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Disclaimer

The Arizona Health Care Cost Containment System Administration (AHCCCS) is providing this material as an informational reference for physician and non-physician practitioner providers.

Although every reasonable effort has been made to assure the accuracy of the information within these pages at the time of posting, the Medicare and Medicaid program is constantly changing, and it is the responsibility of each physician, non-physician practitioner; supplier or provider to remain abreast of the Medicare and Medicaid program requirements.


Important Notice – Third Party Attestation

The Arizona Medicaid Program does not allow third party attestation for Eligible Providers in the Electronic Provider Incentive Payment System (ePIP).

Eligible Providers should actively participate in the attestation process in ePIP.

Eligible providers are responsible for the completeness and accuracy of the information provided in their attestation in ePIP.
About ePIP

The Arizona Medicaid Promoting Interoperability Program (formerly the Electronic Health Record Incentive Program) will provide incentive payments to eligible professionals and eligible hospitals as they demonstrate adoption, implementation, upgrading, or meaningful use of certified EHR technology. This incentive program is designed to support providers in this period of Health IT transition and instill the use of EHRs in meaningful ways to help our nation to improve the quality, safety, and efficiency of patient health care.

This web application is for the Arizona Medicaid Promoting Interoperability Program. Those electing to partake in the program will use this system to register and participate in the program.

Administration:
The Arizona Health Care Cost Containment System (AHCCCS) is responsible for the implementation of Arizona’s Medicaid Promoting Interoperability Program. Until the end of the program, AHCCCS will disburse payments to providers who adopt, implement, upgrade or demonstrate meaningful use of certified EHR technology. For detailed information, visit AHCCCS website

Resources:
Reference materials for Registration and Attestation are available to explain how to complete these modules. Reference guides, eligibility and payment worksheets, links to a list of EHR technology that is certified for this program, and other general resources will help you complete registration and attestation. For detailed information, visit AHCCCS website

Eligible to Participate:
Providers under the AHCCCS Medicaid program are eligible to participate in the Arizona EHR Incentive Program if they meet the program’s requirements. For detailed information, visit AHCCCS website

Eligible Hospitals (EHs)
Medicaid EHs include:
- Acute Care Hospitals (including Critical Access Hospitals and Cancer Hospitals) with at least 10% Medicaid patient volume
- Children’s Hospitals (not required to meet a Medicaid patient volume)

Eligible Professionals (EPs)
Medicaid EPs include:
- Physicians
- Nurse Practitioners
- Certified Nurse - Midwife
- Dentists
- Physicians Assistants who practice in a Federally Qualified Health Center (FQHC) or Rural Health Clinic (RHC) that is led by the Physician Assistant

Additionally, Medicaid EPs must also:
- Have a minimum of 30% Medicaid patient volume
- Have a minimum of 20% or 30% patient volume for Pediatricians, OR
- Practice predominantly in a FQHC or RHC and have at least 30% patient volume attributed to needy individuals

NOTES: EPs may NOT be hospital-based. This is defined as any provider who furnishes 90% or more of their services in a hospital setting (inpatient or emergency department).

Practice predominantly is defined as any provider who furnishes over 50% of their services over a 6-month period at a FQHC/RHC facility.

Providers must complete and submit an attestation in the ePIP System each program year in order to apply for the program.

TIP
Go to the ePIP System by clicking here
Welcome to the ePIP System Home Page

AHCCCS Promoting Interoperability Program (formerly referred to as the EHR Incentive Payment Program)

This is the official web site for the Arizona Promoting Interoperability Program that provides incentive payments to eligible professionals and eligible hospitals as they adopt, implement, upgrade, or demonstrate meaningful use of certified EHR technology.

Your ePIP account is where you interface with the system to maintain your Promoting Interoperability Program information and track your incentive payments.

If you have not already registered with CMS and have not obtained a CMS Registration ID, click here to find out about registering with CMS.

NOTE: The deadline for registration in the Arizona Promoting Interoperability Program was June 30th, 2017 (the end of the 2016 Program Year). No new registrations are being accepted for this program, except for EPs enrolled in another state on or before Program Year 2016 and are transferring to Arizona. Contact the EHR Incentive Payments Team for more information.

The Centers for Medicare & Medicaid Services (CMS) governs the Promoting Interoperability Program. For more information please see the CMS.gov Promoting Interoperability Program.

ePIP Program Announcements

- CMS has re-branded the program as the Promoting Interoperability Program.
- Program Year 2018 will be open from January 1st 2019 thru December 31st 2019
- Stage 3 Meaningful Use in Program Year 2018 is:
  - Beginning in 2011, the Promoting Interoperability Program (formerly the Electronic Health Records (EHR) Incentive Program) was developed to encourage eligible professionals and eligible hospitals to adopt, implement, upgrade (AIU), and demonstrate meaningful use of certified EHR technology.
  - The program is administered voluntarily by states and territories, and will pay incentives through 2021. Eligible professionals are eligible for incentive payments for 5 years, and participation years do not have to be consecutive.
  - The last year that an eligible professional can begin participation is 2016. Incentive payments for eligible professionals under the Medicaid Promoting Interoperability Program are up to $63,750 over 6 years.
  - Eligible professionals can receive an incentive payment for adopting, implementing, or upgrading (AIU) certified EHR technology in their first year of participation. In subsequent years, eligible professionals can receive incentive payments for successfully demonstrating meaningful use.

What are Meaningful Use Stages?

Meaningful use requirements for 2017-2018: EPs with systems certified with a 2014 CEHRT will be attesting to Modified Stage 2 Objectives:

1. Protect electronic health information created or maintained by the CEHRT through the implementation of appropriate technical capabilities.
2. Use clinical decision support to improve performance on high-priority health conditions
3. Use computerized provider order entry for medication, laboratory, and radiology orders directly entered by any licensed health care professional who can enter orders into the medical record per state, local, and professional guidelines
4. Generate and transmit permissible prescriptions electronically (eRx).
5. The EP 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 care record for each transition of care or referral.
6. Use clinically relevant information from CEHRT to identify patient-specific education resources and provide these resources to the patient.
7. The EP who receives a patient from another setting of care or provider of care or believes an encounter is relevant performs medication reconciliation.
8. 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.
9. Use secure electronic messaging to communicate with patients on relevant health information.
10. The EP is in active engagement with a public health agency to submit electronic public health data from CEHRT exception where prohibited and in accordance with applicable law and practice.

Starting with Program Year 2017, providers with systems that have a 2015 CEHRT will be eligible to attest (optional) to Stage 3 Objectives:

1. Protect electronic protected health information (ePHI) created or maintained by the CEHRT through the implementation of appropriate technical, administrative, and physical safeguards.
2. Generate and transmit permissible prescriptions electronically (eRx)
3. Implement clinical decision support (CDS) interventions focused on improving performance on high-priority health conditions.
4. Use computerized provider order entry (CPOE) for medication, laboratory, and diagnostic imaging orders during 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.
5. The EP provides patients (or patient-authorized representative) with timely electronic access to their health information and patient-specific education.
6. Use CEHRT to engage with patients or their authorized representatives about the patient's care.
7. The EP provides a summary of care record when transitioning or referring their patient to another setting of care, receives or retrieves a summary of care record upon the receipt of a transition of 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.
8. The EP is in active engagement with a public health agency or clinical data registry 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.

Detailed documentation for all of these objectives can be found in the EHR Document Library.

The ePIP System Welcome screen consists of six menu navigational topics.

1. Home
2. Log On
3. Register
4. About
5. PI Doc Library
6. Contact Us

ePIP Program Announcement Update:

ePIP is accepting attestations for Program Year 2018 until August 31, 2019 (subject to CMS approval).

TIPS:
Helpful links are located in the footer of the web page.
Registration (Providers Without an ePIP Account)

Regarding Providers without an ePIP Account:

Only providers who already received payment and transferring to Arizona from other states can still set-up an ePIP account.

Providers must agree to the Terms & Conditions in order to register.

Program Year 2016 was the last year for providers to begin participation in the Promoting Interoperability Program.

You must agree by checking the box in order to proceed.

Your NPI number can be verified at the following link:
https://npiregistry.cms.hhs.gov/registry/
Use our PI Document Library to navigate quickly to the Meaningful Use requirements.

Click the link or Click the download button to view details on the 2018 Meaningful Use Objectives for Stage 2 Modified or Stage 3.

For more information on the 2018 Program Requirements at CMS, click here.
Providers who already have an ePIP account must log on in order to access their account.

If you forgot your password, you can reset your password by clicking the link below the Log On button.

Please allow an hour for server to respond to your request.

Go to the ePIP System by clicking here.

Need help? E-mail the Promoting Interoperability Program Team at EHRIncentivePayments@azahcccs.gov or call us at 602-417-4333.
Welcome to Your ePIP Account Home Page

Welcome To Your ePIP Account

Your ePIP account is where you interface with the system to maintain your qualifying information and track your incentive payments. The menu on the left-hand side of this page is where you navigate the various system functions.

The next step after you register is to Attest to create your application to receive your incentive payment. This is where you will input your system’s CMS EHR Certification ID & required patient volume metrics, as well as make your attestation MU (Meaningful Use) of EHR Certified technology.

You may go to Manage My Account at any time to check your information for accuracy and/or to make any changes to the contact information you have furnished. (e.g. Email address, contact person, etc.)

Once you attestation has been submitted, you can navigate to the Payments section to check the processing status of your incentive payments.

ePIP Program Announcements:

- CMS has re-branded the program as the Promoting Interoperability Program
- Program Year 2018 is now open and accepting attestations
- Stage 3 Meaningful Use in Program Year 2018 is optional

HOME

Returns you to this page.

MY ACCOUNT

- Manage My Account: Review & edit your contact information.
- Change My Password: Change the password for your account
- Modify My Security Questions: Create or modify the security questions associated with your account
- Payments: Track your payments for separate program years.
- Manage Documents: Upload supporting documentation for your attestations
- EHR Certificate Validation Tool: Determine if your CEHRT Identifier is valid

ATTEST

Create & maintain attestations for separate program years.

CONTACT US

Contact the AHCCCS EHR Incentive Payments Group

EHR DOCUMENT LIBRARY

A collection of PDF documents from CMS regarding the EHR Incentive Payment Program

TIP

Helpful links are located in the footer of the web page.

The ePIP Account Welcome screen consists of six menu topics to navigate through the attestation.

1. Home
2. My Account
   - Manage My Account
   - Change My Password
   - Modify My Security Questions
   - Payments
   - Manage Documents
   - EHR Certificate Validation Tool
3. Attest
4. Contacts
   - PI Team
   - Other AHCCCS Contacts
5. PI Doc Library
6. Log Off

ePIP Program Announcement Update:

ePIP is accepting attestations for Program Year 2018 until August 31, 2019 (subject to CMS approval).
My Account – How to Manage My Account

My Account page has six drop down navigation menus to help you manage your ePIP Account.

Let’s take a look at:

- Manage My Account
- Change My Password
- Modify My Security Questions
- Payments
- Manage Documents
- EHR Certificate Validation Tool

Manage My Account allows you to add an authorized secondary contact (optional).

This person does not have access to ePIP but is permitted to communicate with the State to answer general program inquiries and to help you gather your documentation for the attestation.

Click Edit My Account to add or update an authorized secondary contact.

TIP

Your data will appear here.

If incorrect or incomplete, follow the instructions below to modify.

Allow 48 hours for an update.
My Account page has six drop down navigation menus to help you manage your ePIP Account.

Let's take a look at:

- Manage My Account
- Change My Password
- Modify My Security Questions
- Payments
- Manage Documents
- EHR Certificate Validation Tool

Manage My Account allows you to add an authorized secondary contact (optional).

This person does not have access to ePIP but is permitted to communicate with the State to answer general program inquiries and to help you gather your documentation for the attestation.

Click Edit My Account to add or update an authorized secondary contact.
My Account – How to Manage My Password

Change Password

Use the form below to change your password. New passwords must meet the complexity requirements listed below.

Password Complexity Requirements:

- Minimum length of nine characters.
- Must contain at least one upper case alpha character. (ex: A)
- Must contain at least one lower case alpha character. (ex: a)
- Must contain at least one numeric character (ex: 1, 2, 3, etc.).
- Must contain at least one special character (!, @, #, $, etc.).
- The password cannot contain three or more consecutive characters. For example: “111” or “aAa” would not be accepted.
- The password cannot have 3 or more characters in common with the user name.

My Account page has six drop down navigation menus to help you manage your ePIP Account.

Let’s take a look at:

- Manage My Account
- Change My Password
- Modify My Security Questions
- Payments
- Manage Documents
- EHR Certificate Validation Tool

Change My Password allows you to modify your password at any time.

Enter your current password and then your new password.

Passwords must meet the complexity requirements displayed on the screen.
You must enter your password to modify your security questions.
## My Account – How to Manage My Payments

### Payment Status History

<table>
<thead>
<tr>
<th>Program Year</th>
<th>Amount</th>
<th>Payment Date</th>
<th>Payment For</th>
</tr>
</thead>
<tbody>
<tr>
<td>2012</td>
<td>$21,250.00</td>
<td>8/26/2013</td>
<td>AU</td>
</tr>
<tr>
<td>Details</td>
<td>Initial Payment made by AHCCCS on 8/26/2013 for $21250.00. Payment reference # 2688</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013</td>
<td>$8,500.00</td>
<td>11/25/2013</td>
<td>MU</td>
</tr>
<tr>
<td>Details</td>
<td>Initial Payment made by AHCCCS on 11/25/2013 for $8500.00. Payment reference # 2989</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2014</td>
<td>$8,500.00</td>
<td>12/23/2015</td>
<td>MU</td>
</tr>
<tr>
<td>Details</td>
<td>Initial Payment made by AHCCCS on 12/23/2015 for $8500.00. Payment reference # 4574</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2016</td>
<td>$8,500.00</td>
<td>7/24/2017</td>
<td>MU</td>
</tr>
<tr>
<td>Details</td>
<td>Initial Payment made by AHCCCS on 7/24/2017 for $8500.00. Payment reference # 6306</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Example Data Only

**Instructions**

Here is where you can track your incentive payments for separate program years. The processing status of your incentive payments will be displayed along with other payment details in the table above.

---

**TIP**

A payment processing status message is displayed to keep you updated.

---

**My Account page has six drop down navigation menus to help you manage your ePIP Account.**

Let’s take a look:

- Manage My Account
- Change My Password
- Modify My Security Questions
- Payments
- Manage Documents
- EHR Certificate Validation Tool

**Payments allow you to view your payment history and processing status.**
My Account – How to Manage My Documents

Example Data Only

Manage Documents

<table>
<thead>
<tr>
<th>Attestation Type</th>
<th>Attestation Year</th>
<th>File Name</th>
<th>Document Type</th>
<th>Name</th>
<th>Size</th>
<th>Uploaded</th>
<th>Delete</th>
</tr>
</thead>
<tbody>
<tr>
<td>MU3</td>
<td>4</td>
<td>PI Total Encounter QTR4</td>
<td>Meaningful Use DHR Report</td>
<td>Total encounters and unique patients during the measure period</td>
<td>27.0 KB</td>
<td>2/29/2017 2:34 PM</td>
<td>Delete</td>
</tr>
<tr>
<td>MU3</td>
<td>4</td>
<td>Summary, Report, ODM, 30606916 to 123116</td>
<td>Meaningful Use DHR Report</td>
<td>ODM Report</td>
<td>97.5 KB</td>
<td>2/29/2017 2:34 PM</td>
<td>Delete</td>
</tr>
<tr>
<td>MU3</td>
<td>4</td>
<td>Core OBL, 100016 to 123116, 0332457</td>
<td>Meaningful Use DHR Report</td>
<td>Core Objectives Report</td>
<td>22.3 KB</td>
<td>2/29/2017 2:34 PM</td>
<td>Delete</td>
</tr>
</tbody>
</table>

My Account page has six drop down navigation menus to help you manage your ePIP Account.

Let’s take a look at:

- Manage My Account
- Change My Password
- Modify My Security Questions
- Payments
- Manage Documents
- EHR Certificate Validation Tool

Manage Documents allows you to upload your documentation that supports your attestation.

Click Create New to upload documents.

Tag your documents by selecting the appropriate label from the drop down list:

- **Attestation Year** – describes the program year for the document
- **Document Type** – describes the type of document you are uploading.

Tip
The EHR Certification Number is a unique alpha-numeric character string assigned by ONC-Authorized Testing & Certification Board after an PI system has been successfully certified.
**Attestation**

*This Screen Shows Example Data Only*

<table>
<thead>
<tr>
<th>Attestation Completed.</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Details</td>
<td>View</td>
</tr>
<tr>
<td>First Year</td>
<td>2012</td>
</tr>
<tr>
<td>Details</td>
<td>View</td>
</tr>
<tr>
<td>Second Year</td>
<td>2013</td>
</tr>
<tr>
<td>Details</td>
<td>View</td>
</tr>
<tr>
<td>Third Year</td>
<td>2014</td>
</tr>
<tr>
<td>Details</td>
<td>View</td>
</tr>
<tr>
<td>Fourth Year</td>
<td>2016</td>
</tr>
<tr>
<td>Details</td>
<td>View</td>
</tr>
<tr>
<td>Fifth Year</td>
<td>2018</td>
</tr>
</tbody>
</table>

The Attest page is where you create your attestation & view your attestation activity.

Providers must attest if they want to participate in the program (maximum of 6 payments).

Please be sure to read the Meaningful Use Stage Review and the Data Requirements.

**Before Submission:**

Click the Create New button to start a new attestation *(new users)*.

Click the Begin button to start a new attestation *(existing users)*.

Click the Edit button to complete your attestation.

**TIP**

**After Submission:**

Click the Re-submit button to modify a previously failed/rejected attestation.

Click the Details button to view the details of your attestation.

Click the View button to see a status of your Attestation Progress.
Attestation Instructions

Welcome to the Attestation page. Arizona Medicaid providers must attest each payment year for the Medicaid Promoting Interoperability Program. Completing the State attestation is a prerequisite for determining the EHR Incentive Program payment.

In your first participation year, you demonstrated that you Adopted, Implemented or Upgraded your system to certified EHR technology. That was the first step in transforming our nation’s health care system to improve quality, safety and efficiency of care to EHR technology.

Attest Options

Depending on the current status of your attestation, please select one of the following actions:

- Begin: Begin Meaningful Use Attestation.
- Edit: Edit a previously started Meaningful Use Attestation that has not yet been submitted.
- Resubmit: Resubmit a failed or rejected attestation.
- Detail: View detail Meaningful Use Attestation that has been submitted and accepted.

* If you are a new user of the Arizona ePIP system, please select the “Create New” option at the top of the page.

Meaningful Use Stage Overview

Meaningful Use attestations require Medicaid Eligible Professionals (EPs) participating in the EHR Incentive Program to successfully demonstrate “meaningful use” of certified EHR technology. The reporting period for Meaningful Use is a minimum of 90 days.

Requirements for Meaningful Use Measures for EPs

- Meaningful Use Stage 2 consists of 10 Meaningful Use Objectives that must be met according to CMS threshold. If an EP meets the criteria for and can claim an exclusion for measures that have that option, then the measure(s) is also considered met.
- Meaningful Use Stage 3 consists of 8 Meaningful Use Objectives that must be met according to CMS threshold. If an EP meets the criteria for and can claim an exclusion for measures that have that option, then the measure(s) is also considered met.

Beginning in Program Year 2017, CMS adopted final policies to align specific CQMs available to EPs participating in the Medicaid EHR Incentive Program with those available to professionals participating in the Merit-based Incentive Payment System (MIPS).

Changes Include:

- The minimum amount of CQMs EPs must attest to has been reduced from 9 CQMs to 6 CQMs
- EPs are no longer required to attest to CQMs that cover a minimum amount of NQS domains
- 11 CQMS have been removed, leaving EPs the option to attest to 53 CQMs instead of 64 CQMs
Attestation Instructions continued

<table>
<thead>
<tr>
<th>Data Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Please be prepared to provide the following information:</td>
</tr>
<tr>
<td>Medicaid Patient Volume</td>
</tr>
<tr>
<td>• Patient Volume Reporting Period (90 days) ¹</td>
</tr>
<tr>
<td>• Hospital-Based Reporting Period (12 months) ¹</td>
</tr>
<tr>
<td>• Patient Volume Methodology (Individual/Aggregate) ²</td>
</tr>
<tr>
<td>• Total Patient Encounters</td>
</tr>
<tr>
<td>• Medicaid Patient Encounters (Medicaid Title XIX)</td>
</tr>
<tr>
<td>• Hospital-Based Patient Encounters (Medicaid Title XIX Inpatient Hospital &amp; Emergency Department)</td>
</tr>
</tbody>
</table>

Notes:
- ¹ Reporting periods are from the prior calendar year that precedes the payment year.
- ² For Individual Patient Volume Methodology:
  - Patient Volume criteria is based on Provider's data
  - Hospital-Based criteria is based on Provider's data

Additional Requirement:
Non-Hospital-Based Criteria:
EPs selecting Medicaid Patient Volume Type cannot be hospital-based. Hospital-Based Patient Encounters are encounters received at an inpatient hospital or an emergency department place of service. Hospital-Based EPs have 90 percent or more of their covered professional services in a hospital setting during the 12-month reporting period.

<table>
<thead>
<tr>
<th>Needy Individual Patient Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Patient Volume Reporting Period ¹</td>
</tr>
<tr>
<td>• Practice Predominantly Reporting Period ¹</td>
</tr>
<tr>
<td>• Patient Volume Methodology</td>
</tr>
<tr>
<td>• Total Patient Encounters</td>
</tr>
<tr>
<td>• Needy Individual Patient Encounters (Medicaid Title XIX CHF* Title XIX &amp; Patients Paying Below Cost)</td>
</tr>
<tr>
<td>• FQHC/RHC Facility Patient Encounters in Practice Predominantly Reporting Period</td>
</tr>
<tr>
<td>• Total Patient Encounters in Practice Predominantly Reporting Period</td>
</tr>
</tbody>
</table>

Notes:
- ¹ Reporting periods:
  - Patient Volume Reporting Period is a 90-day period in prior calendar year
  - Practice Predominantly Reporting Period is a 6-month period in prior calendar year

Additional Requirement:
Practice Predominantly Criteria
EPs selecting Needy Individual Patient Volume Type must practice predominantly at FQHC/RHC facilities. Practice Predominantly EPs have more than 50 percent of patient encounters at FQHC/RHC facilities place of service during the 6-month reporting period.

## AIU Selection

Note: As of the end of Program Year 2016 (June 30th, 2017) the AIU Selection is no longer available

- **Adopted Certified EHR**
  Adoption of an EHR system requires that a provider acquired, purchased or secured access to certified EHR technology.

- **Implemented Certified EHR**
  Implementation of an EHR system requires that a provider installed or commenced utilization of certified EHR technology.

- **Upgraded Certified EHR**
  Upgrade of an EHR system requires that a provider upgraded from existing EHR technology to certified EHR technology or expanded the functionality of existing certified EHR technology.
This is where you will monitor your progress towards completion of your attestation.

Note that the ability to complete the steps on this page is sequential. You must complete the steps in sequence (top down) to access subsequent sections.

The supporting documentation must be uploaded after you complete each step.

Click the Begin button to complete each step.

Click the Continue button to finish a step.

Click the Modify button to change information previously entered.

TIP
Please make certain that your contact detail is always up to date.

You must first update your contact changes in the CMS Registration and Attestation System at the following Link: [Click Here](https://www.azepip.gov/).

Wait at least 48 hours for the information you modified in the CMS Registration and Attestation System to feed to your ePIP account.

Did you know that you can enter an authorized secondary contact in ePIP?

This person does not have access to ePIP but is permitted to communicate with the State to answer general program inquiries and to help you gather your documentation for the attestation.

Go to My Account, Click Manage My Account and Click Edit My Account to update your authorized secondary contact *(optional)*.
Patient Volume Criteria

Patient volume is required each time you apply for the program.

Medicaid Patient Volume is an available option for all providers.

Needy Patient Volume is only an available option for providers practicing in a FQHC, RHC, or Tribal Clinic.

If you are attesting using your group Aggregate patient volume, every provider in the group must also select aggregate”.

Out of State Medicaid Patient encounters can be excluded in the numerator (if not needed to meet the patient volume) but must be reported in the denominator.

Note that inclusion of out of state patient encounters is optional in the numerator and slows the approval process since we must validate with the respective state(s).
# Report Medicaid Patient Volume Data Elements

**Report Patient Volume**

Please enter 90-day patient volume data from the calendar year prior to the Program Year for which you are attesting. For example, a Program Year 2018 attestation should have patient volume data from calendar year 2017.

### Reporting Period

- **Patient Volume Reporting Period Start Date**
- **Patient Volume Reporting Period End Date**

### All Patient Encounters

- **Total Patient Encounters**

**Note:** Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for the rendering provider. The EP must report all Medicaid & Non-Medicaid places of services when reporting the above total (denominator).

### Medicaid Patient Encounters

- **Arizona Medicaid Patient Encounters**

**Note:** Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for the rendering provider. The EP must report all Medicaid Title XIX places of services when reporting the above Medicaid patient encounters (numerator).

### Optional Border States

- **California Medicaid Patient Encounters**
- **Colorado Medicaid Patient Encounters**
- **New Mexico Medicaid Patient Encounters**
- **Nevada Medicaid Patient Encounters**
- **Utah Medicaid Patient Encounters**

Data to determine the Patient Volume includes all Place of Services.

- **TIP**

The numerator is Medicaid Title XIX patient encounters only.

The denominator is All patient encounters [Medicaid and Non-Medicaid].

**Medicaid Patient Volume**

- Is the percentage of Medicaid Title XIX patient encounters in the reporting period.

- Providers selecting this option must also demonstrate that they are not hospital-based.

**Patient Volume Reporting dates**

- Must be a continuous 90-day period selected from the year prior to the program year.

**Out of State Medicaid Patient encounters**

- Can be excluded in the numerator *(if not needed to meet the patient volume)* but must be reported in the denominator.
Providers selecting Medicaid Patient Volume must demonstrate that they are not hospital-based.

The Hospital-based Reporting date is the 12-month period from the year prior to the program year.

Hospital-Based providers have 90% or more of their Medicaid Title XIX patient encounters in a hospital setting defined as:
- Inpatient Hospital [POS 21]
- Emergency Department [POS 23]

Providers may need to obtain patient encounter data from the hospital and should consider requesting it in advance.

---

### Report Hospital-Based Patient Encounters

<table>
<thead>
<tr>
<th>Reporting Period</th>
<th>Start Date</th>
<th>End Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>(12 months in year prior to Program Year)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### All Medicaid Patient Encounters

| EP Total Medicaid Patient Encounters | | |
| Note: Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for the rendering provider. The EP must report all Medicaid Title XIX places of services when reporting the above total (denominator). | | |

### Medicaid Hospital-Based Patient Encounters

| EP Medicaid Inpatient Hospital Patient Encounters [POS21] | | |
| EP Medicaid Emergency Department Patient Encounters [POS23] | | |
| Note: Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for the rendering provider. The EP must report all Medicaid Title XIX Inpatient Hospital (places of service 21) & Emergency Department (places of service 23) only when reporting the hospital-based patient encounters (numerator). | | |

---

Data to determine the Medicaid Hospital-Based includes all Place of Services.

**Numerator** is Medicaid Title XIX IP & ED patient encounters only [POS 21 & POS 23].

**Denominator** is All Medicaid Title XIX patient encounters [All Place of Services].
# Report Needy Patient Volume Data Elements

## Report Patient Volume

<table>
<thead>
<tr>
<th>Reporting Period</th>
<th>(90 days in year prior to Program Year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Volume Reporting Period Start Date</td>
<td></td>
</tr>
<tr>
<td>Patient Volume Reporting Period End Date</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>EP Total Patient Encounters</th>
<th>(90 days in year prior to Program Year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Patient Encounters</td>
<td></td>
</tr>
</tbody>
</table>

**Note:** Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for the rendering provider. The EP must report all Medicaid & Non-Medicaid places of services when reporting the above total (denominator).

## Arizona Encounters

<table>
<thead>
<tr>
<th>Medicaid Title XIX</th>
<th>CHIP Title XXI</th>
<th>Patients Paying Below Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arizona Needy Individual Patient Encounters</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data to determine the Patient Volume includes all Place of Services.

**TIP**

The numerator is Needy Patient Encounters only.

The denominator is All patient encounters [Needy & Non-Needy].

Needy Patient Volume is the percentage of needy patient encounters in the reporting period.

Needy patient encounters are classified as Medicaid Title XIX, CHIP Title XXI & Patients Paying Below Cost (sliding scale) encounters.

Non-Needy patient encounters are Medicare, Private Insurance, Self-Pay, Commercial, etc.

Providers selecting this option must also demonstrate that they practiced predominantly in a FQHC, RHC or Tribal Clinic.

Patient Volume Reporting dates must be a continuous 90-day period selected from the year prior to the program year.
Report Needy Patient Volume Data Elements continued

Here is where you report your Medicaid out of state patient encounters for our Border States (optional if you wish to include in the numerator).

Please note that Out of State Medicaid Patient encounters can be excluded in the numerator (if not needed to meet the patient volume) but must be reported in the denominator.

<table>
<thead>
<tr>
<th>State</th>
<th>Medicaid Title XIX</th>
<th>CHIP Title XXI</th>
<th>Patients Paying Below Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>California Needy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual Patient Encounters</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colorado Needy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual Patient Encounters</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>New Mexico Needy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual Patient Encounters</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nevada Needy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual Patient Encounters</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Utah Needy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual Patient Encounters</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note that inclusion of out of state patient encounters is optional in the numerator and slows the approval process since we must validate with the respective state(s).
Providers selecting Needy Patient Volume must demonstrate that they practiced predominantly in a FQHC, RHC or Tribal Clinic.

Practice Predominantly Reporting dates is a 6-month period from the year prior to the program year.

Practice predominantly providers have more than 50% of their patient encounters in a FQHC, RHC or Tribal Clinic.

Data to determine the Practice Predominantly includes all Place of Services.

Numerator is FQHC, RHC or Tribal Clinic patient encounters only [inside facility].

Denominator is for All Place of Services [inside & outside the facility].
Note that as you complete each step:

- Column on the left changes from “Incomplete” to “Completed” status
- Column on the right changes from “Begin” to “Modify” designation.

Remember that each requirement task must be followed sequentially.

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Program Notes</th>
<th>Program Year Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>The data required for this attestation is grouped into categories. In order to complete your attestation, you must complete ALL of the tasks listed below. Click on the Begin button to start performing a given step. If a step has been started, but not completed, click on the Continue button to finish a step. Once a step is finished, you can click on the Modify button to change any information that was previously entered.</td>
<td>The AHCCCS Promoting Interoperability Program is currently open for Program Year 2018.</td>
<td>We encourage all providers to review the CMS documentation for Program Year 2018 before attesting. These documents are available on the CMS website, or in ePPIP in the ePPIP Document Library.</td>
</tr>
</tbody>
</table>

**TIP**

- Click the Begin button to complete each step.
- Click Continue button to finish a step.
- Click Modify button to change information previously entered.
Attestation Information

(*) Red asterisk indicates a required field.

**EHR Certification number**

- Please provide your EHR Certification number: 

- Please provide the date the system with the EHR Certification number above was implemented:

**EHR Reporting Period**

Program Year: 2018 (selecting your reporting period from Calendar Year 2018)

Please select an EHR Reporting Period of 90 days.

- EHR Reporting Period Start Date

- EHR Reporting Period End Date

**EHR Reporting Period Note:**
This range applies to Meaningful Use Objective Measures. The Meaningful Use EHR Report should align with this data range.

**CQM Reporting Period Note:**
This range applies to Clinical Quality Measures. The CQM Report should align with this data range.

**EHR Locations**

For providers who work at multiple sites, at least 50% of all encounters must take place at a location(s) with a certified EHR technology (CEHRT) system. Please specify:

- Do you work at multiple practice locations?

- Enter the total number of locations:

Eligible professionals who practice in multiple locations must take some additional steps in order to successfully participate in the Medicaid Electronic Health Record (EHR) Incentive Program. Below are links to the CMS Tip Sheets for Stage 2 and Stage 3 outlining these steps.

**Stage 2 Tip Sheet**

<table>
<thead>
<tr>
<th>Address</th>
<th>Suite #</th>
<th>City</th>
<th>State</th>
<th>Zip</th>
</tr>
</thead>
</table>

**Stage 3 Tip Sheet**

- Enter any additional practice address(es) with CEHRT:

<table>
<thead>
<tr>
<th>Address</th>
<th>City</th>
<th>State</th>
<th>Zip</th>
</tr>
</thead>
</table>

| Add |

**Encounters**

- Total patient encounters at all locations during the EHR Reporting Period:

- Total patient encounters at locations with CEHRT during the EHR Reporting Period:

**Note:** CMS defines patient encounters as any encounter where a medical treatment is provided and/or evaluation and management services are provided, except a hospital inpatient department (Place of Service 21) or a hospital emergency department (Place of Service 23). Patient encounters in ambulatory surgical centers would be included for the purpose of this definition.

**Stage 2 (Modified):** At least 50% of unique patients seen at locations with certified EHR technology must have their data in a certified EHR during the EHR reporting period.

**Stage 3:** At least 80% of unique patients seen at locations with certified EHR technology must have their data in a certified EHR during the EHR reporting period.

Please specify:

- Total unique patients during the EHR Reporting Period:

- Total unique patients have their data in a Certified EHR system during the EHR Reporting Period:

**Next** **Cancel**

You are now ready to being attesting to the Meaningful Use portion of the attestation.

First, we will need some general information about your PI system. Be sure to tell us if you have patients that are still maintained on paper records (Non-CEHRT).

You must select your PI Reporting Period start & end date from calendar year 2018 for the Meaningful Use Objectives & Clinical Quality Measures that you are attesting to.

Complete the number of unique patient encounters in your PI reporting period.

Complete the number of unique patients in your PI reporting period.
Program Year 2018 Flexibility Information

Providers have the option of attesting to Stage 2 or Stage 3 depending on their system’s certification (in effect no later than December 31, 2018).

Rules for Stage 3 participation:

- Providers with technology certified to a combination of the 2015 Edition & 2014 Edition (if the mix of certified technologies would not prohibit them from meeting the Stage 3 measures).
- Providers with technology certified for the 2015 Edition.
- Providers in the second year or greater of Meaningful Use participation.

Flexibility:

Based on the CEHRT year entered & your MU Participation Year you have the option of attesting to either Stage 2 or Stage 3.

Providers must review the details of Stage 3 before making a selection.

Click one of the following buttons:

- Attest to Stage 2
- Attest to Stage 3

NOTE: Once a Stage is selected, it cannot be undone without the PI Staff deleting your attestation (will cause re-work for the provider).
Attestation Progress (After Attestation Information)

Note that as you complete each step:

☑ Column on the left changes from “Incomplete” to “Completed” status
☑ Column on the right changes from “Begin” to “Modify” designation.

Remember that each requirement task must be followed sequentially.

TIP

Click the Begin button to complete each step.
Click Continue button to finish a step.
Click Modify button to change information previously entered.
Meaningful Use Objectives for Stage 3 (Optional)

<table>
<thead>
<tr>
<th>Providers with systems certified with a 2015 CEHRT as of 12.31.2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Protect electronic protected health information (ePHI) created or maintained by the certified electronic health record technology (CEHRT) through the implementation of appropriate technical, administrative, and physical safeguards.</td>
</tr>
<tr>
<td>2 Generate and transmit permissible prescriptions electronically (eRx)</td>
</tr>
<tr>
<td>3 Implement clinical decision support (CDS) interventions focused on improving performance on high-priority health conditions.</td>
</tr>
<tr>
<td>4 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.</td>
</tr>
<tr>
<td>5 The eligible professional (EP) provides patients (or patient-authorized representative) with timely electronic access to their health information and patient-specific education.</td>
</tr>
<tr>
<td>6 Use certified electronic health record technology (CEHRT) to engage with patients or their authorized representatives about the patient’s care.</td>
</tr>
<tr>
<td>7 The eligible professional (EP) provides a summary of care record when transitioning or referring their patient to another setting of care, receives or retrieves a summary of care record upon the 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 electronic health record (EHR) using the functions of certified EHR technology (CEHRT).</td>
</tr>
<tr>
<td>8 The eligible professional (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 electronic health record technology (CEHRT), except where prohibited, and in accordance with applicable law and practice.</td>
</tr>
</tbody>
</table>

Welcome to Stage 3

Providers must attest to 8 Meaningful Use Objectives using EHR 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 in the numerator must occur during the PI reporting period.

Stage 3 includes flexibility within certain objectives to allow providers to choose the measures most relevant to their patient population or practice. Stage 3 flexible measures include:

- Coordination of Care & Patient Engagement … You must meet thresholds for at least 2 of 3 measures
- Health Information Exchange… You must meet the thresholds for at least 2 of 3 measures
- Public Health Reporting … You must report on at least 2 of 3 measures.
Stage 3 Objective 1 Measure 1 Protect Patient Health Information

Meaningful Use Objectives - Stage 3 for Program Year 2018

ePIP Measure 1 of 20 - CMS Meaningful Use Objective 1, Measure 1

Protect Patient Health Information

Objective Details:

Protect Patient Health Information: Protect electronic protected health information (ePHI) created or maintained by the CEHRT through the implementation of appropriate technical, administrative, and physical safeguards.

Measure Requirements:

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)(ii) 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.

Additional Information:

- To meet Stage 3 requirements for an HI reporting period in 2018, all providers must use technology certified to the 2015 Edition. A provider who has technology certified to a combination of the 2013 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
- EPs must conduct or review a security risk analysis of CEHRT including addressing encryption/security of data, and implement updates as necessary at least once each calendar year and attest to conducting the analysis or review.
- It is acceptable for the security risk analysis to be conducted outside the HI reporting period; however, the analysis must be unique for each HI reporting period, the scope must include all HI reporting period and it must be conducted within the calendar year of the HI reporting period (January 1st – December 31st).
- An analysis must be done upon installation or upgrade to a new system and a review must be conducted covering each HI reporting period. Any security updates and deficiencies that are identified should be included in the provider’s risk management process and implemented or corrected as dictated by that process.
- The security risk analysis requirement under 45 CFR 164.308(a)(1) must assess the potential risks and vulnerabilities to the confidentiality, availability, and integrity of all ePHI that an organization creates, receives, maintains, or transmits. This includes ePHI in all forms of electronic media, such as hard drives, floppy disks, CDs, DVDs, smart cards or other storage devices, personal digital assistants, transmission media, or portable electronic media.
- All minimum providers should be able to show a plan for preventing or mitigating deficiencies and that steps are being taken to implement that plan.
- The parameters of the security risk analysis are defined in 45 CFR 164.308(a)(1), which was created by the HIPAA Security Rule. Meaningful use does not impose new or expanded requirements on the HIPAA Security Rule nor does it require specific use of every certification and standard that is included in certification of EHR technology. More information on the HIPAA Security Rule can be found at http://www.hhs.gov/ocr/privacy/security/administrative/security/oracle/.

Regulatory References:

- This objective may be found in Section 42 of the code of the federal register at 495.24 (d)(1)(ii) (A) and (B). For further discussion please see 80 FR 52692.
- In order to meet this objective and measure, an HI must possess the capabilities and standards of CEHRT at 45 CFR 170.315 (d)(1) through (d)(4).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measurement requirements for this particular objective. (Please review before attesting to this measure).

For detailed information about the Protect Patient Health Information objective, please click here.

Note: (Please Review before attesting to this measure): Further information about SRA can be found in the CMS SRA Fact Sheet, please click here.

Note: (Please Review before attesting to this measure): Further information about SRA can be found in the AHCCCS SRA Fact Sheet, please click here.

Supporting Documentation Requirements:

The Security Risk Analysis measure requires supporting documentation to be uploaded. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process. If you previously submitted the SRA documentation to Arizona in a prior program year, please submit any updates to those documents for this program year.

The supporting documentation should include the following elements for verification:

- The date that the Security Risk Analysis was completed, reviewed or updated (Please consult the CMS Measure Documentation and the Tip Sheet via the links above to insure that this date falls within the acceptable date range for the program year).
- Risk Analysis document (which should include information verifying the items listed below):
  - Potential threats and vulnerabilities were assessed.
  - An Asset Inventory was performed.
  - An Assessment of current security measures was performed.
  - Likelihood and Potential Impact of a threat occurrence.
  - Level of Risk determined by the assessments above.
- Action Plan document (which should include information verifying the items listed below):
  - What steps has the practice taken to respond to mitigate the identified risks?
  - Who is/are the individual(s) responsible for implementing the required changes?
  - When will the required changes be implemented?

(*) Red asterisk indicates a required field
(>) Gray asterisk indicates a conditionally required field

Measure Entry:

Complete the following information:

- Have you conducted or reviewed a security risk analysis per 45 CFR 164.308(a)(1), including addressing the security (including encryption) of ePHI created or maintained by CEHRT in accordance with requirements under 45 CFR 164.312(a)(2)(ii) and 45 CFR 164.306(d)(3). Did you implement security updates as necessary, and correct identified security deficiencies as part of your risk management process?
  - Yes
  - No

Enter the date you completed your security risk analysis:

TIP:

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 2 Measure 1 Electronic Prescribing (eRx)

Electronic Prescribing (eRx): Generate and transmit permissible prescriptions electronically (eRx).

Measure Requirements:

- More than 60 percent of all permissible prescriptions written by the EP are queried for a drug formulation and transmitted electronically using CEHRT.

Additional Information:

- To meet Stage 3 requirements for an EP reporting period in 2018, 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
- The provider is permitted, but not required, to limit the measure of this objective to those patients whose records are maintained using certified EHR technology (CEHRT).
- Authorizations for items such as durable medical equipment, or other items and services that may require EP authorization before the patient could receive them, are not included in the definition of prescriptions. These are excluded from the numerator and the denominator of the measure.
- Instances where patients specifically request a paper prescription may not be excluded from the denominator of this measure. The denominator includes all prescriptions written by the EP during the PI reporting period.
- As electronic prescribing of controlled substances is not possible, providers may choose to include these prescriptions in their permissible prescriptions where feasible and allowable by state and local law. If a provider chooses to include such prescriptions, he or she must do so uniformly across all patients and across all allowable schedules for the duration of the PI reporting period.
- Over the counter (OTC) medications are excluded from the definition of prescription.
- An EP needs to use CEHRT as the sole means of creating the prescription, and when transmitting to an external pharmacy that is independent of the EP’s organization such transmission must use standards adopted for EHR technology certification.
- EPs should include in the numerator and denominator both types of electronic transmissions (those within and outside the organization) for the measure of this objective.
- For purposes of counting prescriptions “generated and transmitted electronically,” we consider the generation and transmission of prescriptions to occur concurrently if the prescriber and dispenser are the same person and/or are accessing the same record in an integrated EHRI to creating an order in a system that is electronically transmitted to an internal pharmacy.
- Providers can use intranetwork networks that convert information from the certified EHRI into a computer-based fax in order to meet this measure as long as the EP generates an electronic prescription and transmits it electronically using the standards of CEHRT to the intranetwork, and this results in the prescription being filled without the need for the provider to communicate the prescription in an alternative manner.
- Prescriptions transmitted electronically within an organization (the same legal entity) do not need to use the NCQOP standards. However, an EP’s EHRI must meet all applicable certification criteria and be certified as having the capability of meeting the external transmission requirements of § 170.354(b). In addition, the EHRI that is used to transmit prescriptions within the organization would need to be CEHRT. For more information, refer to CMS’s FAQ at https://www.healthit.gov/policy-researchers-implementers/qa/question-12-10-022.
- Providers may limit their effort to query a pharmacy to simply using the function available to them in their CEHRT with no further action required. If a query using the function of their CEHRT is not possible or shows no result, a provider is not required to conduct any further manual or paper-based action in order to complete the query, and the provider may count the prescription in the numerator.
- EPs are practicing at multiple locations are eligible for the exclusion if any of their practice locations that are equipped with CEHRT meet the exclusion criteria.
- EPs that are part of an organization that owns or operates its own pharmacy within the 10-mile radius are not eligible for the exclusion regardless of whether pharmacy can accept electronic prescriptions from EPs outside of the organization.

TIP:

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

The Navigation bar at the bottom will monitor your progress.

For detailed information about the Electronic Prescribing objective, please click here.

Supporting Documentation Requirements:

Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

(*) Red asterisk indicates a required field

Measure Entry:

Exclusion 1: Based on ALL patient records; Any EP who writes fewer than 100 permissible prescriptions during the PI reporting period would be excluded from this requirement. Exclusion from this requirement does not prevent an EP from achieving meaningful use.

- Does this exclusion apply to you?
  - Yes
  - No

Exclusion 2: Based on ALL patient records; Any EP who 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/her PI reporting period would be excluded from this requirement. Exclusion from this requirement does not prevent an EP from achieving meaningful use.

- Does this exclusion apply to you?
  - Yes
  - No

PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology.

This data was extracted from both paper records as well as records maintained using Certified EHR Technology (CEHRT).

This data was extracted only from patient records maintained using certified EHR technology.

Complete the following information:

Numerator: The number of prescriptions in the denominator generated, queried for a drug formulation 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 PI reporting period, or number of prescriptions written for drugs requiring a prescription in order to be dispensed during the PI reporting period.

<table>
<thead>
<tr>
<th>Numerator:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Denominator:</td>
</tr>
</tbody>
</table>
### Stage 3 Objective 3 Measure 1 Clinical Decision Support

**Meaningful Use Objectives - Stage 3 for Program Year 2018**

**ePIP Measure 3 of 20 - CMS Meaningful Use Objective 3, Measure 1**

**Clinical Decision Support - Measure 1 of 2**

#### Objective Details:

**Clinical Decision Support - Measure 1 of 2**: Implement clinical decision support (CDS) interventions focused on improving performance on high-priority health conditions.

#### Measure Requirements:

Implement five clinical decision support interventions related to four or more CQMs at a relevant point in patient care for the entire PI reporting period. Absent four CQMs related to an EHR's scope of practice or patient population, the clinical decision support interventions must be related to high-priority health conditions.

#### Additional Information:

- To meet Stage 3 requirements for an EHR reporting period in 2018, 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
- Providers should implement the CDS intervention at a relevant point in clinical workflows when the intervention can influence clinical decision making before diagnostic or treatment action is taken in response to the intervention.
- Well-designed CDS encompasses a variety of workflow optimized information tools, which can be presented to providers, clinical and support staff, patients, and other caregivers at various points in time. These may include but are not limited to: computerized alerts and reminders for providers and patients; information displays or lists; context-aware knowledge retrieval specifications which provide a standard mechanism to incorporate information from online resources (commonly referred to as infobuttons); clinical guidelines; condition-specific order sets; focused patient data reports and summaries; documentation templates; dagnostic support; and contextually relevant reference information. These functionalities may be deployed on a variety of platforms (that is, mobile, cloud-based, installed).
- The same interventions do not have to be implemented for the entire PI reporting period as long as the threshold of 5 is maintained for the duration of the PI reporting period.
- While the ONC 2015 Edition final rule specifies that the CDS module that is certified to the CDS standard must have certain capabilities to provide or enable CDS for provider use, it does not certify the supports or resources themselves.
- If there are limited CQMs applicable to an EHR's scope of practice, the EP should implement CDS interventions that he or she believes will drive improvements in the delivery of care for the high-priority health conditions relevant to their specialty and patient population. These high-priority conditions must be determined prior to the start of the PI reporting period in order to implement the appropriate CDS to allow for improved performance.
- Drug-drug and drug-allergy interaction alerts are separate from the 5 clinical decision support interventions and do not count toward the 5 required for this first measure.

#### Definition of Terms:

**Clinical Decision Support**: HIT functionality that builds upon the foundation of an EHR to provide persons involved in care processes with general and person-specific information, intelligently filtered and organized, at appropriate times, to enhance health and health care.

**Regulatory References:**

- [This objective may be found in Section 42 of the code of the federal register at 495.24 (a)(2)(i)(A) and (B).](https://www.azepip.gov/)
- [In order to meet this objective and measure, an EP must use the capabilities and standards of CEDHIT at 45 CFR 170.315(a)(9) and (a)(4).](https://www.azepip.gov/)

**The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective.** (Please review before attesting to this measure)

For detailed information about the Clinical Decision Support objective, please click here

#### Supporting Documentation Requirements:

Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

(* Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

### Measure Entry:

**Have you implemented five clinical decision support interventions related to four or more clinical quality measures, or high-priority health conditions?**

- [ ] Yes
- [ ] No

---

**TIP:**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 3 Measure 2 Clinical Decision Support

Meaningful Use Objectives - Stage 3 for Program Year 2018
ePIP Measure 4 of 20 - CMS Meaningful Use Objective 3, Measure 2
Clinical Decision Support - Measure 2 of 2

Objective Details:

Clinical Decision Support - Measure 2 of 2: Implement clinical decision support (CDS) interventions focused on improving performance on high priority health conditions.

Measure Requirements:

The EP has enabled and implemented the functionality for drug-drug and drug-allergy interaction checks for the entire PI reporting period.

Additional Information:

- To meet Stage 3 requirements for an PI reporting period in 2018, 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
- Providers should implement the CDS intervention at a relevant point in clinical workflows where the intervention can influence clinical decision making before diagnostic or treatment action is taken in response to the intervention.
- Well-designed CDS encompasses a variety of workflow optimized information tools, which can be presented to providers, clinical and support staff, patients, and other caregivers at various points in time. These may include but are not limited to: computerized alerts and reminders for providers and patients; information displays or notes; content aware knowledge retrieval specifications which provide a standard mechanism to incorporate information from online resources (commonly referred to as InfoButtons); clinical guidelines; condition-specific order sets, focused patient data reports and summaries; documentation templates; diagnostic support; and contextually relevant reference information. These functionalities may be deployed on a variety of platforms (that is, mobile clients, web-based).
- The same interventions do not have to be implemented for the entire PI reporting period as long as the threshold of 5 is maintained for the duration of the PI reporting period.
- While the ONC 2015 Edition final rule specifies that the CDS module that is certified to the 2015 standard must have certain capabilities to provide or enable CDS for provider use, it does not certify the supports or resources themselves.
- If there is limited CDS applicable to an EP’s scope of practice, the EP should implement CDS interventions that he or she believes will drive improvements in the delivery of care for the high-priority health conditions relevant to their specialty and patient population. These high-priority conditions must be determined prior to the start of the PI reporting period in order to implement the appropriate CDS to allow for improved performance.
- Drug-drug and drug-allergy interaction alerts are separate from the 5 clinical decision support interventions and do not count toward the 5 required for this first measure.

Definition of Terms:

Clinical Decision Support - HIT functionality that builds upon the foundation of an EHR to provide persons involved in care processes with general and person-specific information, intelligently filtered and organized, at appropriate times, to enhance health and health care.

Regulatory References:

- This objective may be found in Section 42 of the code of the federal register at 459.24 (a)(2)(iii)(A) and (B). For further discussion please see 457.628D0
- In order to meet this objective and measure, an EP must use the capabilities and standards of CERHT at 45 CFR 170.316(a)(5) and (a)(6).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please review before attesting to this measure)

For detailed information about the Clinical Decision Support objective, please click here

Supporting Documentation Requirements:

Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attestation process.

(9) Red asterisk indicates a required field
(7) Gray asterisk indicates a conditionally required field

Measure Entry:

Exclusion: Based on ALL patient records: Any EP who writes fewer than 100 medication orders during the PI reporting period would be excluded from this requirement. Exclusion from this requirement does not prevent an EP from achieving meaningful use.

* Does this exclusion apply to you?

Yes ☐ No ☑

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?

Yes ☐ ☑ No ☐

May 10, 2019
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Stage 3 Objective 4 Measure 1 Computerized Provider Order Entry

Meaningful Use Objectives - Stage 3 for Program Year 2018

ePIP Measure 5 of 20 - CMS Meaningful Use Objective 4, Measure 1

Computerized Provider Order Entry - Measure 1 of 2

Objective Details:

Computerized Provider Order Entry - Measure 1 of 3: 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 Requirements:

More than 60 percent of medication orders created by the EP during the PI reporting period are recorded using computerized provider order entry.

Additional Information:

- To support the requirements for an PI reporting period in 2018, 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
- The provider is permitted, but not required, to limit the measure of this objective to those patients whose records are maintained using certified EHR technology (CEHRT).
- The CPOE function must be used to create the first record of the order that becomes part of the patient's medical record and before any action can be taken on the order to count in the numerator.
- In some situations, it may be impossible or inadvisable to wait until the data are collected and uploaded before making the order, such as situations where an intervention is identified and immediately initiated by the provider or initiated immediately after a verbal order by the ordering provider to a licensed healthcare professional under the provider’s direct supervision. Therefore in these situations, and as long as the order is entered using CPOE by a licensed healthcare professional, a certified medical assistant or other appropriately credentialed staff member to create the first record of that order (i.e., becomes part of the patient's medical record), those orders would count in the numerator of the CPOE measure.
- Any licensed healthcare providers and clinical staff credentialed to and with the duties equivalent of a medical assistant or is appropriately credentialed and performs administrative services similar to a medical assistant, but carries a more specific title due to other specialization in their duties or to the specialty of the medical professional they assist, can enter orders into the medical record for purposes of including the order in the numerator for the objective of CPOE if they originate the order per state, local, and professional guidelines.
- It is up to the provider to determine the proper credentialing, training, and duties of the medical staff entering the orders as long as they fit within the guidelines prescribed. Credentialing for a medical assistant must come from an organization other than the organization employing the medical assistant.
- An EP must satisfy all three measures for this objective through a combination of meeting the thresholds and exclusions (or both).
- Orders involving tele-health or remote communication (such as phone orders) may be included in the numerator as long as the order entry otherwise meets the requirements of the objective and measures.
- Providers may exclude orders that are predetermined for a given patient characteristics or for a given procedure (also known as “protocol” or “standing orders”) from the calculation of CPOE numerators and denominators. Note this does not require providers to exclude this category of orders from their numerator and denominator (77 FR 52694).
- CPOE is the entry of the order into the patient’s EHR that uses a specific function of CEHRT. CPOE does not otherwise specify how the order is filled or otherwise carried out.

Definition of Terms:

Computerized Provider Order Entry (CPOE) - A provider’s use of computer assistance to directly enter medical orders (for example, medications, consultations with other providers, laboratory services, imaging studies, and other ancillary services) from a computer or mobile device.

Laboratory Order - An order for any service provided by a laboratory that could not be performed by a non-laboratory.

Radiology Order - An order for any imaging service that uses electronic product radiation. The EP can include orders for other types of imaging services that do not rely on electronic product radiation in this definition as long as the policy is consistent across all patients and for the entire PI reporting period.

Regulatory References:

- This objective may be found in Section 42 of the code of the federal register at 495.24 (d)(4)(i)(A) and (B). For further discussion please see 80 FR 62840
- In order to meet this objective and measure, an EP must use the capabilities and standards of CEHRT at 45 CFR 170.315(a)(1) through (3).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measurement process.

For detailed information about the Computerized Provider Order Entry objective, please click here

Supporting Documentation Requirements:

Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attestation process.

(*) Red asterisk indicates a required field
(+) Gray asterisk indicates a conditionally required field

Measure Entry:

Exclusion: Based on ALL patient records. Any EP who writes fewer than 100 medication orders during the PI reporting period would be excluded from this requirement. Exclusion from this requirement does not prevent an EP from achieving meaningful use.

* Does this exclusion apply to your?  
  Yes  No
  * PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology
  * This data was extracted from both paper records as well as records maintained using CEHRT
  * This data was extracted only from patient records maintained using CEHRT

Complete the following information:

Numerator: The number of medication orders in the denominator during the PI reporting period that are recorded using CPOE.

Denominator: The number of medication orders created by the EP during the PI reporting period.

TIP:
Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page. Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 4 Measure 2 Computerized Provider Order Entry

Meaningful Use Objectives - Stage 3 for Program Year 2018

Computerized Provider Order Entry - Measure 2 of 3: 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 Requirements:

- More than 60 percent of laboratory orders created by the EP during the PI reporting period are recorded using computerized provider order entry.

Additional Information:

- To meet Stage 3 requirements for an PI reporting period in 2018, 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 prevent them from meeting the Stage 3 measures. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.

- The provider is permitted, but not required, to limit the measure of this objective to those patients whose records are maintained using certified EHR technology (CEHRT).

- The CPOE function must be used to create the first record of the order that becomes part of the patient’s medical record and before any action can be taken on the order to count in the numerator.

- In some situations, it may be impossible or infeasible to wait until a record of the order has been created. For example, situations where an intervention is identified and immediately initiated by the provider, or initiated immediately after a verbal order by the ordering provider to a licensed healthcare professional under his/her direct supervision. Therefore in these situations, so long as the order is entered using CPOE by a licensed healthcare professional, certified medical assistant or other appropriately credentialed staff member to create the first record of that order as it becomes part of the patient’s medical record, these orders would count in the numerator of the CPOE measure.

- Any licensed healthcare professionals and clinical staff credentialed to and with the duties equivalent of a medical assistant or is appropriately credentialed and performs assistive services similar to a medical assistant, but carries a more specific title due to either specialization of their duties or to the specialty of the medical professional they assist, can enter orders in the medical record for purposes of including the order in the numerator for the objectives of CPOE. If they can originate the order per state, local and professional guidelines. It is up to the provider to determine the proper credentialing, training, and duties of the medical staff entering the orders as long as they fit within the guidelines prescribed. Credentialing for a medical assistant must come from an organization rather than the organization employing the medical assistant.

- An EP must satisfy all three measures for this objective through a combination of meeting the thresholds and exclusions (or both).

- Orders involving tele-health or remote communication (such as phone orders) may be included in the numerator, as long as the order entry otherwise meets the objective of the measure and requirements.

- Providers may exclude orders that are predetermined for a given set of patient characteristics or for a given procedure (also known as “protocols” or “standing orders”) from the calculation of CPOE numerators and denominators. Note this does not require providers to exclude this category of orders from their numerator and denominator (77 F.3296).

- CPOE is the entry of this objective into the patient’s EHR that uses a specific function of CEHRT. CPOE does not otherwise specify how the order is filled or otherwise carried out.

Definition of Terms:

Computerized Provider Order Entry (CPOE): A provider’s use of computer assistance to directly enter medical orders (for example, medications, consultations with other providers, laboratory services, imaging studies, and other auxiliary services) from a computer or mobile device.

Diagnostic Imaging: Includes other imaging tests such as ultrasonic, magnetic resonance, and computed tomography in addition to traditional radiography.

Laboratory: A facility for the biological, microbiological, serological, chemical, immunological, hematological, biophysical, cytological, pathological, or other examination of from the human body for the purpose of providing information for the diagnosis, prevention, or treatment of any disease or impairment of, or the assessment of the health of, human beings. These examinations also include procedures to determine measure, or otherwise describe the presence or absence of various substances or organisms in the body. Facilities only collecting or preparing specimens (or both) or only serving as a mailing service and not performing testing are not considered laboratories.

Radiology: An order for any imaging service that uses electrical product radiation. The EP can include orders for other types of imaging services that do not rely on electrical product radiation in this definition as long as the service is consistent across all patients and for the entire PI reporting period.

TIP: Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

Adding Documentation Requirements:

Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

(1) Red asterisk indicates a required field

(1) Gray asterisk indicates a conditionally required field

Measure Entry:

Exclusion: Based on ALL patient records: Any EP who writes fewer than 100 medication orders during the PI reporting period would be excluded from this requirement. Exclusion from this requirement does not prevent an EP from achieving meaningful use.

- Does this exclusion apply to you?

  Yes ☐ No ☑

- If you select Yes, please select one of the following:

  - PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology.
  
    - This data was extracted from both paper records as well as records maintained using Certified EHR Technology (CEHRT) ☐
    
    - This data was extracted only from paper records maintained using certified EHR technology ☐

  - NURSING: The number of laboratory orders in the denominator during the PI reporting period that are recorded using CPOE Denominator: The number of laboratory orders created by the EP during the PI reporting period.

  - Numerator:

  - Denominator:

The Navigation bar at the bottom will monitor your progress.
### Stage 3 Objective 4 Measure 3 Computerized Provider Order Entry

#### Measure 3 Complete all required fields.

- **If you select the exclusions, you must upload documentation to support that separately.**

- **If you are not certain how to run the radiology orders using CPOE report, you may need to contact your CEHRT vendor.**

The Navigation bar at the bottom will monitor your progress.
Stage 3 Objective 5 Measure 1 Patient Electronic Access

**Objective Details:**

**Patient Electronic Access to Health Information - Measure 1 of 2:** The EP provides patients (or patient-authorized representatives) with timely electronic access to health information and patient-specific education.

**Measure Requirements:**

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 their or his/her health information.
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 CDRT.

**Additional Information:**

- To implement an API, the provider would need to host and enable the API functionality such that any application chosen by a patient would enable the patient to gain access to their individual health information provided that the application is configured to meet the technical specifications of the API. Providers may not prohibit patients from using any application, including third-party applications, which meet the technical specifications of the API, including the security requirements of the API. Providers are expected to provide patients with detailed instructions on how to authenticate their access through the API and provide the patient with supplemental information on available applications that leverage the API.

- Similar to how providers support patient access to VST capabilities, providers should continue to have identity verification processes to ensure that a patient using an application, which is leveraging the API, is provided access to their health information.

- In circumstances where there is no information available to patients or one of the fields previously noted, either because the EP can be excluded from recording such information or because there is no information to record (for example, no medication allergies or laboratory tests), the EP may have an indication that the information is not available and still meet the objective and its associated measures.

- The patient must be able to access this information on demand, such as through a patient portal or personal health record (PHR) or by other online or electronic means. We note that while a covered entity may be able to fully satisfy a patient’s request for information through VST, the measure does not replace the covered entity’s responsibilities to meet the broader requirements under HIPAA to provide an individual, upon request, with access to PHI in a designated record set.

- Providers should also be aware that while meaningful use is limited to the capabilities of CDRT to provide online access there may be patients who cannot access their EHRs electronically because of a disability. Providers who are not covered entities must provide individuals with disabilities equal access to information and appropriate auxiliary aids and services as provided in the applicable statutes and regulations.

- For Measure 1, providers must offer all four functionalities (view, download, transmit, and access through API) to their patients. And, patient health information needs to be made available to each patient in view, download, and transmit within 24 hours of the information being available to the provider for view and any time that information is generated within the first three months or three years.

- A patient who has multiple encounters during the PI reporting period, or even in subsequent PI reporting periods in future years, needs to be provided access for each encounter where they are seen by the EP.

- If a patient elects to opt out of participation, that patient must still be included in the denominator.

- If a patient elects to ‘opt out’ of participation, the provider may omit that patient in the numerator if the patient is provided all of the necessary information to subsequently access their information, obtain access through a patient-authorized representative, or otherwise opt back-in without further follow-up action required by the provider.

**Measure Description:**

The measure includes all encounters within the PI reporting period that is a full calendar year, or if it is less than a full calendar year, within the calendar year in which the PI reporting period occurs (between January 1st and December 31st).

**Measure Data Source:**

- Paper-based records are no longer allowed or required to be counted for measure 2 calculations. Providers may still provide paper-based educational materials for their patients, with patient (or authorized representative) accepting their health information online.

**TIP:**

- Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 5 Measure 2 Patient Electronic Access

Meaningful Use Objectives - Stage 3 for Program Year 2018 ePIP Measure 9 of 20 - CMS Meaningful Use Objective 5, Measure 2
Patient Electronic Access to Health Information - Measure 2 of 2

Objective Details:
Patient Electronic Access to Health Information - Measure 2 of 2: The EP provides patients (or patient-authorized representative) with timely electronic access to their health information and patient-specific education.

Measure Requirements:
The EP must use clinically relevant information from CEMHR 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 PI reporting period.

Additional Information:
To meet Stage 3 requirements for an PI reporting period in 2018, 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. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.

- To implement an API, the provider would need to fully enable the API functionality such that any application chosen by a patient would enable the patient to gain access to their individual health information provided that the application is configured to meet the technical specifications of the API. Providers may not prohibit patients from using any application, including third-party applications, which meet the technical specifications of the API, including the security requirements of the API. Providers are expected to provide patients with detailed instructions on how to authenticate their access through the API and provide the patient with supplemental information on available applications that leverage the API.
- Similar to how providers support patient access to VIT capabilities, providers should continue to have identity verification processes to ensure that a patient using an application, which is leveraging the API, is provided access to their health information.
- In circumstances where there is no information available to populate one or more of the fields previously listed, either because the EP can be excluded from recording such information electronically for reasons such as the EP is a limited service provider (for example, no medication allergies or laboratory tests), the EP may have an indication that the information is not available and still meet the objective and its associated measures.
- The patient must be able to access this information on demand, such as through a patient portal or personal health record (PHR) or by other online electronic means. We note that while an exclusion applies may be able to fully satisfy a patient’s request for information through VIT, the measure does not replace the covered entity’s responsibilities to meet the broader requirements under HIPAA to provide an individual, upon request, with access to PHI in a designated record set.
- Providers should also be aware that while meaningful use is limited to the capabilities of CEMHR to provide online access, there may be patients who cannot access their EHR electronically because of a disability. Providers who are covered by civil rights laws must provide individuals with disabilities equal access to information and appropriate auxiliary aids and services as provided in the applicable statutes and regulations.
- For Measure 1, providers must offer all four functionalities (view, download, transmit, and access through API) to their patients. And, patient health information needs to be be made available to each patient for view, download, and transmit within 48 hours of the information being available to the provider for each and every time that information is generated whether the patient has been “seen” for three months or for three years.
- A patient has multiple encounters during the PI reporting period, or even in subsequent PI reporting periods in future years, needs to be provided access for each encounter where they are seen by the EP.
- If a patient elects “not to participate” their patient must still be included in the denominator.
- If a patient elects “not to participate” the provider may count that patient in the numerator if the patient is provided all of the necessary information to subsequently access their information, obtain access through a patient authorized representative, or otherwise opt-out in further follow-up action required by the provider.
- For Measure 2, beginning in 2017, actions included in the numerator must occur within the PI 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 PI reporting period occurs (between January 1st and December 31st).
- Paper-based actions are no longer allowed or required to be counted for Measure 2 calculations. Providers may still provide paper-based educational materials for their patients, we are just no longer allowing them to be included in measure calculations.

Definition of terms:
- Application Programming Interface (API): A set of programming protocols established for multiple purposes. APIs may be created by a provider or provider organization to provide the patient with access to their health information through a third-party application with more flexibility than is often found in many current patient portals.
- Provide Access: When a patient possesses all of the necessary information needed to view, download, or transmit their information. This could include providing patients with instructions on how to access their health information, the websites address they must visit for online access, a unique and registered username or password, instructions on how to create a login, or any other instructions, tools, or materials that patients need in order to view, download, or transmit their information.
- View: The patient (or authorized representative) accessing their health information online.
- Download: The movement of information from online to physical electronic media.
- Transmit: This may be any means of electronic transmission according to any transport standard(s) (SMTP, FTP, REST, SOAP, etc.). However, the relocation of physical electronic media (for example, USB, CD) does not qualify as electronic transmission.
- Business Days: Business days are defined as Monday through Friday excluding federal or state holidays on which the EP or their respective administrative staffs are unavailable.
- Diagnoses: All data derived to diagnose and treat disease. Examples include, but are not limited to, blood tests, microbiology, urinalysis, pathology tests, radiology, cardiac imaging, nuclear medicine tests, and pulmonary function tests.
- Regulatory References:
- This objective may be found in Section 42 of the code of the federal register at 45 CFR 170.315(a)(1) and (b). For further discussion please see IBM IR D/S61866
- In order to meet this objective and measure, an EP must use the capabilities and standards of CEMHR at 45 CFR 170.315(a)(1) and (a)(2) and (a)(3)

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please review before attesting to this measure) For detailed information about the Patient Electronic Access objective, please click here

Note: (Please review before attesting to this measure) Further information about Patient Electronic Access objective can be found in the CMS Tip Sheet, please click here

Supporting Documentation Requirements:
Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attest process.

- Red asterisk indicates a required field
- Gray asterisk indicates a conditionally required field

Measure Entry:
- Exclusion: An EP may exclude from the measure if they have no office visits during the PI reporting period.
- Does this exclusion apply to you? Yes No
- This exclusion applies to patients who:
- Do not complete the following information:
- Do not have at least one EP visit during the PI reporting period
- The number of patients in the denominator (or patient-authorized representative) who view, download, or transmit to a third party their health information
- Denominator: Number of unique patients seen by the EP during the PI reporting period
- Yes No

For additional information, please visit: https://www.azepip.gov/
Stage 3 Objective 6 Measure 1 Coordination of Care

Measure Requirements:
For an PM reporting period in 2018, 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.

1. View, download or transmit to a third party their health information.
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 EHR.

3. A combination of 1 and 2.

Additional Information:
To meet Stage 3 requirements for an IT reporting period in 2018, all providers must use technology certified to the 2015 Edition. A provider who has technology certified to a cut-off date before the 2015 Edition and 2016 Edition may postpone adoption to the Stage 3 measures. If the cut-off date certified technology would not meet the Stage 3 measures. A provider who has technology certified to the 2014 Edition may not delay to Stage 3.

For the numerators for Measures 1 and 2, beginning in 2019, the activity should occur within the IT 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 IT reporting period occurs (between January 1st and December 31st).

Providers must attest to all these measures and must meet the requirements for at least two measures to meet the objective.

There are six actions a patient might take as part of Measure 1:
1. Name and contact information
2. Download their information
3. Transact their information to a third party
4. Access their information through an API
These actions may overlap, but a provider is able to count any actions in the single numerator. Therefore, for the first measure, a provider may meet a combined threshold for VST and API actions, or if their technology functions overlap, then any view, download, transmit, or API actions taken by the patient using CHS-IT should count toward the threshold.

- In order to meet the objectives, the following information must be available within 4 business days of the information being made available to the EP:
  - Patient name
  - Provider name and office contact information
  - Current and past problem list
  - Procedures
  - Laboratory test results
  - Current medication list and medication history
  - Current smoking status and tobacco history
  - Vital signs (height, weight, blood pressure, BMI, growth charts)
  - Allergies
  - Demographic information (preferred language, sex, race, ethnicity, date of birth)
  - Care plan, including goals and instructions

Any known care team members, including the primary care provider (PCP) of record

- An EP can only access additional information and uploads with the information that is known to the care team.

- Measure 1 includes provider-initiated communications (when a provider sends a message to a patient or the patient’s authorized representative), and provider-to-provider communications only if the patient is included. A provider can only count messages in the numerator when the provider participates in the communication (e.g. any patient-initiated communication only if the provider responds to the patient. Note: Providers are not required to respond to every message received if no response is necessary.

- For Measure 2, the forms of data that would satisfy the measure are broad. It may include, but is not limited to, social service data, data generated by a patient or a patient’s authorized representative, electronic or paper documents, health-related data from the use of mobile applications for tracking health and nutrition, home healthcare records, records of clinical care received, and other forms of data that provide the basis for creating and maintaining the patient record.

TIP: Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

The navigation bar at the bottom will monitor your progress.

Click the hyperlink on the ePIP screen to learn more about this requirement.

Stage 3 Screen 10
Coordination of Care

Measure 1
Complete all required fields.

If you select the exclusions, you upload documentation to support that separately.
Stage 3 Objective 6 Measure 2 Coordination of Care

Meaningful Use Objectives - Stage 3 for Program Year 2018

ePIP Measure 11 of 20: CMS Meaningful Use Objective 6, Measure 2
Coordination of Care through Patient Engagement - Measure 2 of 3

Objective Details:
Coordination of Care through Patient Engagement - Measure 2 of 3: Use CEHRT to engage with patients or their authorized representatives about the patient's care.

Measure Requirements:
For an PI reporting period in 2018, more than 5 percent of all unique patients seen by the EHR during the PI 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.

Additional Information:
- To meet the measure requirements for an PI reporting period in 2018, all providers must use technology certified to the 2015 Edition. A provider who has technology certified to a combination of the 2015 Editions and 2014 Edition may demonstrate compliance to the Stage 3 requirements. If the mix of certified technologies would prohibit them from meeting the measure requirements, then the provider must demonstrate compliance to the Stage 2 requirements.
- For the numerator for measures 1 and 2, beginning in 2017, the action must occur within the PI 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 measure is reported.
- Providers must attest to all three measures and must attest the thresholds for at least two measures to meet the objective.
- There are four actions a patient might take as part of Measure 1:
  1. View their information
  2. Answer their information
  3. Access their information through an API
  4. Upload their information to a third party
- To meet the measure requirements, providers must demonstrate that the required actions are taking place for at least two unique patients seen by the PI reporting period. The data collected for this objective may be used to inform the Measurement System for the 2020 PI Reporting Period Challenge.
- These actions may overlap, but a provider is allowed to count any and all actions in the single numerator. Therefore, for the first measure, a provider may meet a combined threshold from Stage 2 and 3 actions, or if their technology function creates, then any view, download, transmit, or API actions taken by the patient using CEHRT would count toward the threshold.
- In order to meet the objective, the following information must be available within 4 business days of the information being made available to the PI.
  - Patient name
  - Provider's name and title or role
  - Contact information (phone)
  - Current and past problem list
  - Allergies
  - Laboratory test results
  - Immunization record and vaccination history
  - Current immunization allergy list and medication allergy history
  - Vital signs (height, weight, blood pressure, BMI growth charts)
  - Smoking status
  - Demographic information (preferred language, sex, race, ethnicity, date of birth)
  - Care plan items( ), including goals and medications

- Any electronic exchange of patient care messages including the primary care provider (PCP) or resource

- An eIP can make available additional information and still align with the objective.
- Measure 2 includes provider-initiated communications (when a provider sends a message to a patient or the patient’s authorized representative), and provider-to-provider communications if the patient is included. A provider can only count messages in the numerator when the provider's part B in the communication is an any patient-initiated communication, regardless of who initiated the message.
- For Measure 3, the types of data that would satisfy the measure are broad. It may include, but is not limited to, social service data, generated by a patient or a patient’s authorized representative, advance directive, insurance data, race and ethnicity data, social security data, race and ethnicity data, social security data, resource health monitoring data, and disease monitor data. In addition, the sources of data vary and may include mobile applications for tracking health and nutrition, health devices with tracking capabilities such as activity trackers or blood pressure monitors, wearable devices such as activity trackers, or patient-generated health data, and other methods or regular sources generated health data.
- For the Patient navigation Health data measure, the data may not be identifiable: the patient providers to the PI or care location during the office visit as such data does not meet the patient identifiable criteria to warrant care coordination and patient engagement in a wide range of settings outside the provider’s immediate scope of practice.
- For Measure 3, we do not specify the manner in which providers are required to incorporate the data. Providers may work with their EHR developers to establish the methods and processes that best fit their practice and needs. For example, if data provided can be easily inserted into a structured format (as in an existing field within the EHR, such as a C-CDUA or care provider) or care provider data appears as data that is entered or selected from within the EHR (such as a D-CDUA or care provider data appears as a data that is entered or selected from within the EHR).

- For the Patient navigation Health data measure, the data may not be identifiable: the patient providers to the PI or care location during the office visit as such data does not meet the patient identifiable criteria to warrant care coordination and patient engagement in a wide range of settings outside the provider’s immediate scope of practice.

Applications (relevant components): A set of pre-existing protocols established for multiple purposes. APIs may be enabled by a provider or provider organization to provide the patient access to their health information through a third-party application that meets privacy and security requirements.

- View: The patient or authorized representative accessing their health information online.
- Download: The movement of information from servers to personal electronic media.
- Transmission: This may be any means of electronic transmission according to any transport standard(s) (CMTP, FTP, HTTP, ROAP, etc.). However, the inclusion of physical transmission (for example, LHI, CO) does not qualify as transmission.

- Patient Generated Health data - Data generated by a patient or their authorized representative, includes any source of health data, data generated by a patient or a patient’s authorized representative, advance directives, patient medical device data, patient health monitoring data, and wellness monitor data.

- Secure Messaging: Any electronic communication between a provider and patient that ensures only those parties can access the communication. This electronic message could be sent via the electronic messaging function of an ePIP, an online patient portal or any other electronic means.

- Unique Patient: A patient is a person known to be an EHR (even if there is no IP report for them). For purposes of measurement, patient is only counted once in the denominator for the measure. All the measures comparing on the term “unique patient” relate to what is contained in the patient’s record. Some of this information may be used to identify, or be used for purposes of measurement, the provider as well as patient encounters. This is especially true for providers who use electronic health records that are part of the same EHR reporting period.

- Regulatory References:
  - This objective may be found in Section 42 of the code of the federal register at 45F 24.68(a)(4)(vii) and (viii). For further discussion please see 80 FR 62851
  - The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular measure.

For detailed information about the Coordination of Care through Patient Engagement objective, please click here.

Supporting Documentation Requirements:
Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attestation process.

- (*) Asterisk indicates a required field
- (C) Gray asterisk indicates a conditionally required field

Exclusions:
- Any IP may exclude from the measure if they have no office visits during the PI reporting period.

- Does this exclusion apply to you?
  - No
  - Yes

Exclusion: Any IP 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 PI reporting period may exclude the measure.

- Does this exclusion apply to you?
  - No
  - Yes

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 PI reporting period.

- Number of unique patients seen by the PI during the PI reporting period.

- Denominator:

Meaningful Use Objective: Navigation

<table>
<thead>
<tr>
<th>Measure</th>
<th>Short Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
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<td>2</td>
<td>Coordination of Care</td>
<td>Complete all required fields.</td>
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</table>

The Navigation bar at the bottom will monitor your progress.

TIP: Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 6 Measure 3 Coordination of Care

Meaningful Use Objectives - Stage 3 for Program Year 2018
ePIP Measure 12 of 20 - CMS Meaningful Use Objective 6 Measure 3 Coordination of Care through Patient Engagement - Measure 3 of 3

Objectives: Coordination of Care through Patient Engagement - Measure 3 of 3: Use CEHRT to engage with patients or their authorized representatives about the patient's care.

Measure Requirements: 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 PI reporting period.

Additional Information:
• For the Stage 3 requirements for an EP reporting period in 2018, all providers must use technology certified to the 2015 Edition, a provider who has technology certified to a combination of the 2014 Edition and 2015 Edition may potentially attest to the Stage 3 requirements, if the mix of certified technologies would not prohibit them from meeting the Stage 3 measure. A provider who has technology certified to the 2014 Edition may not attest to Stage 3.
• For the numerator for measures 1 and 2, beginning in 2017, the action must occur within the PI reporting period if that is a full calendar year, or if it is less than a full calendar year, the period in which the reporting period occurs (between January 1 and December 31) is used.
• Providers must attest to all three measures and must meet the thresholds for at least two measures to meet the objective.
• The measure is applicable if they have done at least 200 completed attestation entries. If a provider has not met the 200 required entries, they cannot attest to this measure.
• The measure includes the CEHRT and the CEHRT must be open for the entire 30 minute encounter.
• The CEHRT must have the ability to view and update the health information entered by the patient or their authorized representative.
• The CEHRT must have the ability to view and update the health information entered by the patient or their authorized representative.
Stage 3 Objective 7 Measure 1 – 3 Health Information Exchange

Additional Information:

- To meet Stage 3 requirements, all providers must use technology certified to the 2015 Edition for the Health Information Exchange objective.
- For Measure 1 and 3, providers may continue to limit the denominator to those patients whose records are maintained using CEHRT for measures with a denominator other than unique patients seen by the EP during the PI reporting period.
- For Measure 1, beginning in 2017, in order to count in the numerator, the exchange must occur within the PI 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 PI reporting period occurs.
- For Measure 1, the referring provider must have reasonable certainty of receipt by the receiving provider to count the action toward the measure. A provider must have this may include confirmation of receipt or that a query of the summary of care record has occurred in order to count the action in the numerator.
- Apart from the three fields noted as required for the summary of care record (i.e., current problem list, current medication list, and current medication allergy list), in circumstances where there is no information available to populate one or more of the fields listed (because the EP does not record such information or because there is no information to record), the EP may leave the field(s) blank and still meet the objective and its associated measure.
- A provider must have the ability to transmit all data pertaining to laboratory test results in the summary of care document, but may work with their system developer to establish clinically relevant parameters for the most appropriate results for the given transition or referral.
- A provider who limits the transmission of laboratory test result data in a summary of care document must send the full results upon request (i.e. all lab results as opposed to a subset).
- The exchange must comply with the privacy and security protocols for ePHI under HIPAA.
- In cases where the providers share access to an EHR, a transition or referral may still count toward the measure if the referring provider creates the summary of care document using CEHRT and sends the summary of care document electronically. If a provider chooses to include such transitions to providers where access to the EHR is shared, they must do so universally for all patient and all transitions or referrals.
- For Measure 1, the initiating provider must send a C-CDA document that the receiving provider would be capable of electronically incorporating as a C-CDA on the receiving end. In other words, a provider sends a C-CDA and the receiving provider converts the C-CDA into a PDF or a fax or some other format. The sending provider may still count the transition or referral in the numerator. If the sending provider converts the file to a format the receiving provider could not electronically receive and incorporate as a C-CDA, the initiating provider may not count the transition in their numerator.
- For the purposes of defining the cases in the denominator for Measure 2, we stated that what constitutes “unavailable” and, therefore, may be excluded from the denominator, will be that a provider:
  - Requested an electronic summary of care record to be sent and did not receive an electronic summary of care document, and
  - The provider either:
    - Queried at least one external source via HEI functionality and did not locate a summary of care for the patient, or the provider does not have access to HEI functionality to support such a query, or
    - Confirmed that HEI functionality supporting query for summary of care documents was not operational in the provider's geographic region and not available within the provider's EHR network as of the start of the PI reporting period.
- For Measure 2, a record cannot be considered to be incorporated if it is discarded without the reconciliation of clinical information or if it is stored in a manner that is not accessible for provider use within the EHR.
- For Measure 3, the process may include both automated and manual reconciliation to allow the receiving provider to work with both the electronic data provided with any necessary review, and to work directly with the patient to reconcile their health information.
- For Measure 3, if no update is necessary, the process of reconciliation may consist of simply verifying that fact or reviewing a record received on referral and determining that such information is merely duplicative of existing information in the patient record.
- Non-medical staff may conduct reconciliation under the direction of the provider so long as the provider or other credentialed medical staff is responsible and accountable for reviewing the information and for the assessment of and action on any relevant CDR.

Definitions of Terms

Transition of Care: The movement of a patient from one setting of care (hospital, ambulatory primary care practice, ambulatory specialty care practice, long-term care, home health, rehabilitation facility) to another. At a minimum this includes all transitions of care and referrals that are ordered by the EP.

Summary of Care Record: All summary of care documents used to meet this objective must include the following information if the provider knows it:

- Patient name
- Referring or transitioning provider’s name and office contact information (EP only)
- Procedures
- Encounter diagnosis
- Immunizations
- Laboratory test results
- Vital signs (height, weight, blood pressure, BMI)
- Smoking status
- Functional status, including activities of daily living, cognitive and disability status
- Demographic information (preferred language, sex, race, ethnicity, date of birth)
- Care plan, including goals and instructions
- Care team including the primary care provider of record and any additional known care team members beyond the referring or transitioning provider and the receiving provider
- Reason for referral (EP only)
- Current problem list (Providers may also include historical problems at their discretion)*
- Current medication list
- Current medication allergy list

*Note: An EP must verify that the fields for current problem list, current medication list, and current medication allergy list are not blank and include the most recent information known by the EP as of the time of generating the summary of care document or include a notation of no current problem, medication, and/or medication allergies.

Current problem lists - At a minimum a list of current and active diagnoses.

Active/current medication list - A list of medications that a given patient is currently taking.

Active/current medication allergy list - A list of medications to which a given patient has known allergies.

Allergy - An exaggerated immune response or reaction to substances that are generally not harmful.

Care Plan - The structure used to define the management actions for the various conditions, problems, or issues. A care plan must include at a minimum the following components: problem (the focus of the care plan), goal (the target outcome) and any instructions that the provider has given to the patient. A goal is a defined target or measure to be achieved in the process of patient care (an expected outcome).
Stage 3 Objective 7 Measure 1 Health Information Exchange

Objective Details:
Health Information Exchange - Measure 1 of 3: The EP provides a summary of care record when transitioning or referring their patient to another setting of care, receives or retrieves a summary of care record upon the receipt of a transition or referral or upon the first patient encounter with a new provider, and incorporates summaries of care information from other providers into their EHR using the functions of CEHRT.

Measure Requirements:
For more than 50 percent of transitions of care and referrals, the EP that transitions or refers their patient to another setting of care provider or care of provider, creates a summary of care record using CEHRT and electronically exchanges the summary of care record.

Regulatory References:
- This objective may be found in Section 49 of the code of the federal register at 493.24 (d)(7)(A) and (B). For further discussion please see 80 FR 62601
- In order to meet this objective and measure, an EP must use the capabilities and standards of CEHRT at 493.24 (d)(7)(A) through (d)(2) and (d)(6) through (d)(8).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure). For more information regarding the Health Information Exchange objective, please click here

Supporting Documentation Requirements:
Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHRR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attestation process.

(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

Measure Entry:
Exclusion: Any EP who transfers a patient to another setting or refers a patient to another provider less than 100 times during the PI reporting period.
Does this exclusion apply to you?
○ Yes ○ No
Exclusion: 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 4Mpbs broadband availability according to the latest information available from the FCC on the first day of the PI reporting period may exclude the measures.

Does this exclusion apply to you?
○ Yes ○ No

*PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology
○ This data was extracted from both paper records as well as records maintained using Certified EHR Technology (CEHRT).
○ This data was extracted only from patient records maintained using certified EHR technology.

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 CEHRT and exchanged electronically.
Denominator: Number of transitions of care and referrals during the PI reporting period for which the EP was the transferring or referring provider.

* Numerator:

* Denominator:

Meaningful Use Objectives - Navigation
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Meaningful Use Objective Summary

TIP
Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 7 Measure 2 Health Information Exchange

Health Information Exchange - Measure 2 of 3

Objective Details:
Health Information Exchange - Measure 2 of 3: The EP provides a summary of care record when transitioning or referring their patient to another setting of care, receives or retrieves a summary of care record upon the 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.

Measure Requirements:
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.

Regulatory References:
- This objective may be found in Section 42 of the code of the federal register at 45FR 620811 and (B). For further discussion please see 80 FR 620811
- In order to meet this objective and measure, an EP must use the capabilities and standards of CEHRT at 45 CFR 170.315 (b)(1) through (b)(3) and (b)(6) through (b)(8).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

For detailed information about the Health Information Exchange objective, please click here

Supporting Documentation Requirements:
Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the “Attestation Progress” page as a required step in the attestation process.

(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

Measure Entry:
Exclusion: 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 PI reporting period is excluded from this measure.
- Does this exclusion apply to you?
  - Yes
  - No

Exclusion: 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 4 Mbps broadband availability according to the latest information available from the FCC on the first day of the PI reporting period may exclude the measures.
- Does this exclusion apply to you?
  - Yes
  - No

PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology
- This data was extracted from both paper records as well as records maintained using Certified EHR Technology (CEHRT).
- This data was extracted only from paper records maintained using certified EHR technology.

Complete the following information:
Numerator: Number of patient encounters in the numerator 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 PI 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.

Numerator:

Denominator:

Meaningful Use Objectives - Navigation
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18

TIP
Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.
Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 7 Measure 3 Health Information Exchange

Meaningful Use Objectives - Stage 3 for Program Year 2018

ePIP Measure 15 of 20 - CMS Meaningful Use Objective 7, Measure 3
Health Information Exchange - Measure 3 of 3

Objective Details:
Health Information Exchange - Measure 3 of 3: The EP provides a summary of care record when transitioning or referring their patients to another setting of care, receives or retrieves a summary of care record upon the 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 CEHR.

Measure Requirements:
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.

Regulatory References:
- This objective may be found in Section 42 of the code of the federal register at 45CFR 170.315(b)(1) through (b)(8) and (a)(8) through (a)(8).
- In order to meet this objective and measure, an EP must use the capabilities and standards of CEHR at 45 CFR 170.315(b)(1) through (b)(6) and (a)(8) through (a)(8).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

For detailed information about the Health Information Exchange objective, please click here

Note: (Please Review before attesting to this measure): For more information regarding the Health Information Exchange objective, please click here

Supporting Documentation Requirements:
Meaningful Use Objective Measures require supporting documentation. The supporting documentation for this measure should be included in your Meaningful Use EHR Report. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

Measure Entry:
Exclusion: 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 PI reporting period is excluded from this measure.

* Does this exclusion apply to you?
   - [ ] Yes   [ ] No

PATIENT RECORDS: Please select whether the data used to support this measure was extracted from ALL patient records or only from patient records maintained using certified EHR technology
   - [ ] This data was extracted from both paper records as well as records maintained using Certified EHR Technology (CEHR).
   - [ ] This data was extracted only from patient records maintained using certified EHR Technology.

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 PI reporting period for which the EP was the recipient of the transition or referral or had never before encountered the patient.

* Numerator:

* Denominator:

Meaningful Use Objectives - Navigation

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 8 Measure 1 – 5 Public Health and Clinical Data Registry Reporting

**Additional Information:**

- To meet all the measures within the public health objective, EPs must use CEHRT and the standards included in the [2015 Edition](https://www.azepip.gov/) proposed rule. CMS anticipates that as new public health registries and clinical data registries are created, ONC and CMB will work with the public health community and clinical specialty societies to develop ONC-certified electronic reporting standards for those registries so providers have the option to count participation in those registries under the measures for this objective.
- EPs must attest to at least two measures from the Public Health Reporting Objective, Measures 1 through 5.
- If public health agencies have not declared 6 months before the start of the PM reporting period whether the registry they are offering will be ready on January 1 of the upcoming year for use by providers seeking to meet PM reporting periods in that upcoming year, a provider can claim an exclusion.
- An exclusion for a measure does not count toward the total of two measures. Instead, in order to meet this objective, an EP would need to meet two of the total number of measures available to them. If the EP qualifies for multiple exclusions and the remaining number of measures available to the EP is less than two, the EP can meet the objective by meeting all of the remaining measures available to them and claiming the applicable exclusions. Available measures include ones for which the EP does not qualify for an exclusion.
- For Measure 1, provider’s health IT system may layer additional information on the immunization history, forecast, and still successfully meet this measure.
- Bi-directionally provides that certified health IT must be able to receive and display a consolidated immunization history and forecast in addition to sending the immunization record.
- For Measure 1, an exclusion does not apply if an entity designated by the immunization registry or immunization information system can receive electronic immunization data submissions. For example, if the immunization registry cannot accept the data directly or in the standards required by CEHRT, but it has designated a Health Information Exchange to do so on their behalf and the Health Information Exchange is capable of accepting the information in the standards required by CEHRT, the provider could not claim the second exclusion.
- For Measure 2, because syndromic surveillance reporting is more appropriate for urgent care settings and eligible hospitals, we removed this measure for eligible professionals for Stage 3 with the exception of providers who are practicing in urgent care settings. Note: some states have chosen to waive the urgent care setting requirement. Please contact your state Medicaid agency for more information.
- For Measure 2, an exclusion does not apply if an entity designated by public health agency can receive electronic syndromic surveillance data submissions. For example, if the public health agency cannot accept the data directly or in the standards required by CEHRT, but it has designated a Health Information Exchange to do so on their behalf and the Health Information Exchange is capable of accepting the information in the standards required by CEHRT, the provider could not claim the second exclusion.
- Measure 3, Electronic Case Reporting is not required until 2019, since we believe that the standards will be mature and that jurisdictions will be able to accept these types of data by that time.
- For Measure 4, EPs may choose to report more than one public health registry to meet the number of measures required to meet the objective.
- For Measure 4, a provider may count a specialized registry (such as a prescription drug monitoring) if the provider achieved the phase of active engagement defined under Active Engagement Option 3: Production, including production data submission with the specialized registry, in a prior year under the applicable requirements of the EHR Incentive Programs for that year.
- For Measure 5, EPs may choose to report more than one clinical data registry to meet the number of measures required to meet the objective.
- For Measure 5, the definition of jurisdiction is general, and the scope may be local, state, regional or at the national level. The definition will be dependent on the type of registry to which the provider is reporting. A registry that is “bureaucratic” would be considered a registry at the national level and would be included for purposes of this measure.
- Providers who have previously registered, tested, or began ongoing submission of data to registry do not need to test/perform the process beginning at active engagement option 1.
- The provider may simply attest to the active engagement option which most closely reflects their current status.
- In determining whether an EP meets the first exclusion, the registries in question are those sponsored by the public health agencies with jurisdiction over the area where the EP practices and national medical societies covering the EP’s scope of practice. Therefore, an EP must complete two actions in order to determine available registries or claim an exclusion.
  - Determine if the jurisdiction (state, territory, etc.) endorses or sponsors a registry; and,
  - Determine if a National Specialty Society or other specialty society with which the provider is affiliated endorses or sponsors a registry.
- If a provider is part of a group which submits data to a registry, but the provider does not contribute to that data (for example they do not administer immunizations), the provider should not attest to meeting the measure but instead should select the exclusion. The provider may then select a different more relevant measure to meet.
- If a provider does the actions that results in a data element for a registry in the normal course of their practice and is in active engagement to submit a registry, but simply has no cases for the reporting period, the provider is not required to take the exclusion and may attest to meeting the measure.
- CMB has published a central repository for public health agency (PHA) and clinical data registry (CDR) reporting. That central repository is available at [https://www.cms.gov/Regulations-and-Guidance/Legislation/EHRIncentivePrograms/CentralizedRepository.html](https://www.cms.gov/Regulations-and-Guidance/Legislation/EHRIncentivePrograms/CentralizedRepository.html)

**Definition of Terms:**

**Active engagement** means that the provider is in a process of moving towards sending “production data” to a public health agency or clinical data registry, or is sending production data to a public health agency or clinical data registry.

**Active Engagement Option 1 - Completed Registration to Submit Data:** The EP registered to submit data with the PHA or, where applicable, the CDR to which the Information is being submitted; registration was completed within 45 days after the start of the PM reporting period; and the EP is awaiting an invitation from the PHA or CDR 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 PM reporting period.

**Active Engagement 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 CDR within 30 days; failure to respond twice within an PM reporting period will result in the provider not meeting the measure.

**Active Engagement Option 3 - Production:** The EP has completed testing and validation of the electronic submission and is electronically submitting production data to the PHA or CDR.

**Production data** refers to data generated through clinical processes involving patient care, and it is used to distinguish between data and “test data” which may be submitted for the purposes of enrolling in and testing electronic data transfers.

**Tip:** Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report. Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 8 Measure 1 Public Health and Clinical Data Registry Reporting

Measure Requirements:
The EP is in active engagement with a public health agency or clinical data registry 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.

Supporting Documentation Requirements:

The Public Health Objective Measures require supporting documentation to be uploaded. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

Please provide supporting documentation outlining your active engagement with the Immunization Registry. If you are choosing one of the available exclusions please provide documentation to support your exclusion choice.

(*) Red asterisk indicates a required field
(°) Gray asterisk indicates a conditionally required field

TIP:
Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 8 Measure 2 Public Health and Clinical Data Registry Reporting

Meaningful Use Objectives - Stage 3 for Program Year 2018

Measure 17 of 20 - CMS Meaningful Use Objective 8, Measure 2

Public Health and Clinical Data Registry Reporting - Measure 2 of 5

Objective Details:

Public Health and Clinical Data Registry Reporting - Measure 2 of 5: The EP is in active engagement with a public health agency or clinical data registry to submit electronic public health data is a meaningful way using certified EHR technology, except where prohibited, and in accordance with applicable law and practice.

Measure Requirements:

The EP is in active engagement with a public health agency to submit syndromic surveillance data from an urgent care setting.

Regulatory References:

- This objective may be found in Section 43 of the code of the federal register at 49§ 34 (d)(B)(A) and (B). For further discussion please see 80 FR 62979.
- In order to meet this objective and measure, an EP must use the capabilities and standards of CCHRT at 45 CFR 170.315 (g)(1), (g)(2), (g)(4), (g)(5), (g)(6) and (g)(7).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

For detailed information about the Public Health and Clinical Data Registry Reporting objective, please click here

Note: (Please Review before attesting to this measure): For more information regarding the Public Health Reporting objective, please click here

Supporting Documentation Requirements:

The Public Health Objective Measures require supporting documentation to be uploaded. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

Please provide supporting documentation outlining your active engagement with the Syndromic Surveillance Registry. If you are choosing one of the available exclusions please provide documentation to support your exclusion choice.

(*) Red asterisk indicates a required field
(+) Gray asterisk indicates a conditionally required field

Measure Entry:

Exclusion: Is not in a category of providers from which ambulatory syndromic surveillance data is collected by their jurisdiction’s syndromic surveillance system.

* Does this exclusion apply to you?
  - Yes ☐ No ☑

Exclusion: 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 CCHRT definition at the start of the PI reporting period.

* Does this exclusion apply to you?
  - Yes ☐ No ☑

Exclusion: 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 PI reporting period.

* Does this exclusion apply to you?
  - Yes ☑ No ☐

Complete the following information:

* Are you in active engagement with a public health agency to submit syndromic surveillance data?
  - Yes ☐ No ☑

TIP:

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.

Stage 3 Screen 17

Public Health and Clinical Data Registry Reporting

☑ Measure 2

Complete all required fields.

If you select the exclusions, you must upload documentation to support that separately.

If you are in active engagement to submit immunization data to a public health agency, you must upload documentation to support that separately.

The Navigation bar at the bottom will monitor your progress.

Meaningful Use Objectives - Navigation

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Stage 3 Objective 8 Measure 3 Public Health and Clinical Data Registry Reporting

Meaningful Use Objectives - Stage 3 for Program Year 2018
ePIP Measure 18 of 20: CMS Meaningful Use Objective 8, Measure 3
Public Health and Clinical Data Registry Reporting - Measure 3 of 5

Objective Details:
Public Health and Clinical Data Registry Reporting - Measure 3 of 5: The EP is in active engagement with public health agency or clinical data registry 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.

Measure Requirements:
The EP is in active engagement with a public health agency to submit case reporting of reportable conditions.

Regulatory References:
- This objective may be found in Section 42 of the code of the federal register at 45FR24 (d)(8)(A) and (B). For further discussion please see 80 FR 62070
- In order to meet this objective and measure, an EP must use the capabilities and standards of CHERT at 45CFR 170.315 (I)(1), (I)(2), (I)(4), (I)(5), (I)(6) and (I)(7).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)
For detailed information about the Public Health and Clinical Data Registry Reporting objective, please click here

(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

Measure Entry:

CMS has indicated that Stage 3, Objective 8, Measure 3 is not required for Program Year 2018. You are not required to attest to this measure in 2018. Please click Save & Continue below to go to the next measure.

Meaningful Use Objectives - Navigation
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
Meaningful Use Objectives Summary

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Stage 3 Objective 8 Measure 4 Public Health and Clinical Data Registry Reporting

**Objective Details:**

Public Health and Clinical Data Registry Reporting - Measure 4 of 5: The EP is in active engagement with a public health agency or clinical data registry 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.

**Measure Requirements:**

The EP is in active engagement with a public health agency to submit data to public health registries.

**Regulatory References:**

- This objective may be found in Section 42 of the code of the federal register at 495.24 (d)(8)(1)(A) and (B). For further discussion please see 80 FR 63670.
- In order to meet this objective and measure, an EP must use the capabilities and standards of CDR/RT at 45 CFR 170.315 (f)(7), (f)(2), (f)(4), (f)(5), (f)(6) and (f)(7).

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

For detailed information about the Public Health and Clinical Data Registry Reporting objective, please click here

Note: (Please Review before attesting to this measure): For more information regarding the Public Health Reporting objective, please click here

**Supporting Documentation Requirements:**

The Public Health Objective Measures require supporting documentation to be uploaded. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

Please provide supporting documentation outlining your active engagement with the Public Health Registry. If you are choosing one of the available exclusions please provide documentation to support your exclusion choice.

(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

**Measure Entry:**

Exclusion: Does not diagnose or directly treat any disease or condition associated with a public health registry in their jurisdiction during the PI reporting period

- Does this exclusion apply to you?
  - Yes
  - No

Exclusion: 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 CDR/RT definition at the start of the PI reporting period.

- Does this exclusion apply to you?
  - Yes
  - No

Exclusion: 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 PI reporting period.

- Does this exclusion apply to you?
  - Yes
  - No

**TIP:**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page.

Click the hyperlink on the ePIP screen to learn more about this requirement.

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Stage 3 Objective 8 Measure 5 Public Health and Clinical Data Registry Reporting

Meaningful Use Objectives - Stage 3 for Program Year 2018
ePIP Measure 20 of 20 - CMS Meaningful Use Objective 8, Measure 5
Public Health and Clinical Data Registry Reporting - Measure 5 of 5

Objective Details:
Public Health and Clinical Data Registry Reporting - Measure 5 of 5: The EP is in active engagement with a public health agency or clinical data registry 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.

Measure Requirements:
The EP is in active engagement to submit data to a clinical data registry.

Regulatory References:
• This objective may be found in Section 42 of the code of the federal register at 495.24 (d)(3)(i)(A) and (B). For further discussion please see 80 FR 62870
• In order to meet this objective and measure, an EP must use the capabilities and standards of CHERT at 45 CFR 170.315 (f)(1), (f)(2), (f)(4), (f)(5), (f)(6) and (f)(7)

The Centers for Medicare and Medicaid Services (CMS) provides documentation to guide you through the measure requirements for this particular objective. (Please Review before attesting to this measure)

For detailed information about the Public Health and Clinical Data Registry Reporting objective, please click here
Note: (Please Review before attesting to this measure): For more information regarding the Public Health Reporting objective, please click here

Supporting Documentation Requirements:
The Public Health Objective Measures require supporting documentation to be uploaded. The link for uploading this documentation will appear on the "Attestation Progress" page as a required step in the attestation process.

Please provide supporting documentation outlining your active engagement with the Clinical Data Registry. If you are choosing one of the available exclusions please provide documentation to support your exclusion choice.
(*) Red asterisk indicates a required field
(*) Gray asterisk indicates a conditionally required field

Measure Entry:

Exclusion: Does not diagnose or directly treat any disease or condition associated with a clinical data registry in their jurisdiction during the PI reporting period.
• Does this exclusion apply to you?
  ○ Yes ○ No

Exclusion: 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 CHERT definition at the start of the PI reporting period.
• Does this exclusion apply to you?
  ○ Yes ○ No

Exclusion: Operates in a jurisdiction where no clinical 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 PI reporting period.
• Does this exclusion apply to you?
  ○ Yes ○ No

Complete the following information:
• Are you in active engagement with a public health agency to submit data to a clinical data registry?
  ○ Yes ○ No

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.
Click the hyperlink on the ePIP screen to learn more about this requirement.

The Navigation bar at the bottom will monitor your progress.
Attestation Progress (After Objective Measures)

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Program Year Notes</th>
<th>Program Year Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>The data required for this attestation is grouped into categories. In order to complete your attestation, you must complete ALL of the tasks listed below. Click on the <code>Begin</code> button to start performing a given step. If a step has been started, but not completed click on the <code>Continue</code> button to finish a step. Once a step is finished you can click on the <code>Modify</code> button to change any information that was previously entered.</td>
<td>The AHCCCS Preserving Interoperability Program is currently open for Program Year 2018.</td>
<td>We encourage all providers to review the CMS documentation for Program Year 2018 before attesting. These documents are available at the CMS EHR website, or in ePIP in the ePIP Document Library.</td>
</tr>
</tbody>
</table>

When you complete a step and the status has changed from “Begin” to “Modify”, you can close the program and it will automatically save your work.

You can return later and modify previous steps in this section.

Click the `Begin` button to complete each step.

Click the `Continue` button to finish a step.

Click the `Modify` button to change information previously entered.
Clinical Quality Measures

<table>
<thead>
<tr>
<th>National Quality Strategy (NQS) Domains</th>
<th>Number CQMs Available</th>
</tr>
</thead>
<tbody>
<tr>
<td>1  Person and Caregiver-Centered Experience and Outcomes</td>
<td>4</td>
</tr>
<tr>
<td>2  Patient Safety</td>
<td>5</td>
</tr>
<tr>
<td>3  Communication and Care Coordination</td>
<td>2</td>
</tr>
<tr>
<td>4  Community/Population Health</td>
<td>11</td>
</tr>
<tr>
<td>5  Efficiency and Cost Reduction</td>
<td>4</td>
</tr>
<tr>
<td>6  Effective Clinical Care</td>
<td>29</td>
</tr>
</tbody>
</table>

Clinical Quality Measures (CQMs) Selection:

Providers are required to report on 6 of 55 separate CQMs from any of the National Quality Strategy domains.

Select the CQMs that best apply to your scope of practice.

The CQM Reporting Period is a 90-day period selected from 2018.

If your certified EHR technology does not contain patient data for at least 6 CQMs:

☑ Report the CQMs for which there is patient data

☑ Report the remaining required CQMs as "zero denominators" as displayed by your certified EHR technology.

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Clinical Quality Measures for Person and Caregiver-Centered Experience & Outcomes

<table>
<thead>
<tr>
<th>Objective</th>
<th>Measure</th>
<th>Selected</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS 157v6 \ NQF 0384 - Oncology: Medical and Radiation – Pain Intensity Quantified</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>☐</td>
</tr>
<tr>
<td>CMS 66v6 - Functional Status Assessment for Total Knee Replacement</td>
<td>Percentage of patients 18 years of age and older who received an elective primary total knee arthroplasty (TKA) and completed a functional status assessment within 90 days prior to the surgery and in the 270-365 days after the surgery</td>
<td>☐</td>
</tr>
<tr>
<td>CMS 56v6 - Functional Status Assessment for Total Hip Replacement</td>
<td>Percentage of patients 18 years of age and older who received an elective primary total hip arthroplasty (THA) and completed a functional status assessment within 90 days prior to the surgery and in the 270-365 days after the surgery</td>
<td>☐</td>
</tr>
<tr>
<td>CMS 90v7 - Functional Status Assessments for Congestive Heart Failure</td>
<td>Percentage of patients 18 years of age and older with congestive heart failure who completed initial and follow-up patient-reported functional status assessments</td>
<td>☐</td>
</tr>
</tbody>
</table>

Select the CQMs that best apply to your scope of practice.

4 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
### Patient Safety

#### Objective

<table>
<thead>
<tr>
<th>Measure</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS 156v6</td>
<td>Percentage of patients 65 years of age and older who were ordered high-risk medications. Two rates are reported.</td>
</tr>
<tr>
<td>CMS 139v6</td>
<td>Percentage of patients 65 years of age and older who were screened for future fall risk during the measurement period.</td>
</tr>
<tr>
<td>CMS 68v7</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 using all immediate resources available on the date of the encounter. This list must include all known prescriptions, over-the-counter, herbals, and vitamin/mineral/dietary (nutritional) supplements AND must contain the medications name, dosage, frequency and route of administration.</td>
</tr>
<tr>
<td>CMS 132v6</td>
<td>Percentage of patients aged 16 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 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>
</tr>
<tr>
<td>CMS 177v6</td>
<td>Percentage of patient visits for those patients aged 6 through 17 years with a diagnosis of major depressive disorder with an assessment for suicide risk.</td>
</tr>
</tbody>
</table>

Select the CQMs that best apply to your scope of practice.

5 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

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**TIP**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Clinical Quality Measures for Communication and Care Coordination

<table>
<thead>
<tr>
<th>Objective</th>
<th>Measure</th>
<th>Selected</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS 50v6 - 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>☐</td>
</tr>
<tr>
<td>CMS 142v6 \ NQF 0089 - 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>☐</td>
</tr>
</tbody>
</table>

Select the CQMs that best apply to your scope of practice.

2 of 55 CQMs is available under this domain.

The Navigation bar at the bottom will monitor your progress.

---

TIP

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
### Clinical Quality Measures for Community / Population Health

<table>
<thead>
<tr>
<th>Objective</th>
<th>Measure</th>
<th>Selected</th>
</tr>
</thead>
</table>
| CMS 15w6 \ NQF 0024 - Weight Assessment and Counseling for Nutrition and Physical Activity for Children and Adolescents | Percentage of patients 3-17 years of age who had an outpatient visit with a Primary Care Physician (PCP) or Obstetrician / Gynecologist (OB/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 |          |
| CMS 138w6 \ NQF 0028 - Preventive Care and Screening: Tobacco Use: Screening and Cessation Intervention | Percentage of patients aged 18 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.  
  Three Rates are Reported:  
  - Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months.  
  - Percentage of patients aged 18 years and older who were screened for tobacco use and identified as a tobacco user who received tobacco cessation intervention.  
  - Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received tobacco cessation intervention if identified as a tobacco user. |          |
| CMS 153w6 \ NQF 0033 - Chlamydia Screening for Women                       | 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 |          |
| CMS 117w6 \ NQF 0038 - Childhood Immunization Status                     | Percentage of children 2 years of age who had four diphtheria, tetanus and acellular pertussis (DTaP); three polo (IPV), one measles, mumps and rubella (MMR); three H influenza type B (HIB); three hepatitis B (Hep B); one chicken pox (VAR); four pneumococcal conjugate (PCV); one hepatitis A (Hep A); two or three rotavirus (RV); and two influenza (flu) vaccines by their second birthday |          |
| CMS 147w7 \ NQF 0041 - Preventive Care and Screening: Influenza Immunization | 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. |          |

Select the CQMs that best apply to your scope of practice.

11 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

---

**TIP**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Clinical Quality Measures for Community / Population Health cont’d.

- **CMS 27v7 \ NQF 0418** - Preventive Care and Screening: Screening for Clinical Depression and Follow-Up Plan
  - Percentage of patients aged 12 years and older screened for clinical depression on the date of the encounter using an age appropriate standardized screening tool AND if positive, a follow up plan is documented on the date of the positive screen.

- **CMS 69v6 \ NQF 0421** - Preventive Care and Screening: Body Mass Index (BMI) Screening and Follow-Up Plan
  - Percentage of patients aged 18 years and older with a BMI documented during the current encounter or during the previous twelve months AND with a BMI outside of normal parameters, a follow-up plan is documented during the encounter or during the previous twelve months of the current encounter. Normal Parameters: Normal Parameters: Age 18 years and older BMI => 18.5 and < 25 kg/m²

- **CMS 82v5 \ NQF 1401** - Maternal depression screening
  - The percentage of children who turned 6 months of age during the measurement year, who had a face-to-face visit between the clinician and the child during child’s first 6 months, and who had a maternal depression screening for the mother at least once between 0 and 6 months of life.

- **CMS 22v6** - Preventive Care and Screening: Screening for High Blood Pressure and Follow-Up Documented
  - Percentage of patients aged 18 years and older seen during the reporting period who were screened for high blood pressure AND a recommended follow-up plan is documented based on the current blood pressure (BP) reading as indicated.

- **CMS 79v6** - Children Who Have Dental Decay or Cavities
  - Percentage of children, ages 0-20 years, who have had tooth decay or cavities during the measurement period.

- **CMS 127v6 \ NQF 0043** - Pneumonia Vaccination Status for Older Adults
  - Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine.

---

**TIP**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.

Select the CQMs that best apply to your scope of practice.

11 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.
Clinical Quality Measures for Efficiency and Cost Reduction

<table>
<thead>
<tr>
<th>Efficiency and Cost Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Objective</td>
</tr>
<tr>
<td>CMS 146v6 \ NQF 0002 -</td>
</tr>
<tr>
<td>Appropriate Testing for</td>
</tr>
<tr>
<td>Children with Pharyngitis</td>
</tr>
<tr>
<td>CMS 160v7 \ NQF 0052 - Use</td>
</tr>
<tr>
<td>of Imaging Studies for Low</td>
</tr>
<tr>
<td>CMS 154v6 \ NQF 0069 -</td>
</tr>
<tr>
<td>Appropriate Treatment for</td>
</tr>
<tr>
<td>Children with Upper Respiratory Infection (URI)</td>
</tr>
<tr>
<td>CMS 129v7 \ NQF 0389 -</td>
</tr>
<tr>
<td>Prostate Cancer: Avoidance of</td>
</tr>
<tr>
<td>Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients</td>
</tr>
</tbody>
</table>

Efficiency and Cost Reduction

Select the CQMs that best apply to your scope of practice.

4 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
### Clinical Quality Measures for Effective Clinical Care

<table>
<thead>
<tr>
<th>Objective</th>
<th>Measure</th>
<th>Selected</th>
</tr>
</thead>
</table>
| CMS 13v6 \ NQF 0004 - Initiation and Engagement of Alcohol and Other Drug Dependence Treatment | Percentage of patients 13 years of age and older with a new episode of alcohol and other drug (AOD) dependence who received the following: Two rates are reported:  
  - Percentage of patients who initiated treatment within 14 days of the diagnosis.  
  - 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. |          |
| CMS 16v6 \ NQF 0018 - Controlling High Blood Pressure                      | Percentage of patients 18-85 years of age who had a diagnosis of hypertension and whose blood pressure was adequately controlled (<140/90mmHg) during the measurement period.                                           |          |
| CMS 125v6 - Breast Cancer Screening                                       | Percentage of women 50-74 years of age who had a mammogram to screen for breast cancer.                                                                                                                |          |
| CMS 124v6 \ NQF 0032 - Cervical Cancer Screening                         | Percentage of women 21-64 years of age who were screened for cervical cancer using either of the following criteria:  
  - Women age 21-64 who had cervical cytology performed every 3 years.  
  - Women age 30-64 who had cervical cytology/human papillomavirus (HPV) co-testing performed every 5 years.                                             |          |
| CMS 130v6 \ NQF 0034 - Colorectal Cancer Screening                      | Percentage of adults 50-75 years of age who had appropriate screening for colorectal cancer.                                                                                                             |          |
| CMS 127v6 \ NQF 0043 - Pneumococcal Vaccination Status for Older Adults | Percentage of patients 65 years of age and older who have ever received a pneumococcal vaccine.                                                                                                         |          |
| CMS 131v6 \ NQF 0055 - Diabetes: Eye Exam                               | Percentage of patients 18-78 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. |          |

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Select the CQMs that best apply to your scope of practice.

29 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.
Clinical Quality Measures for Effective Clinical Care continued

<table>
<thead>
<tr>
<th>Measure Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS 123v6 NQF 0056</td>
<td>Diabetes: Foot Exam</td>
</tr>
<tr>
<td>CMS 122v6 NQF 0059</td>
<td>Diabetes: Hemoglobin A1c (HbA1c) Poor Control (&gt;9%)</td>
</tr>
<tr>
<td>CMS 134v6 NQF 0062</td>
<td>Diabetes: Medical Attention for Nephropathy</td>
</tr>
<tr>
<td>CMS 164v6 NQF 0068</td>
<td>Ischemic Vascular Disease (IVD): Use of Aspirin or Another Antiplatelet</td>
</tr>
<tr>
<td>CMS 145v6 NQF 0070</td>
<td>Coronary Artery Disease (CAD): Beta-Blocker Therapy—Prior Myocardial Infarction (MI) or Left Ventricular Systolic Dysfunction (LVEF &lt;40%)</td>
</tr>
<tr>
<td>CMS 139v6 NQF 0081</td>
<td>Heart Failure (HF): Angiotensin-Converting Enzyme (ACE) Inhibitor or Angiotensin Receptor Blocker (ARB) Therapy for Left Ventricular Systolic Dysfunction (LVSD)</td>
</tr>
</tbody>
</table>

The percentage of patients 18-75 years of age with diabetes (type 1 and type 2) who received a foot exam (visual inspection and sensory exam with mono filament and a pulse exam) during the measurement year.

Percentage of patients 18-75 years of age with diabetes who had hemoglobin A1c > 9.0% during the measurement period.

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.

Percentage of patients 18 years of age and older who were diagnosed with 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 documentation of use of aspirin or another antiplatelet during the measurement period.

Percentage of patients aged 18 years 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 <40% who were prescribed beta-blocker therapy.

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) < 40% who were prescribed ACE inhibitor or ARB therapy either within a 12 month period when seen in the outpatient setting OR at each hospital discharge.

Select the CQMs that best apply to your scope of practice.

29 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

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Click the hyperlink on the ePIP screen to learn more about this requirement.
Clinical Quality Measures for Effective Clinical Care continued

CMS 136v7, NQF 0108 - Follow-Up Care for Children Prescribed ADHD Medication (ADD)
- Percentage of children 6-12 years of age and newly dispensed a medication for attention-deficit/ hyperactivity disorder (ADHD) who had appropriate follow-up care. Two rates are reported:
  - Percentage of children who had one follow-up visit with a practitioner with prescribing authority during the 30-Day Initiation Phase.
  - Percentage of children who remained on ADHD medication for at least 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.

CMS 169v6, NQF 0405 - Bipolar Disorder and Major Depression: Appraisal for alcohol or chemical substance use
- Percentage of patients with depression or bipolar disorder with evidence of an initial assessment that includes an appraisal for alcohol or chemical substance use.

CMS 52v6, NQF 0405 - HIV/AIDS: Pneumocystis Jiroveci Pneumonia (PCP) Prophylaxis
- Percentage of patients aged 6 weeks and older with a diagnosis of HIV/AIDS who were prescribed Pneumocystis jiroveci pneumonia (PCP) prophylaxis.

CMS 133v6, NQF 0565 - Cataracts: 20/40 or Better Visual Acuity within 90 Days Following Cataract Surgery
- 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.

CMS 158v6 - Pregnant women that had HBsAg testing
- This measure identifies pregnant women who had a HBsAg (hepatitis B) test during their pregnancy.

CMS 159v6, NQF 0710 - Depression Remission at Twelve Months
- Adult patients age 18 and older with major depression or dysthymia and an initial PHQ-9 score > 9 who demonstrate remission at twelve months defined as PHQ-9 score less than 5. This measure applies to both patients with newly diagnosed and existing depression whose current PHQ-9 score indicates a need for treatment.

Effective Clinical Care

Select the CQMs that best apply to your scope of practice.

29 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

TIP
Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
Clinical Quality Measures for Effective Clinical Care continued

<table>
<thead>
<tr>
<th>Measure Code</th>
<th>Measure Description</th>
<th>Available</th>
</tr>
</thead>
</table>
| CMS 144v6 \
NQF 0083 | Heart Failure (HF): Beta-Blocker Therapy for Left Ventricular Systolic Dysfunction (LVSD)                                                                                                                                 |           |
| CMS 143v6 \
NQF 0086 | 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) < 40% who were prescribed beta-blocker therapy either within a 12 month period when seen in the outpatient setting OR at each hospital discharge. |           |
| CMS 147v6 \
NQF 0088 | Diabetic Retinopathy: Documentation of Presence or Absence of Macular Edema and Level of Severity of Retinopathy                                                                                                     |           |
| CMS 161v6 \
NQF 0104 | 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. |           |
| CMS 128v6 \
NQF 0105 | Anti-depressant Medication Management                                                                                                                                                                                |           |

**Effective Clinical Care**

Select the CQMs that best apply to your scope of practice.

29 of 55 CQMs are available under this domain.

The Navigation bar at the bottom will monitor your progress.

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Click the hyperlink on the ePIP screen to learn more about this requirement.
### Clinical Quality Measures for Effective Clinical Care continued

| CMS 160v6 | Depression Utilization of the PHQ-9 Tool |
| CMS 74v7 | Primary Care Provider Interventions Offered by Primary Care Providers, including Dentists |
| CMS 149v6 | Dementia: Cognitive Assessment |
| CMS 65v7 | Hypertension: Improvement in Blood Pressure |
| CMS 34v1 | Statin Therapy for the Prevention and Treatment of Cardiovascular Disease |

- **CMS 160v6**: 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.
- **CMS 74v7**: Percentage of children, age 9-20 years, who received a fluoride varnish application during the measurement period.
- **CMS 149v6**: 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.
- **CMS 65v7**: Percentage of patients aged 18-85 years of age with a diagnosis of hypertension whose blood pressure improved during the measurement period.
- **CMS 34v1**: Percentage of the following patients - all considered at high risk of cardiovascular events - who were prescribed or were on statin therapy during the measurement period:
  - Adults aged >= 21 years who were previously diagnosed with or currently have an active diagnosis of clinical atherosclerotic cardiovascular disease (ASCVD), OR
  - Adults aged >= 21 years who have ever had a fasting or direct low-density lipoprotein cholesterol (LDL-C) level >= 190 mg/dL or were previously diagnosed with or currently have an active diagnosis of familial or pure hypercholesterolemia; OR
  - Adults aged 40-75 years with a diagnosis of diabetes with a fasting or direct LDL-C level of 70-189 mg/dL.

- **CMS 645v1**: Bone density evaluation for patients with prostate cancer and receiving androgen deprivation therapy. Patients determined as having prostate cancer who are currently starting or undergoing androgen deprivation therapy (ADT), for an anticipated period of 12 months or greater and who receive an initial bone density evaluation. The bone density evaluation must be prior to the start of ADT or within 3 months of the start of ADT.

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**TIP**

Make sure that you upload all documents that support the above entries in your attestation. You can do so on the Attestation Progress page under Meaningful Use PI Report.

Click the hyperlink on the ePIP screen to learn more about this requirement.
## Appendices

<table>
<thead>
<tr>
<th>Appendix</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Medicaid Patient Volume Report Layout</td>
</tr>
<tr>
<td>B</td>
<td>Medicaid Hospital-Based Report Layout</td>
</tr>
<tr>
<td>C</td>
<td>Needy Patient Volume Report Layout</td>
</tr>
<tr>
<td>D</td>
<td>Needy Practice Predominantly Report Layout</td>
</tr>
<tr>
<td>E</td>
<td>Definitions</td>
</tr>
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<td>F</td>
<td>Frequently Asked Questions</td>
</tr>
<tr>
<td>G</td>
<td>Electronic Funds Transfer – ACH Form Instructions</td>
</tr>
<tr>
<td>H</td>
<td>Electronic Funds Transfer – ACH Form</td>
</tr>
<tr>
<td>I</td>
<td>Contacts</td>
</tr>
</tbody>
</table>
Appendix A – Medicaid Patient Volume Report Layout

Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for each rendering provider.

The Medicaid Patient Volume calculation using **all** places of services is:

- **Numerator:** Medicaid Title XIX Patient Encounters
- **Denominator:** All Patient Encounters [Medicaid + Non-Medicaid]
  
  - Non-Medicaid includes CHIP Title XXI (KidsCare), Medicare, Private Insurance, Self-Pay, Commercial, Sliding Scale, etc.

**Reporting Period** is a continuous 90-day period in the prior calendar year.

<table>
<thead>
<tr>
<th>Description</th>
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</tr>
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<tbody>
<tr>
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<tr>
<td>Patient Name</td>
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</tr>
<tr>
<td>Payer Financial Class</td>
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</tr>
<tr>
<td>Medicaid, CHIP (KidsCare), Medicare, Private Insurance, Self-Pay, Commercial, etc.</td>
<td>Alpha</td>
</tr>
<tr>
<td>Correctional Facilities: Use Medicaid or Non-Medicaid description</td>
<td></td>
</tr>
<tr>
<td>Payer Name (if applicable specify Health Plan Name)</td>
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</tr>
<tr>
<td>Payer Health Plan ID / Site ID (Medicaid or CHIP)</td>
<td>Numeric</td>
</tr>
<tr>
<td>Payer Medicaid/CHIP Coordination of Benefits</td>
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</tr>
<tr>
<td>⚫ For Medicaid Title XIX: Enter Medicaid Primary, Medicaid Secondary, Medicaid Tertiary, etc.</td>
<td>Alpha</td>
</tr>
<tr>
<td>⚫ For CHIP (KidsCare) Title XXI: Enter CHIP Primary, CHIP Secondary, CHIP Tertiary, etc.</td>
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</tbody>
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*Correctional Facility is a practice location for providers rendering care to inmates in a prison, jail, reformatory, work farm, detention center, or any other similar facility maintained by Federal, State or local authorities for the purpose of confinement or rehabilitation of adult or juvenile criminal offenders. **NOTE:** Incarceration & Release Date must be included in your report.
Appendix B – Medicaid Hospital-Based Report Layout

Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for each rendering provider.

The Medicaid Hospital-Based calculation using all Medicaid Title XIX places of service only is:

- **Numerator:** Medicaid Title XIX Hospital-Based Patient Encounters [Place of Service 21 & 23 Only]
- **Denominator:** All Medicaid Title XIX Patient Encounters [All Place of Services]

Reporting Period is a continuous 12-month period in the prior calendar year.

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Appendix C – Needy Patient Volume Report Layout

Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for each rendering provider.

The Needy Patient Volume calculation using all places of services is:

- **Numerator (Needy Patient Encounters):**
  - Needy includes Medicaid Title XIX, CHIP Title XXI (KidsCare) & Patients Paying Below Cost (Sliding Scale)

- **Denominator: All Patient Encounters [Needy + Non-Needy]**
  - Non-Needy includes Medicare, Private Insurance, Self-Pay, Commercial, etc.

Reporting Period is a continuous 90-day period in the prior calendar year.

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Appendix D – Needy Practice Predominantly Report Layout

Patient Encounters are measured by counting unique visits based on date of service per provider per patient. Multiple claims for the same patient on the same day are counted as one visit for each rendering provider.

The Practice Predominantly calculation using all places of services is:
- Numerator: All FQHC/RHC/Tribal Clinic Patient Encounters [Place of Services inside facility only]
- Denominator: All Total Patient Encounters [All Place of Services inside & outside facility]

Reporting Period is a continuous 6-month period in the prior calendar year.

<table>
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## Appendix E – Definitions

### Attestation

The attestation process allows the providers to attest to the Promoting Interoperability Program’s as they demonstrate adoption, implementation, upgrade (AIU), or meaningful use of EHR technology. **AIU attestations are not available after 2016.**

### Promoting Interoperability (PI)

A longitudinal electronic record of patient health information generated by one or more encounters in any care delivery setting. Included in this information are patient demographics, progress notes, problems, medications, vital signs, past medical history, immunizations, laboratory data and radiology reports. The Promoting Interoperability automates and streamlines the clinician’s workflow. The Promoting Interoperability has the ability to generate a complete record of a clinical patient encounter - as well as supporting other care-related activities directly or indirectly via interface - including evidence-based decision support, quality management, and outcomes reporting.

### Eligible Professionals (EP)

Physicians (Doctor of Medicine, Doctor of Osteopathy), Dentists, Nurse Practitioners, Certified Nurse Midwives and Physician Assistants (PA) practicing in a FQHC/RHC/Tribal Clinic led by the PA.

### ePIP

An online application that interfaces with the CMS Registration and Attestation system and the Prepaid Medicaid Management Information System (PMMIS) to allow providers to complete applications for the Medicaid Promoting Interoperability (PI) Program for Arizona.

### Meaningful Use

Use of certified EHR technology (CEHRT) to Improve quality, safety, efficiency, & reduce health disparities; Engage patients & families in their health care; Improve care coordination; Improve population & public health and all the while maintaining privacy and security.

### Meaningful Use Exclusion

A reason or reasons associated with a Meaningful Use objective that can be selected, if applicable, to exempt a provider from having to meet the measure

### Meaningful Use Exemption

Found mainly in the Clinical Quality Measures, this counts the number of members that were seen by a provider during the Meaningful Use Reporting Period, but were not eligible to be included in the measure being reported.

### Meaningful Use Stages

- **Stage 1 Data Capture & Information Sharing:** Requirements focus on electronic data capture and information sharing with the patient or other health care professionals.
- **Stage 2 / Stage 2 Modified Advanced Clinical Processes:** Requirements focus on expanding Stage 1 requirements by emphasizing patient engagement and care coordination. Improvements to ease reporting requirements and align with other quality reporting programs (Stage 2 Modified).
- **Stage 3 Improved Outcome:** Requirements focus on using CEHRT to improve health outcomes.

### Patient Volume Methodology

Method in which an EP reports his/her patient encounters. Individual is the sum of patient encounters for a single EP. Aggregate is the sum of patient encounters for the entire practice (includes all providers).

### Program Year

The calendar year in which a provider is attesting. Providers can participate and receive payment up to a maximum of 6 years.

### Registration

The registration process allows the provider to participate in the Promoting Interoperability Program. Providers must complete a federal and state level registration process. **Only providers transferring from other States are permitted to register to set-up an ePIP account after Program Year 2016.**
<table>
<thead>
<tr>
<th>Q1</th>
<th>Can I switch between Medicare and Medicaid programs?</th>
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<tbody>
<tr>
<td></td>
<td>Providers can switch between the Medicare and Medicaid programs any time before they receive their first incentive payment. Eligible Professionals can switch one time (before 2015) between the Medicare and Medicaid Incentive Programs if they have received one incentive payment.</td>
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</tbody>
</table>

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<th>Can I skip a year after I have started the Promoting Interoperability program?</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Eligible Professionals (EPs) in the Medicaid Promoting Interoperability (PI) program can skip a year without a Medicaid penalty. It is not necessary to notify Medicaid that you are skipping a year. When you return, you continue with the next payment year.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q3</th>
<th>Are physicians who work in hospitals eligible to receive Medicaid Promoting Interoperability (PI) payments?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Physicians who furnish substantially all, defined as 90% or more, of their covered professional services in an inpatient (POS 21) and emergency department (POS 23) of a hospital are not eligible for incentive payments under the Medicare and Medicaid Promoting Interoperability (PI) Programs.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q4</th>
<th>Is my practice eligible to apply &amp; receive payments through the Medicare and Medicaid Promoting Interoperability (PI) Programs?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No, your practice cannot apply for payment. Attestations are submitted by individual Eligible Professionals (EPs) who can voluntarily re-assign payment to their practice.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q5</th>
<th>Will Promoting Interoperability Payments be subject to audit?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Incentive payments made to Eligible Professionals under the Medicaid Promoting Interoperability (PI) Program is subject to audit by the Promoting Interoperability Programs. AHCCCS is responsible for conducting the audit for your attestation. Unless otherwise indicated, you will be contacted by AHCCCS with instructions when you are selected for the State audit. PI audit questions can be directed to the Promoting Interoperability Post Payment Audit Team at: <a href="mailto:EHRPost-PayAudits@azahcccs.gov">EHRPost-PayAudits@azahcccs.gov</a> or 602.417.4440</td>
</tr>
</tbody>
</table>
Appendix F – Frequently Asked Questions regarding Registration

<table>
<thead>
<tr>
<th>Q6</th>
<th>How often do I need to Register?</th>
</tr>
</thead>
</table>
|    | You need to Register **once** in order to participate in the Promoting Interoperability Program. Thereafter, you must keep your registration information updated in each system.  
|    | When updating information in your CMS registration, make sure that you “re-submit” your Registration information and allow 24 – 48 hours to feed to ePIP.  
|    | Each time you attest, it is recommended that you review and update the “Contact Information” in both systems as needed. |

<table>
<thead>
<tr>
<th>Q7</th>
<th>I registered in the CMS Registration &amp; Attestation System but my registration is still showing ‘Send for State Approval’. How can I troubleshoot the problem?</th>
</tr>
</thead>
</table>
|    | After completing the registration in the CMS Registration and Attestation System, allow 24 to 48 hours for your registration information to transfer from that system to Arizona’s Electronic Provider Incentive Payment System (ePIP).  
|    | If your CMS registration status shows ‘**Sent for State Approval**’, please send an inquiry to Medicaid at EHRIncentivePayments@azahcccs.gov for assistance.  
|    | **If your CMS registration status shows ‘Registration Started/Modified/In Progress’, please re-submit your CMS registration.** |

<table>
<thead>
<tr>
<th>Q8</th>
<th>Can providers participating in the Medicare or Medicaid Promoting Interoperability (PI) Programs update their information (for example, if an address was mistakenly entered)? If so, will the State receive an update or full refresh of this information for its Medicaid Promoting Interoperability (PI) Program?</th>
</tr>
</thead>
</table>
|    | Yes, providers who have registered for the Medicare or Medicaid Promoting Interoperability (PI) Programs may correct errors or update information through the registration module on the CMS registration website [https://ehrincentives.cms.gov/hitech/login.action](https://ehrincentives.cms.gov/hitech/login.action)  
|    | The updated registration information will be sent to the State. |

<table>
<thead>
<tr>
<th>Q9</th>
<th>I previously received an Promoting Interoperability payment from another Medicaid State and have since moved to Arizona. Can I continue to participate in the program?</th>
</tr>
</thead>
</table>
|    | Yes, you can continue to participate in the Arizona Medicaid Promoting Interoperability (PI) Program.  
|    | First you must update your changes in the CMS Registration & Attestation System and then register in the State’s Registration & Attestation System to create your ePIP account. |
### Q10
I am ready to start a new attestation but I do not see that option when I log in to ePIP. What are the possible reasons for such?

If a payment decision has not been issued for the prior Program Year in which you attested, you cannot begin a new Program Year attestation.

If your previous attestation was denied or rejected, you may need to have your attestation refreshed.

In any instance if you cannot start a new Program Year, please email the Promoting Interoperability Program team at EHRIncentivePayments@azahcccs.gov.

---

### Q11
How do I know if my Promoting Interoperability (PI) system is certified?

The Medicare and Medicaid Promoting Interoperability (PI) Programs require the use of certified EHR technology, as established by a set of standards and certification criteria.

EHR technology needs to be certified by an ONC-Authorized Testing and Certification Body (ONC-ATCB) in order to qualify for incentive payments. The Certified Health IT Product List (CHPL) is available at [http://www.healthit.hhs.gov/CHPL](http://www.healthit.hhs.gov/CHPL). Providers must maintain the proper certification requirements & submit the required documentation to demonstrate that their EHR technology is properly certified.

---

### Q12
How do we submit documentation to support the attestation?

ePIP is the State’s repository for storing your attestation information. Providers are required to upload their documentation at the time of attestation.

Passwords should follow standard operating procedures to prevent access to your ePIP accounts.

The ePIP website, [https://www.azepip.gov/](https://www.azepip.gov/), has a Hypertext Transfer Protocol Secure (HTTPS) feature which has a built in communications protocol for secure communication over a computer network. Therefore, documents uploaded to ePIP are secure and encrypted.

---

### Q13
How can I change my attestation information after I have attested for the Medicaid Promoting Interoperability (PI) Program?

If you discover that the information you entered during your Medicaid attestation was not complete and accurate for some reason, please email Medicaid at EHRIncentivePayments@azahcccs.gov.
### Appendix F – Frequently Asked Questions regarding Meaningful Use

<table>
<thead>
<tr>
<th>Q14</th>
<th><strong>What is the deadline for Medicaid Eligible Professionals to submit attestations for Program Year 2018?</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Eligible Professionals participate in the Medicaid Promoting Interoperability (PI) Programs on a calendar year basis. Generally, the Medicaid attestation deadline is 90-days following the end of the calendar year. At this time, the deadline for Program Year 2018 has been extended to <strong>August 31, 2019</strong>.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q15</th>
<th><strong>What are the reporting periods for Eligible Professionals participating in the Promoting Interoperability (PI) Program?</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>For Program Year 2018, the reporting periods are as follows:</td>
</tr>
<tr>
<td></td>
<td><strong>Volume (select a period from 2017):</strong></td>
</tr>
<tr>
<td></td>
<td>Patient Volume - a continuous 90-day period in the prior calendar year</td>
</tr>
<tr>
<td></td>
<td>Hospital-Based - a 12-month period in the prior calendar year</td>
</tr>
<tr>
<td></td>
<td>Practice Predominantly - continuous 6-month period in the prior calendar year</td>
</tr>
<tr>
<td></td>
<td><strong>Meaningful Use (select a period from 2018):</strong></td>
</tr>
<tr>
<td></td>
<td>The Promoting Interoperability reporting period for the Meaningful Use Objectives &amp; the Clinical Quality Measures is a continuous 90-day period within the calendar year.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q16</th>
<th><strong>Under the Medicare and Medicaid Promoting Interoperability (PI) Program, who is responsible for demonstrating meaningful use of certified EHR technology, the provider or the vendor?</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>To receive an Promoting Interoperability payment, the Eligible Professional is responsible for demonstrating meaningful use of certified EHR technology under both the Medicare and Medicaid Promoting Interoperability (PI) programs.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q17</th>
<th><strong>Is there a penalty if I start the Promoting Interoperability program and do not attest to Meaningful Use?</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Providers who have a Medicare patient population and have not attested to Meaningful Use will have a reduction in Medicare payments.</td>
</tr>
<tr>
<td></td>
<td>Providers that do not serve Medicare members are not penalized if they do not attest or if they withdraw from the Medicaid Promoting Interoperability (PI) Program after receiving an incentive payment.</td>
</tr>
</tbody>
</table>
### Appendix F – Frequently Asked Questions regarding Payment

<table>
<thead>
<tr>
<th>Q18</th>
<th>I am choosing to reassign my PI payment to my practice. Will I have any financial liability if I do so?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>The State of Arizona issues 1099s to the Payee (recipient) of the Promoting Interoperability funds. If you have reassigned your payment to your practice, you will not personally receive a 1099. For more information on 1099s, visit the AHCCCS website at <a href="https://www.azahcccs.gov/PlansProviders/CurrentProviders/EHR/">https://www.azahcccs.gov/PlansProviders/CurrentProviders/EHR/</a>. Click the Payment drop down and see <strong>IMPORTANT TAX INFORMATION</strong>.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q19</th>
<th>How is the Eligible Professional payment amounts determined?</th>
</tr>
</thead>
</table>
|     | Medicaid EPs can receive a maximum of $63,750 over a six year period.  
**Note:** There are special eligibility & payment options for Pediatricians. |

<table>
<thead>
<tr>
<th>Q20</th>
<th>How often are payments made?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Payments are disbursed once per month via Electronic Funds Transfer.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q21</th>
<th>Are payments from the Medicare and Medicaid Promoting Interoperability (PI) Programs subject to federal income tax?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>We note that nothing in the Act excludes such payments from taxation or as tax-free income. Therefore, it is our belief that incentive payments would be treated like any other income. Providers should consult with a tax advisor or the Internal Revenue Service regarding how to properly report this income on their filings.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q22</th>
<th>Are payments from the Medicare and Medicaid Promoting Interoperability (PI) Programs subject to recoupments?</th>
</tr>
</thead>
</table>
|     | Both Medicare and Medicaid are required to recoup any or all portions of the Promoting Interoperability payment if any of the following conditions are determined:  
- Provider or Payee received an improper payment  
- Provider does not meet the requirements of the program  
- Evidence of fraud and abuse |

<table>
<thead>
<tr>
<th>Q23</th>
<th>How long will it take to receive a payment?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>We must first perform the pre-payment audit. The Promoting Interoperability Team strives to complete within eight (8) weeks of attestation during off peak periods. Delays are experienced when waiting for missing information, resolving issues, during peak periods, training or staffing changes.</td>
</tr>
</tbody>
</table>
# Appendix G – Electronic Funds Transfer ACH Form Instructions

## Provider Information

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provider Name</td>
<td>The legal name of the provider.</td>
<td>Required</td>
</tr>
<tr>
<td>Doing Business As Name (DBA)</td>
<td>The legal name of the provider.</td>
<td>Optional</td>
</tr>
<tr>
<td>Provider Address Street</td>
<td>The number and street name where the provider can be contacted.</td>
<td>Required</td>
</tr>
<tr>
<td>City</td>
<td>City associated with provider address.</td>
<td>Required</td>
</tr>
<tr>
<td>State/Province</td>
<td>State or province associated with provider address.</td>
<td>Required</td>
</tr>
<tr>
<td>Zip Code/Postal Code</td>
<td>Zip Code/Postal Code.</td>
<td>Required</td>
</tr>
</tbody>
</table>

## Provider Identifiers Information

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Federal Tax Identification Number (FTIN) or Employer Identification Number (EIN)</td>
<td>A Federal Tax Identification Number also known as an Employer Identification Number (EIN) used to identify a business entity. Needs 9 digits.</td>
<td>Required</td>
</tr>
<tr>
<td>National Provider Identifier (NPI)</td>
<td>A Health Insurance Portability and Accountability Act (HIPAA) - Required when provider has been enumerated with an NPI.</td>
<td>Optional</td>
</tr>
<tr>
<td>Trading Partner ID</td>
<td>AHCCCS Provider ID, 6 digits - 2 digits.</td>
<td>Required</td>
</tr>
</tbody>
</table>

## Provider Contact Information

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provider Name</td>
<td>Name of a contact in provider office for handling EFT issues.</td>
<td>Required</td>
</tr>
<tr>
<td>Title</td>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td>Tel Number</td>
<td>Number associated with contact person.</td>
<td>Optional</td>
</tr>
<tr>
<td>Tel Number Ext</td>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td>Email Address</td>
<td>An electronic mail address at which ARHCCCS might contact the provider.</td>
<td>Optional</td>
</tr>
<tr>
<td>Fax Number</td>
<td>A number at which the provider can be paged.</td>
<td>Optional</td>
</tr>
</tbody>
</table>

## Provider Agent Information - If Applicable

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provider Agent Name</td>
<td>Name of provider’s authorized agent.</td>
<td>Required</td>
</tr>
<tr>
<td>Agent Address</td>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td>Street</td>
<td>The number and street name where the provider can be contacted.</td>
<td>Required</td>
</tr>
<tr>
<td>City</td>
<td>City associated with provider address.</td>
<td>Required</td>
</tr>
<tr>
<td>State/Province</td>
<td>State or province associated with provider address.</td>
<td>Required</td>
</tr>
<tr>
<td>Zip Code/Postal Code</td>
<td>Zip Code/Postal Code.</td>
<td>Required</td>
</tr>
<tr>
<td>Provider Agent Contact Name</td>
<td>Name of a contact in provider office for handling EFT issues.</td>
<td>Required</td>
</tr>
<tr>
<td>Tel Number</td>
<td>Number associated with contact person.</td>
<td>Optional</td>
</tr>
<tr>
<td>Tel Number Ext</td>
<td></td>
<td>Optional</td>
</tr>
<tr>
<td>Email Address</td>
<td>An electronic mail address at which ARHCCCS might contact the provider.</td>
<td>Optional</td>
</tr>
<tr>
<td>Fax Number</td>
<td>A number at which the provider can be paged.</td>
<td>Optional</td>
</tr>
</tbody>
</table>
# Appendix G – Electronic Funds Transfer ACH Form Instructions (continued)

## FINANCIAL INSTITUTION INFORMATION

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Financial Institution Name / Address</td>
<td>Address associated with financial institution</td>
<td>Required</td>
</tr>
<tr>
<td>State</td>
<td>Street address associated with receiving deposits/financial institution name field</td>
<td>Required</td>
</tr>
<tr>
<td>City</td>
<td>City associated with receiving deposits/financial institution address field</td>
<td>Required</td>
</tr>
<tr>
<td>State/Province Code</td>
<td>State/Province Code associated with the State</td>
<td>Required</td>
</tr>
<tr>
<td>Code</td>
<td>6 to 8 Character Code</td>
<td>Required</td>
</tr>
<tr>
<td>Tax Number</td>
<td>A contact telephone number at the provider's bank</td>
<td>Optional</td>
</tr>
<tr>
<td>Tax Number Ext</td>
<td></td>
<td>Optional</td>
</tr>
</tbody>
</table>

## SECTION 6

### Opening Number

A unique identifier of the financial institution where the provider maintains an account to which payments are to be deposited

### Account Number

The type of account the provider wishes to receive EFT payments, e.g., Checking, Savings

### Account Number Linkage to Provider Identifier

Provider's account number at the financial institution to which EFT payments are to be deposited

### Provider Federal Tax Identification Number (TIN)

Provider preference for grouping (bulking) claim payments – must match preference for UB-0020-835 remittance advice

### National Provider Identifier (NPI)

Optional – required if NPI is not applicable

## SECTION 7

### Reason for Submission

- **New Enrollment**: Required
- **Change Enrollment**: Required
- **Cancel Enrollment**: Required

**Enrollment Submission**

- **Voided Check**: A voided check is attached to provide confirmation of identification/account numbers
- **Bank Letter**: A letter or bank letterhead that normally certifies the account number, routing and account numbers

## AUTHORIZATION

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Required/Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td>Authorized Signature</td>
<td>The signature of an individual authorized by the provider or its agent to initiate, modify or terminate an enrollment</td>
<td>Required</td>
</tr>
<tr>
<td>Joint Name of Authorized Signer</td>
<td>The printed name of the person submitting the form</td>
<td>Required</td>
</tr>
<tr>
<td>Date</td>
<td>The title of person signing the form</td>
<td>Optional</td>
</tr>
<tr>
<td>Submission Date</td>
<td>The date on which the enrollment is submitted - CCYY/MM/DD</td>
<td>Required</td>
</tr>
<tr>
<td>Requested Date</td>
<td>The date on which the requested action is to begin - CCYY/MM/DD</td>
<td>Required</td>
</tr>
</tbody>
</table>

For a full, printable PDF of this document, please click on the following link, [Click Here](https://www.azepip.gov/).
Appendix H – Electronic Funds Transfer ACH Form Sample

<table>
<thead>
<tr>
<th>STATE OF ARIZONA – ARIZONA HEALTH CARE COST CONTAINMENT SYSTEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electronic Funds Transfer (EFT) Authorization Agreement</td>
</tr>
<tr>
<td>Name: AHCCCS Provider (DBA)</td>
</tr>
<tr>
<td>Provider Name:</td>
</tr>
<tr>
<td>Provider Address:</td>
</tr>
<tr>
<td>Provider Federal Tax Identification Number (TIN) or Employee Identification Number (EIN)</td>
</tr>
<tr>
<td>National Provider Identifier (NPI)</td>
</tr>
<tr>
<td>PROVIDER CONTACT INFORMATION</td>
</tr>
<tr>
<td>Provider Contact Name:</td>
</tr>
<tr>
<td>Title:</td>
</tr>
<tr>
<td>Telephone Number &amp; Extension:</td>
</tr>
<tr>
<td>Email Address:</td>
</tr>
<tr>
<td>Fax Number:</td>
</tr>
<tr>
<td>PROVIDER AGENT INFORMATION - APPLICABLE</td>
</tr>
<tr>
<td>Provider Agent Name:</td>
</tr>
<tr>
<td>Provider Agent Address:</td>
</tr>
<tr>
<td>Provider Agent Contact Name:</td>
</tr>
<tr>
<td>Title:</td>
</tr>
<tr>
<td>Telephone Number &amp; Extension:</td>
</tr>
<tr>
<td>Email Address:</td>
</tr>
<tr>
<td>Fax Number:</td>
</tr>
<tr>
<td>FINANCIAL INSTITUTION INFORMATION</td>
</tr>
<tr>
<td>Financial Institution Name:</td>
</tr>
<tr>
<td>Financial Institution Address:</td>
</tr>
<tr>
<td>Financial Institution Telephone Number &amp; Extension:</td>
</tr>
<tr>
<td>Financial Institution Routing Number:</td>
</tr>
<tr>
<td>Type of Account at Financial Institution:</td>
</tr>
<tr>
<td>Provider’s Account Number with Financial Institution:</td>
</tr>
<tr>
<td>Account Number Linkage to Provider Identifier:</td>
</tr>
<tr>
<td>Provider’s Federal Tax Identification Number:</td>
</tr>
<tr>
<td>National Provider Identifier Number:</td>
</tr>
<tr>
<td>SUBMISSION INFORMATION</td>
</tr>
<tr>
<td>Reason for Submission:</td>
</tr>
<tr>
<td>Include with Enrollment Submission:</td>
</tr>
<tr>
<td>OR Bank Letter - A letter on bank letterhead that formally certifies the account owners routing and account numbers</td>
</tr>
<tr>
<td>AUTHORIZATION</td>
</tr>
<tr>
<td>I, the undersigned, authorize the Arizona Department of Administration (ADOA), General Accounting Office (GAO) and the Arizona Health Care Cost Containment System (AHCCCS) to process payment to me via Automated Clearing House (ACH) deposits. The State of Arizona and AHCCCS shall deposit the ACH deposits in the financial institution and account designated above.</td>
</tr>
<tr>
<td>I authorize the State of Arizona to notify me of any changes to my account information.</td>
</tr>
<tr>
<td>By signing this authorization agreement, I authorize the State of Arizona to debit my account for any amount due to the State of Arizona.</td>
</tr>
<tr>
<td>By signing this authorization agreement, I authorize the State of Arizona to credit my account for any amount due to me, including interest.</td>
</tr>
</tbody>
</table>

For a full, printable PDF of this document, please click on the following link, [Click Here](https://www.azepip.gov/)
## Appendix I – Contact Us

<table>
<thead>
<tr>
<th>Need Help with:</th>
<th>Contact Us:</th>
</tr>
</thead>
</table>
| Medicaid Promoting Interoperability (PI) Program | AHCCCS PI Pre-Payment Staff  
602-417-4333  
Email: EHRIncentivePayments@azahcccs.gov  
Website: Arizona Medicaid EHR Incentive Program |
| | AHCCCS Promoting Interoperability Post Payment Staff  
602-417-4440  
Email: EHRPost-PayAudits@azahcccs.gov |

<table>
<thead>
<tr>
<th>Having Trouble with:</th>
<th>Help is Available:</th>
</tr>
</thead>
</table>
| CMS Registration process | CMS Information Center  
888-734-6433  
Website: CMS Medicare and Medicaid EHR Incentive Programs |
| AHCCCS Provider Number, NPI, or TIN | AHCCCS Provider Registration  
602-417-7670 (option 5) Maricopa County  
800-794-6862 Outside Maricopa County  
800-523-0231 Out-of-State  
Website: AHCCCS Provider Registration Unit |
| Electronic Funds Transfer (EFT) | AHCCCS Finance  
602-417-5500  
Website: Automated Clearing House (ACH) Vendor Authorization Form |
| ePIP System | AHCCCS PI Staff  
602-417.4333  
Website: ePIP Systems for Registration & Attestation |
| No-Cost Education & Assistance for HIT / HIE | Arizona Health-e Connection (AzHeC)  
602-688-7200  
Email: ehr@azhec.org |
Thank you for your interest in the Promoting Interoperability Program

Website: Arizona Medicaid EHR Incentive Program

602.417.4333

EHRIncentivePayments@azahcccs.gov