For more than 80 percent of transitions or referrals received and patient encounters in which the provider has never before encountered the patient, the eligible hospital or CAH performs a clinical information reconciliation. The provider must implement clinical information reconciliation for the following three clinical information sets: (1) Medication Review of the patient’s medication, including the name, dosage, frequency, and route of each medication. (2) Medication allergy. Review of the patient’s known medication allergies. (3) Current Problem list. Review of the patient’s current and active diagnoses.

**Complete the following information:**

- **Numerator:** The number of transitions of care or referrals in the denominator where the following three clinical information reconciliations were performed: Medication list, medication allergy list, and current problem list.
- **Denominator:** Number of transitions of care or referrals during the EHR reporting period for which the eligible hospital or CAH inpatient or emergency department (POS 21 or 23) was the recipient of the transition or referral or has never before encountered the patient.
Public Health and Clinical Data Registry Reporting

Objective: The eligible hospital or CAH is an active participant in a Public Health Agency (PHA) or clinical data registry (CDR) to submit electronic public health data in a meaningful way using certified EHR technology, except where prohibited, and in accordance with applicable laws and practices.

In order to meet this objective, eligible hospital or CAH would need to meet four of the total number of measures available to them. Reporting an exclusion for a measure does not qualify towards meeting the objective unless the eligible hospital or CAH can report on fewer than four measures. If an EHR or CAH can report on fewer than four measures, the eligible hospital or CAH must report on any possible measures and claim the applicable exclusions for the remaining measures. If no measures remain available, the eligible hospital or CAH can meet the objective by claiming applicable exclusions for all measures.

For Measure 4, eligible hospital or CAH may choose to report more than one public health registry to meet the number of measures required to meet the objective. For Measure 5, eligible hospital or CAH may choose to report more than one clinical data registry to meet the number of measures required to meet the objective.

Select "I will report on this measure" to report for the specific measure. Select "I will claim exclusion for this measure" to claim exclusion for the specific measure.

Active Engagement: If your state has one public health agency that manages registration for all the public health measures, you may use this option to select the level of active engagement to apply to all measures with reporting to a Public Health Agency. Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

Option 1 - Completed Registration to Submit Data: The eligible hospital or CAH registered to submit data with the PHA or, where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period, and the eligible hospital or CAH is enrolled in the PHA to begin testing and validation. This option allows providers to meet the measure when the PHA or the CDR has limited resources to initiate the testing and validation process. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

Option 2 - Testing and Validation: The eligible hospital or CAH is in the process of testing and validation of the CDR and in accordance with data. Providers must respond to requests from the PHA or, where applicable, the clinical data registry within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

Option 3 - Production: The eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

<table>
<thead>
<tr>
<th>Measure</th>
<th>I will report on this measure</th>
<th>I will claim exclusion for this measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure 1 - Immunization Registry Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 2 - Syndromic Surveillance Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 3 - Electronic Case Reporting (not required until 2015)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 4 - Public Health Registry Reporting (Registry #1)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 5 - Public Health Registry Reporting (Registry #2)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 6 - Public Health Registry Reporting (Registry #3)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 7 - Clinical Data Registry Reporting (Registry #1)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 8 - Clinical Data Registry Reporting (Registry #2)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 9 - Clinical Data Registry Reporting (Registry #3)</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Measure 10 - Electronic Reportable Laboratory Data Reporting</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Please select the "Previous Screen" button to go back or the "Save & Continue" button to proceed.
Measure 1 - Immunization Registry Reporting

Measure: The EH or CAH is in active engagement with a Public Health Agency (PHA) to submit immunization data and receive immunization forecasts and histories from the public health immunization registry immunization information system.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- [ ] Does not administer any immunizations to any of the populations for which data is collected by its jurisdiction's immunization registry or immunization information system during the EHR reporting period.
- [ ] Operates in a jurisdiction for which no immunization registry or immunization information system is capable of accepting the specific standards required to meet the CEHRT definition at the start of the EHR reporting period.
- [ ] Operates in a jurisdiction where no immunization registry or immunization information has declared readiness to receive immunization data as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- [ ] Option 1 - Completed Registration to Submit Data: The eligible hospital or CAH registered to submit data with the PHA to which the information is being submitted; registration was completed within 30 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- [ ] Option 2 - Testing and Validation: The eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or, where applicable, the clinical data registry within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

- [ ] Option 3 - Production: The eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA.

Attach Files

The following attachments are optional:

- Other Attachment

File Name | Subject | Remove
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No records to display.

Add File | Remove Selected

Please select the 'Previous screen' button to go back or the 'Save & Continue' button to proceed.

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Measure 2 - Syndromic Surveillance Reporting

Measure: The eligible hospital or CAH is in active engagement with a public health agency to submit syndromic surveillance data from an urgent care setting.

Exclusion Criteria: Meeting one or more of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Does not have an emergency or urgent care department;
- Operates in a jurisdiction in which no public health agency is capable of receiving electronic syndromic surveillance data from eligible hospital or CAH in the specific standards required to meet the CEHRT definition at the start of the EHR reporting period;
- Operates in a jurisdiction where no public health agency has declared readiness to receive syndromic surveillance data from eligible hospital or CAH as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or, where applicable, the clinical data registry within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

- Option 3 - Production: the eligible hospital or CAH is in active engagement with a public health agency to submit electronic reportable laboratory results has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

Attach Files

The following attachments are optional:

- Other Attachment

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No records to display.

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Please select the ‘Previous screen’ button to go back or the ‘Save & Continue’ button to proceed.

Previous Screen | Save & Continue
Measure 4 - Public Health Registry Reporting

Measure: The eligible hospital or CAH is in active engagement with a Public Health Agency (PHA) to submit data to public health registries.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply:

- Does not diagnose or directly treat any disease or condition associated with a public health registry
- Operates in a jurisdiction for which no public health agency is capable of receiving electronic registry transactions in the specific standards required to meet the CCHRT definition at the start of the EHR reporting period.
- Operates in a jurisdiction where no public health registry for which the eligible hospital or CAH is eligible has declared readiness to receive electronic registry transactions as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or where applicable, the clinical data registry within 30 days. Failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.
- Option 3 - Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

Registry Name: [please enter name of registry]

Attach Files

The following attachments are optional:

- Other Attachment

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No records to display.

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Please select the "Previous screen" button to go back or the "Save & Continue" button to proceed.
### Measure 5 - Clinical Data Registry Reporting

**Measure:** The eligible hospital or CAH is in active engagement to submit data to a clinical data registry (CDR).

**Exclusion Criteria:** Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- [ ] Does not diagnose or directly treat any disease or condition associated with a clinical data registry in their jurisdiction during the EHR reporting period.
- [ ] Operates in a jurisdiction for which no clinical data registry is capable of accepting electronic registry transactions in the specific standards required to meet the CCHIT definition at the start of the EHR reporting period.
- [ ] Operates in a jurisdiction where no clinical data registry for which the eligible hospital or CAH is eligible has declared readiness to receive electronic registry transactions as of 6 months prior to the start of the EHR reporting period.

**Active Engagement:** Select the level of active engagement that you demonstrate for this measure. Only one level may be selected.

- **Option 1:** Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHA, or where applicable, the clinical data registry to which the information is being submitted; registration was completed within 60 days after the start of the EHR reporting period; and the eligible hospital or CAH is awaiting an invitation from the PHA to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.

- **Option 2:** Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHA or where applicable, the clinical data registry within 30 days; failure to respond twice within an EHR reporting period would result in the provider not meeting the measure.

- **Option 3:** Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or clinical data registry.

| Registry Name: | [please enter name of registry] |

#### Attach Files

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- **Other Attachment**

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Measure 6 - Electronic Reportable Lab Results Reporting

Measure: The eligible hospital or CAH is in active engagement with a public health agency to submit electronic reportable laboratory results.

Exclusion Criteria: Meeting any of the following criteria qualifies for the exclusion for this measure. Select all that apply.

- Does not perform or order laboratory tests that are reportable in their jurisdiction during the EHR reporting period;
- Operates in a jurisdiction for which no public health agency is capable of accepting the specific ELR standards required to meet the CCHIT definition at the start of the EHR reporting period;
- Operates in a jurisdiction where no public health agency has declared readiness to receive electronic reportable laboratory results from an eligible hospital or CAH as of 6 months prior to the start of the EHR reporting period.

Active Engagement: Select the level of active engagement you demonstrate for this measure. Only one level may be selected.

- Option 1 - Completed Registration to Submit Data: the eligible hospital or CAH registered to submit data with the PHE, or where applicable, the clinical data registry to which the information is being submitted, registration was completed within 60 days after the start of the EHR reporting period, and the eligible hospital or CAH is awaiting an invitation from the PHE to begin testing and validation. Providers that have registered in previous years do not need to submit an additional registration to meet this requirement for each EHR reporting period.
- Option 2 - Testing and Validation: the eligible hospital or CAH is in the process of testing and validation of the electronic submission of data. Providers must respond to requests from the PHE or, where applicable, the clinical data registry within 30 days. Failure to respond twice within an EHR reporting period period would result in the provider not meeting the measure.
- Option 3 - Production: the eligible hospital or CAH has completed testing and validation of the electronic submission and is electronically submitting production data to the PHE or clinical data registry.

Registry Name: [please enter name of registry]

Attach Files

The following attachments are optional:

- Other Attachment

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Subject
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Clinical Quality Measures

EHs and CAHs must report on a total of sixteen (16) Clinical Quality Measures. If an eligible hospital’s or CAH’s CEHRT does not contain patient data for at least 10 CQMs, then the EH or CAH must report the CQMs for which there is patient data and report the remaining required CQMs as “zero denominators” as displayed by the EH’s or CAH’s CEHRT.

Eligible hospitals and CAHs that have 5 or fewer discharges per quarter in the same quarter as their reporting period, or 20 or fewer discharges per full PY reporting period for which data is being electronically submitted as defined by the CQMs denominator population are exempted for reporting the CQMs.

Clinical Quality Measures Summary

<table>
<thead>
<tr>
<th>CMS Measure ID</th>
<th>Title</th>
<th>Description</th>
<th>NQF (not final if this column will be Dropped)</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>CM0553</td>
<td>AMI-1e: Primary PCI Received within 90 Minutes of Hospital Arrival</td>
<td>Acute myocardial infarction (AMI) patients with ST segment elevation on LSEST or on the ECG closest to arrival time receiving primary PCI during the hospital stay with a time from hospital arrival to fibrinolysis of 90 minutes or less.</td>
<td>0.03</td>
<td></td>
</tr>
<tr>
<td>CM0552</td>
<td>ED-2: Median time from ED arrival to ED departure for discharged ED patients</td>
<td>Median time from emergency department arrival to time of discharge from the emergency room for patients discharged from the emergency department.</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>CM0526</td>
<td>CMC-1: Home Management Plan of Care (HMPG) Document Given to Patient/Guardian</td>
<td>An assessment that there is documentation in the medical record that a Home Management Plan of Care (HMPG) document was given to the patient or their legal guardian.</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>CM0556</td>
<td>Emergency Department (ED) - 1 Emergency Department Throughput - Median time from ED arrival to ED departure for admitted ED patients</td>
<td>Median time from emergency department arrival to time of departure from the emergency room for patients admitted to the facility from the emergency department.</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>CM0511</td>
<td>ED-2: Emergency Department Throughput - Median time from ED admission to ED departure for admitted patients</td>
<td>Median time (minutes) from admit decision time to time of departure from the emergency department for emergency department patients admitted to inpatient status.</td>
<td>0.087</td>
<td></td>
</tr>
<tr>
<td>CM551</td>
<td>HMGP-1a: Hearing screening prior to hospital discharge</td>
<td>This measure assesses the proportion of infants that have been screened for hearing loss before hospital discharge.</td>
<td>0.84</td>
<td></td>
</tr>
<tr>
<td>CM5113</td>
<td>PC-01: Vital Sign Measurement Prior to 30 Completed Week 1st</td>
<td>Patients with vital sign measurements completed.</td>
<td>0.409</td>
<td></td>
</tr>
<tr>
<td>CM509</td>
<td>PC-02: Exclusive Breast Milk Feeding</td>
<td>Exclusive breast milk feeding during the newborn’s entire hospitalization.</td>
<td>0.403</td>
<td></td>
</tr>
<tr>
<td>CM05104</td>
<td>Stroke-2: Ischemic stroke - Discharged on antiplatelet therapy</td>
<td>Ischemic stroke patients prescribed antithrombotic therapy at discharge.</td>
<td>0.435</td>
<td></td>
</tr>
<tr>
<td>CM071</td>
<td>Stroke-3: Ischemic stroke - Administration of Therapy for Ischemic Stroke Patients on Admission</td>
<td>Ischemic stroke patients with medical history of ischemic stroke patients who are prescribed antithrombotic therapy at hospital discharge.</td>
<td>0.426</td>
<td></td>
</tr>
<tr>
<td>CM0512</td>
<td>Stroke-4: Ischemic stroke - Administration of Therapy for Ischemic Stroke Patients on Admission</td>
<td>Ischemic stroke patients with medical history of ischemic stroke patients who are prescribed antithrombotic therapy by the end of hospital stay.</td>
<td>0.439</td>
<td></td>
</tr>
<tr>
<td>CM05126</td>
<td>Stroke-5: Ischemic stroke - Administration of Therapy for Ischemic Stroke Patients on Admission</td>
<td>Ischemic stroke patients with medical history of ischemic stroke patients who are prescribed antithrombotic therapy at hospital discharge.</td>
<td>0.436</td>
<td></td>
</tr>
</tbody>
</table>

Appendix K: Meaningful Use Screenshots
| CMS107 | Stroke-IL Ischemic or hemorrhagic stroke - Stroke education | Ischemic or hemorrhagic stroke patients or their caregivers who were given educational materials during hospital stay addressing all of the following: activation of emergency medical system, need for follow-up after discharge, medications prescribed at discharge, risk factors for stroke, and warning signs and symptoms of stroke. |  |
|--------|----------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------|  |
| CMS102 | Stroke-IO Ischemic or hemorrhagic stroke - Assessment for Rehabilitation | Ischemic or hemorrhagic stroke patients who were assessed for rehabilitation services. | 4641 |
| CMS108 | Venous Thromboembolism (VTE) - VTE prophylaxis | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after hospital admission or surgery and date for surgeries that start the day of or the day after hospital admission. | 5371 |
| CMS190 | VTE-2 Intensive Care Unit (ICU) VTE prophylaxis | This measure assesses the number of patients who received VTE prophylaxis or have documentation why no VTE prophylaxis was given the day of or the day after the initial admission (or transfer) to the Intensive Care Unit (ICU) or surgery and date for surgeries that start the day of or the day after ICU admission (or transfer). | 5372 |
Appendix L: DOM Connectivity and Interoperability Strategy

(Retired with 2017 SMHP)

This document defined the DOM Connectivity & Interoperability Strategy from its inception and was updated through calendar year 2016. The IOP strategy is now only of value as an historical document so with the 2017 SMHP and subsequent versions the contents of Appendix L have been removed for efficiency. The Appendix L header page, Table of Contents, Table of Tables and Table of Figures have been retained for anyone considering review of the retired document available in the CMS approved 2016 SMHP.

DOM Connectivity & Interoperability Strategy

As-Is, To-Be and Roadmap Report

2016
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1 Introduction and Overview ......................................................................................... Error! Bookmark not defined.

2 DOM Connectivity and Interoperability Strategy – Assessment of As-Is Environment Error! Bookmark not defined.
   2.1 As-Is DOM Infrastructure ......................................................................................... Error! Bookmark not defined.
      2.1.1 Background ....................................................................................................... Error! Bookmark not defined.
      2.1.2 Connectivity ..................................................................................................... Error! Bookmark not defined.
   2.2 As-Is MMIS, MEDS/X Eligibility Systems, and SLR ........................................ Error! Bookmark not defined.
      2.2.1 Background ....................................................................................................... Error! Bookmark not defined.
      2.2.2 Connectivity ..................................................................................................... Error! Bookmark not defined.
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   2.3 As-Is DOM Clinical Data Infrastructure (CDI) ...................................................... Error! Bookmark not defined.
      2.3.1 Background ....................................................................................................... Error! Bookmark not defined.
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      2.4.1 Background ....................................................................................................... Error! Bookmark not defined.
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   2.6 Connectivity to Federal Agencies .......................................................................... Error! Bookmark not defined.

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   3.1 The State of Mississippi DOM Ecosystem To-Be .................................................. Error! Bookmark not defined.
      3.1.1 High-Level Architecture for DOM Ecosystem .................................................. Error! Bookmark not defined.
      3.1.2 Desired Characteristics of the DOM Ecosystem .............................................. Error! Bookmark not defined.
   3.2 Business and Technical Considerations .................................................................. Error! Bookmark not defined.
      3.2.1 Technical Requirements and Guidance ............................................................ Error! Bookmark not defined.
      3.2.2 Adoption of Various Federal and Industry Standards and Technology Error! Bookmark not defined.
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3.7 To-Be Mississippi State Department of Health Interoperability ... Error! Bookmark not defined.

3.8 To-Be Other State Agency Interoperability .............................. Error! Bookmark not defined.

3.9 To-Be Federal Agency Interoperability and Surrounding State HIE Interoperability .......... Error! Bookmark not defined.

3.10 To-Be for DOM Interoperability Platform with support for NwHIN (HealthWay CONNECT-compliant) as a Connectivity Methodology ............................................. Error! Bookmark not defined.

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4.4 Roadmap for DOM MES, MEDS/X Eligibility Systems, and SLR ...... Error! Bookmark not defined.

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4.4.2 Implementation Path for MEDS/X ........................................ Error! Bookmark not defined.

4.4.3 Implementation Path for SLR .................................................. Error! Bookmark not defined.

4.5 Roadmap for DOM CDI System ................................................ Error! Bookmark not defined.

4.5.1 Implementation Path for DOM CDI System ............................ Error! Bookmark not defined.

4.6 Roadmap for Mississippi Health Information Network (MS-HIN) Interoperability ............ Error! Bookmark not defined.

4.6.1 Implementation Path .............................................................. Error! Bookmark not defined.

4.7 Roadmap for Mississippi State Department of Health Interoperability ...... Error! Bookmark not defined.

4.7.1 Implementation Path ............................................................ Error! Bookmark not defined.

4.8 Roadmap Other State Agency Interoperability ............................. Error! Bookmark not defined.

4.8.1 Implementation Path ............................................................ Error! Bookmark not defined.

4.9 Roadmap for Federal Agency and Surrounding State HIE Interoperability .......... Error! Bookmark not defined.

4.9.1 Implementation Path ............................................................ Error! Bookmark not defined.
4.10 Roadmap for DOM Interoperability Platform support of NwHIN (HealtheWay CONNECT) as a Connectivity Methodology ......................................................... Error! Bookmark not defined.

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**Appendix M: CMS Guidelines Cross-Reference**

The following tables identify the sections of this document where specific SMHP document requirements, primarily the CMS Guidelines, are addressed. An asterisk, “*”, indicates the requirement is considered optional by CMS.

### Cross Reference from CMS Guidelines to Section 3 – Current HIT Landscape Assessment – The “As-Is” Environment:

<table>
<thead>
<tr>
<th>CMS Guidelines Section A: The State’s “As-Is” HIT Landscape</th>
<th>Location in Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. What is the current extent of EHR adoption by practitioners and by hospitals? How recent is this data? Does it provide specificity about the types of EHRs in use by the State’s providers? Is it specific to just Medicaid or an assessment of overall statewide use of EHRs? Does the SMA have data or estimates on eligible providers broken out by types of provider? Does the SMA have data on EHR adoption by types of provider (e.g., children’s hospitals, acute care hospitals, pediatricians, nurse practitioners, etc.)?</td>
<td>Section 3.1</td>
</tr>
<tr>
<td>2. To what extent does broadband internet access pose a challenge to HIT/E in the State’s rural areas? Did the State receive any broadband grants?</td>
<td>Section 3.7</td>
</tr>
<tr>
<td>3. Does the State have Federally-Qualified Health Center networks that have received or are receiving HIT/EHR funding from the Health Resources Services Administration (HRSA)? Please describe.</td>
<td>Section 3.11</td>
</tr>
<tr>
<td>4. Does the State have Veterans Administration or Indian Health Service clinical facilities that are operating EHRs? Please describe.</td>
<td>Section 3.12</td>
</tr>
<tr>
<td>5. What stakeholders are engaged in any existing HIT/E activities and how would the extent of their involvement be characterized?</td>
<td>Section 3.1 and Section 3.9 of SMHP version 1.1</td>
</tr>
<tr>
<td>6. * Does the SMA have HIT/E relationships with other entities? If so, what is the nature (governance, fiscal, geographic scope, etc) of these activities?</td>
<td>Section 3.4. Yes, clinical data interoperability between Medicaid and large health systems for C-CDA exchange in real-time.</td>
</tr>
<tr>
<td>7. Specifically, if there are health information exchange organizations in the State, what is their governance structure and is the SMA involved? ** How extensive is their geographic reach and scope of participation?</td>
<td>Section 3.9, 4.7 Public data about utilization of the HIE is not available.</td>
</tr>
<tr>
<td>8. Please describe the role of the MMIS in the SMA’s current HIT/E environment. Has the State coordinated their HIT Plan with their MITA transition plans and if so, briefly describe how.</td>
<td>Section 3.5</td>
</tr>
<tr>
<td>9. What State activities are currently underway or in the planning phase to facilitate HIE and EHR adoption? What role does the SMA play? Who else is currently involved? For example, how are the regional extension centers (RECs) assisting Medicaid eligible providers to implement EHR systems and achieve meaningful use?</td>
<td>Section 3.5</td>
</tr>
<tr>
<td><strong>CMS Guidelines Section A: The State’s “As-Is” HIT Landscape, continued</strong></td>
<td><strong>Location in Document</strong></td>
</tr>
<tr>
<td>---</td>
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</tr>
<tr>
<td>10. Explain the SMA’s relationship to the State HIT Coordinator and how the activities planned under the ONC-funded HIE cooperative agreement and the Regional Extension Centers (and Local Extension Centers, if applicable) would help support the administration of the EHR Incentive Program.</td>
<td>Section 3.9</td>
</tr>
<tr>
<td>11. What other activities does the SMA currently have underway that will likely influence the direction of the EHR Incentive Program over the next five years?</td>
<td>Section 4.1 We plan to use available data to identify providers that once participated and have dropped out of the EHR Incentive Program through the years. We plan to ramp up education and outreach efforts that will help participants better utilize their new or existing EHRs.</td>
</tr>
<tr>
<td>12. Have there been any recent changes (of a significant degree) to State laws or regulations that might affect the implementation of the EHR Incentive Program? Please describe.</td>
<td>No changes to State Laws that might impact the EHR Incentive Program</td>
</tr>
<tr>
<td>13. Are there any HIT/E activities that cross State borders? Is there significant crossing of State lines for accessing health care services by Medicaid beneficiaries? Please describe.</td>
<td>Section 4.4 No State border initiatives currently. Significant crossing of State lines to areas such as New Orleans and Memphis by Medicaid beneficiaries.</td>
</tr>
<tr>
<td>14. What is the current interoperability status of the State Immunization registry and Public Health Surveillance reporting database(s)?</td>
<td>Section 3.9. Public Health infrastructure is a part of the State HIE, MS-HIN, and accessible by MS-HIN users.</td>
</tr>
<tr>
<td>15. If the State was awarded an HIT-related grant, such as a Transformation Grant or a CHIPRA HIT grant, please include a brief description.</td>
<td>No such award.</td>
</tr>
</tbody>
</table>

*May be deferred  
**The first part of this question may be deferred but States do need to include a description of their HIE(s); geographic reach and current level of participation.
Cross Reference from CMS Guidelines to Section 4 – To-Be:

<table>
<thead>
<tr>
<th>CMS Guidelines Section B: The State’s “To-Be” Landscape</th>
<th>Location in Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Looking forward to the next five years, what specific HIT/E goals and objectives does the SMA expect to achieve? Be as specific as possible; e.g., the percentage of eligible providers adopting and meaningfully using certified EHR technology, the extent of access to HIE, etc.</td>
<td>Section 4.3</td>
</tr>
<tr>
<td>2. *What will the SMA’s IT system architecture (potentially including the MMIS) look like in five years to support achieving the SMA’s long term goals and objectives? Internet portals? Enterprise Service Bus? Master Patient Index? Record Locator Service?</td>
<td>Section 4.2, 4.3</td>
</tr>
<tr>
<td>3. How will Medicaid providers interface with the SMA IT system as it relates to the EHR Incentive Program (registration, reporting of MU data, etc.)?</td>
<td>Section 4.1 and Blue Print in Section 5</td>
</tr>
<tr>
<td>4. Given what is known about HIE governance structures currently in place, what should be in place by 5 years from now in order to achieve the SMA’s HIT/E goals and objectives? While we do not expect the SMA to know the specific organizations will be involved, etc., we would appreciate a discussion of this in the context of what is missing today that would need to be in place five years from now to ensure EHR adoption and meaningful use of EHR</td>
<td>Section 4.7, 4.3</td>
</tr>
<tr>
<td>5. What specific steps is the SMA planning to take in the next 12 months to encourage provider adoption of certified EHR technology?</td>
<td>Section 4.1 We will continue utilizing existing technology and deploy updated SLR releases as required by future CMS regulatory changes.</td>
</tr>
<tr>
<td>6. *If the State has FQHCs with HRSA HIT/EHR funding, how will those resources and experiences be leveraged by the SMA to encourage EHR adoption?</td>
<td>Section 4.9, 4.3</td>
</tr>
<tr>
<td>7. *How will the SMA assess and/or provide technical assistance to Medicaid providers around adoption and meaningful use of certified EHR technology?</td>
<td>Section 4.1 The SMA will no longer work to get new providers into the EHR Provider Incentive Program after the conclusion of Program Year 2016. However, we plan to continue providing resources, education and support for remaining participants, helping them better utilize their existing and new EHR systems to meet Meaningful Use</td>
</tr>
<tr>
<td>8. *How will the SMA assure that populations with unique needs, such as children, are appropriately addressed by the EHR Incentive Program?</td>
<td>Plans to assure that specific populations are appropriately addresses by the EHR Incentive Program have not been designed at this time.</td>
</tr>
<tr>
<td>CMS Guidelines Section B: The State’s “To-Be” Landscape, continued</td>
<td>Location in Document</td>
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</tr>
<tr>
<td>9. If the State included in a description of a HIT-related grant award (or awards) in Section A, to the extent known, how will that grant, or grants, be leveraged for implementing the EHR Incentive Program, e.g., actual grant products, knowledge/lessons learned, stakeholder relationships, governance structures, legal/consent policies and agreements, etc.?</td>
<td>No such award</td>
</tr>
<tr>
<td>10. Does the SMA anticipate the need for new or State legislation or changes to existing State laws in order to implement the EHR Incentive Program and/or facilitate a successful EHR Incentive Program (e.g., State laws that may restrict the exchange of certain kinds of health information)? Please describe.</td>
<td>There is not an expectation for state regulatory changes in the near future that could impact the EHR Incentive Program.</td>
</tr>
<tr>
<td>Please include other issues that the SMA believes need to be addressed, institutions that will need to be present and interoperability arrangements that will need to exist in the next five years to achieve its goals.</td>
<td>Section 4.3</td>
</tr>
</tbody>
</table>

*This question may be deferred if the timing of the submission of the SMHP does not accord with when the long-term vision for the Medicaid IT system is decided. It would be helpful though to note if plans are known to include any of the listed functionalities / business processes.

** May be deferred.
## Cross Reference from CMS Guidelines to Section 5 – Provider Incentive Program Blueprint:

<table>
<thead>
<tr>
<th>CMS Guidelines Section C: Activities Necessary to Administer and Oversee the EHR Incentive Payment Program</th>
<th>Location in Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. How will the SMA verify that providers are not sanctioned, are properly licensed/qualified providers?</td>
<td>Section 5.3.2 Verification or validation of professional licensing uses our Provider Master File which is updated weekly from MMIS data</td>
</tr>
<tr>
<td>2. How will the SMA verify whether EPs are hospital-based or not?</td>
<td>Section 5.2.2.1.3</td>
</tr>
<tr>
<td>3. How will the SMA verify the overall content of provider attestations?</td>
<td>Section 5.4</td>
</tr>
<tr>
<td>4. How will the SMA communicate to its providers regarding their eligibility, payments, etc.?</td>
<td>Section 5.4</td>
</tr>
<tr>
<td>5. What methodology will the SMA use to calculate patient volume?</td>
<td>Section 5.5</td>
</tr>
<tr>
<td>6. (a) What data sources will the SMA use to verify patient volume for EPs and acute care hospitals?</td>
<td>Section 5.5.1.2</td>
</tr>
<tr>
<td>6. (b) How will the SMA verify adopt, implement or upgrade of certified electronic health record technology by providers?</td>
<td></td>
</tr>
<tr>
<td>7. (a) How will the SMA verify that EPs at FQHC/RHCs meet the practices predominately requirement?</td>
<td>Section 5.2.2.1.1</td>
</tr>
<tr>
<td>7. (b) How will the SMA verify meaningful use of certified electronic health record technology for providers’ second participation years?</td>
<td>Section 5.4.2</td>
</tr>
<tr>
<td>8. Will the SMA be proposing any changes to the MU definition as permissible per rule-making? If so, please provide details on the expected benefit to the Medicaid population as well as how the SMA assessed the issue of additional provider reporting and financial burden.</td>
<td>Section 5.4.2</td>
</tr>
<tr>
<td>9. How will the SMA verify providers’ use of certified electronic health record technology?</td>
<td>Section 5.4</td>
</tr>
<tr>
<td>10. How will the SMA collect providers’ meaningful use data, including the reporting of clinical quality measures? Does the State envision different approaches for the short-term and a different approach for the longer-term?</td>
<td>Section 5.4</td>
</tr>
<tr>
<td>11. * How will this data collection and analysis process align with the collection of other clinical quality measures data, such as CHIPRA?</td>
<td>Section 5.9</td>
</tr>
<tr>
<td>12. What IT, fiscal and communication systems will be used to implement the EHR Incentive Program?</td>
<td>Section 5.9</td>
</tr>
<tr>
<td>13. What IT systems changes are needed by the SMA to implement the EHR Incentive Program?</td>
<td>Section 5.9 and Section 4.1.1 enhanced in SMHP version 1.1</td>
</tr>
<tr>
<td>14. What is the SMA’s IT timeframe for systems modifications?</td>
<td>Section 5.9 enhanced in SMHP version 1.1</td>
</tr>
<tr>
<td>15. When does the SMA anticipate being ready to test an interface with the CMS National Level Repository (R&amp;A)?</td>
<td>Section 5.5.2 All interfaces between MS SLR and CMS have been tested, approved and deployed (D16, D18, E7, E8, etc...)</td>
</tr>
<tr>
<td>CMS Guidelines</td>
<td>Section C: Activities Necessary to Administer and Oversee the EHR Incentive Payment Program, continued</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>16. What is the SMA’s plan for accepting the registration data for its Medicaid providers from the CMS R&amp;A (e.g., mainframe to mainframe interface or another means)?</td>
<td>Section 5.3.3</td>
</tr>
<tr>
<td>17. What kind of website will the SMA host for Medicaid providers for enrollment, program information, etc?</td>
<td>Section 5.9.1</td>
</tr>
<tr>
<td>18. Does the SMA anticipate modifications to the MMIS and if so, when does the SMA anticipate submitting an MMIS I-APD?</td>
<td>DOM is preparing an annual update to the MMIS IAPD as well as an update to the MES IAPD. No HIT costs will occur.</td>
</tr>
<tr>
<td>19. What kinds of call centers/help desks and other means will be established to address EP and hospital questions regarding the incentive program?</td>
<td>Section 5.10.1.2</td>
</tr>
<tr>
<td>20. What will the SMA establish as a provider appeal process relative to: a) the incentive payments, b) provider eligibility determinations, and c) demonstration of efforts to adopt, implement or upgrade and meaningful use certified EHR technology?</td>
<td>Section 5.12</td>
</tr>
<tr>
<td>21. What will be the process to assure that all Federal funding, both for the 100 percent incentive payments, as well as the 90 percent HIT Administrative match, are accounted for separately for the HITECH provisions and not reported in a commingled manner with the enhanced MMIS FFP?</td>
<td>Section 5.10.2</td>
</tr>
<tr>
<td>22. (a) What is the SMA’s anticipated frequency for making the EHR Incentive payments (e.g., monthly, semi-monthly, etc.)? (b) What will be the process to assure that Medicaid provider payments are paid directly to the provider (or an employer or facility to which the provider has assigned payments) without any deduction or rebate?</td>
<td>Section 5.6</td>
</tr>
<tr>
<td>23. What will be the process to assure that Medicaid payments go to an entity promoting the adoption of certified EHR technology, as designated by the State and approved by the US DHHS Secretary, are made only if participation in such a payment arrangement is voluntary by the EP and that no more than 5 percent of such payments is retained for costs unrelated to EHR technology adoption?</td>
<td>Section 5.10.1.1</td>
</tr>
<tr>
<td>24. What will be the process to assure that there are fiscal arrangements with providers to disburse incentive payments through Medicaid managed care plans does not exceed 105 percent of the capitation rate per 42 CFR Part 438.6, as well as a methodology for verifying such information?</td>
<td>Not Done in State of MS</td>
</tr>
<tr>
<td>25. What will be the process to assure that all hospital calculations and EP payment incentives (including tracking EPs’ 15% of the net average allowable costs of certified EHR technology) are made consistent with the Statute and regulation?</td>
<td>This requirement is no longer relevant</td>
</tr>
<tr>
<td>26. What will be the role of existing SMA contractors in implementing the EHR Incentive Program – such as MMIS, PBM, fiscal agent, managed care contractors, etc.?</td>
<td>Section 5.10.1</td>
</tr>
<tr>
<td>CMS Guidelines  Section C: Activities Necessary to Administer and Oversee the EHR Incentive Payment Program, continued</td>
<td>Location in Document</td>
</tr>
<tr>
<td>---------------------------------------------------------------</td>
<td>---------------------</td>
</tr>
<tr>
<td>27. * States should explicitly describe what their assumptions are, and where the path and timing of their plans have dependencies based upon: The role of CMS (e.g., the development and support of the National Level Repository; provider outreach/help desk support) The status/availability of certified EHR technology The role, approved plans and status of the Regional Extension Centers The role, approved plans and status of the HIE cooperative agreements State-specific readiness factors</td>
<td>Section 6.3</td>
</tr>
</tbody>
</table>

*May be deferred
Cross Reference from CMS Guidelines to Section 5 – Provider Incentive Program Blueprint:

<table>
<thead>
<tr>
<th>CMS Guidelines Section D: The State’s Audit Strategy**</th>
<th>Location in Document***</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. (a) What will be the SMA’s methods to be used to avoid making improper payments? (Timing, selection of which audit elements to examine pre or post-payment, use of proxy data, sampling, how the SMA will decide to focus audit efforts etc):</td>
<td>Section 5.3.1, 5.3.3, 5.5.2, 5.10.1.1 prepayment checks</td>
</tr>
<tr>
<td>1. (b) Describe the methods the SMA will employ to identify suspected fraud and abuse, including noting if contractors will be used. Please identify what audit elements will be addressed through pre-payment controls or other methods and which audit elements will be addressed post-payment.</td>
<td>Section All sections listed for 1(a) and 5.3.2, 5.5.1.1, 5.5.1.2 5.6, 5.10.1.2.1, 5.10.2 prepayment actions</td>
</tr>
<tr>
<td>2. How will the SMA track the total dollar amount of overpayments identified by the State as a result of oversight activities conducted during the FFY?</td>
<td>Section 5.10.2, 5.7, 5.14 payment reporting</td>
</tr>
<tr>
<td>3. Describe the actions the SMA will take when fraud and abuse is detected.</td>
<td>Section 5.11.3</td>
</tr>
<tr>
<td>4. Is the SMA planning to leverage existing data sources to verify meaningful use (e.g., HIEs, pharmacy hubs, immunization registries, public health surveillance databases, etc.)? Please describe.</td>
<td>Appendix I - describes the requirement surrounding the State’s Immunization roll to provide documentation of registration.</td>
</tr>
<tr>
<td>5. Will the State be using sampling as part of audit strategy? If yes, what sampling methodology will be performed?* (i.e. probe sampling; random sampling)</td>
<td>Appendix J Audit Strategy will be submitted separately and confidentially</td>
</tr>
<tr>
<td>6. **What methods will the SMA use to reduce provider burden and maintain integrity and efficacy of oversight process (e.g., above examples about leveraging existing data sources, piggy-backing on existing audit mechanisms/activities, etc)?</td>
<td>Section 3.8, 4.1.1, 5.10.2</td>
</tr>
<tr>
<td>7. Where are program integrity operations located within the State Medicaid Agency, and how will responsibility for EHR incentive payment oversight be allocated?</td>
<td>Section 5.10.2 enhanced in SMHP version 1.1</td>
</tr>
</tbody>
</table>

*The sampling methodology part of this question may be deferred until the State has formulated a methodology based upon the size of their EHR incentive payment recipient universe.

** The Comprehensive Audit Strategy is referenced as Appendix J of the SMHP. However, Appendix J only contains the statement: *Appendix J will be submitted to CMS separate from this SMHP update to maintain confidentiality.* None of the post-payment audit information is contained in the public facing SMHP.

***SMHP content referenced in this section applies to Pre-payment actions and are not confidential.
## Cross Reference from CMS Guidelines to Section 6 – HIT Roadmap:

<table>
<thead>
<tr>
<th>CMS Guidelines Section E: The State’s HIT Roadmap</th>
<th>Location in Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. <em>Provide CMS with a graphical as well as narrative pathway that clearly shows where the SMA is starting from (As-Is) today, where it expects to be five years from now (To-Be), and how it plans to get there.</em></td>
<td>Section 4.3, 6.1</td>
</tr>
<tr>
<td>2. What are the SMA’s expectations re provider EHR technology adoption over time? Annual benchmarks by provider type?</td>
<td>Section 6.6.1, Table 6-1 enhanced in SMHP version 1.1</td>
</tr>
<tr>
<td>3. Describe the annual benchmarks for each of the SMA’s goals that will serve as clearly measurable indicators of progress along this scenario.</td>
<td>Sections 6.6.1, 6.6.2 and 6.6.2 replaced in SMHP version 1.1</td>
</tr>
<tr>
<td>4. Discuss annual benchmarks for audit and oversight activities.</td>
<td>Appendix J is the Audit Strategy, which is submitted as a separate document.</td>
</tr>
</tbody>
</table>

CMS is looking for a strategic plan and the tactical steps that SMAs will be taking or will take successfully implement the EHR Incentive Program and its related HIT/E goals and objectives. We are specifically interested in those activities SMAs will be taking to make the incentive payments to its providers, and the steps they will use to monitor provider eligibility including meaningful use. We also are interested in the steps SMAs plan to take to support provider adoption of certified EHR technologies. We would like to see the SMA’s plan for how to leverage existing infrastructure and/or build new infrastructure to foster HIE between Medicaid’s trading partners within the State, with other States in the area where Medicaid clients also receive care, and with any Federal providers and/or partners.

*Where the State is deferring some of its longer-term planning and benchmark development for HIT/E in order to focus on the immediate implementation needs around the EHR Incentive Program, please clearly note which areas are still under development in the SMA’s HIT Roadmap and will be deferred.*