# 2020 OHIC PCMH Quality Measures and Integrated Behavioral Health Reporting

This document is intended only to allow practices to prepare their submission for the PCMH Quality Measures and Integrated Behavioral Health Reporting survey. To complete this survey, please fill out the <u>online survey</u>, or contact Cory King at OHIC (<u>Cory.King@ohic.ri.gov</u>) for an Excel template to use in lieu of the online survey.

#### **General Instructions:**

- Practices should submit data for the October 1, 2019 September 30, 2020 performance period. Practices should <u>only</u> submit data for an alternate performance period if they are unable to pull data for the desired period.
- If practices are submitting data through the online survey, they will be sent a PDF of their submission after completing the survey. Practices will be unable to save partially complete surveys, so practices should gather all of the needed data before entering their survey response.
- Practices should complete the online survey if they are entering data for five practice sites or less. If practices are entering in information for more than five practices, they can either:
  - Fill out the survey as many times as needed in order to enter in information for all practice sites (e.g., a seven-site practice can fill out the survey twice and include data for five sites the first time and two sites the second time), or
  - Contact Cory King for an Excel template to use in lieu of the online survey.
- In 2020, OHIC has make the following changes to this survey: (1) updated measure specifications and dates to align with the national versions of these specifications, (2) temporarily removed the Screening for Clinical Depression and Follow-up Plan measure and added the Lead Screening in Children measure, (3) included questions that ask what practices are doing to capture depression screening information and (4) included telehealth.
  - Practices can apply telehealth encounters for numerator compliance or for identifying patient populations (denominator compliance), even if the specification says otherwise, when reporting data for the PCMH Measure Set for the 10/1/2019 9/30/2020 measurement period. Practices, however, are not required to include telehealth encounters when reporting data. As a reminder, all measures in the PMCH Measure Set are reporting only for the 10/1/2019 9/30/2020 measurement period.
- Please note that for 2020, OHIC has decided to move all measures to "reporting-only" status, as it is anticipated that COVID-19 will have a significant impact on quality measure performance. Therefore, practices only need to report performance on the appropriate measures from the PCMH Measure Set in order to meet this component of the PCMH definition. For this same reason, OHIC will use data from the October 1, 2018 September 30, 2019 performance period to assess improvement for the October 1, 2020 September 30, 2021 performance period.

#### **Practice Site Information:**

- Please fill out your contact information.
  - o First Name
  - Last Name
  - o Title
  - Email Address
  - o Phone Number
- Please fill out the contact information for the practice site. If your practice site is a CTC-RI participant, please use the same site name as employed in your CTC-RI data submissions.
  - Name of Practice Site
  - Street Address
  - City
  - State
  - o Zip
  - Site Contact Person's Name
  - Site Contact Person's Email Address
  - o Phone Number
  - Fax Number
- Did this practice site respond to the OHIC 2019 PCMH Measures Survey? You will be prompted to enter the practice site's OHIC PCMH ID Number if the practice site did respond to the OHIC 2019 PCMH Measures Survey. Otherwise, OHIC will assign this practice site with an OHIC PCMH ID Number after reviewing the results of the 2020 PCMH Measures Survey.
  - Yes
  - o No
  - o [If yes] What is the OHIC PCMH ID Number for this practice site? A practice site has an OHIC PCMH ID Number if it has previously responded to the OHIC 2019 PCMH Measures Survey. See the bottom of this web page for more information: http://www.ohic.ri.gov/ohic-reformandpolicy-pcmhinfo.php.
- For what health plans is your practice site a contracted provider (*check all that are applicable*)?
  - Blue Cross Blue Shield of Rhode Island
  - o Neighborhood Health Plan of Rhode Island
  - o Tufts Health Plan
  - UnitedHealthcare
- What is the Tax Identification Number (TIN) for this practice site?
- What are the NPI numbers for all clinicians at this site managing a patient panel (*list each MD's*, *NP's*, *PA's*)?
- Which of the below specialties best indicates the primary care specialty(ies) of this practice site?
  - o Internal Medicine, Family Practice, or General Practice
  - o Pediatric Practice
  - o Both

- Are more than 50% of your practice site's patients covered by Medicaid or uninsured?
  - o Yes
  - o No
- In what transformation year is your practice?
  - Year 1: Practice joined CTC (including PCMH Kids) on or after January 1, 2020 or practice is not participating in any formal transformation initiative
  - Year 2: Practice joined CTC (including PCMH Kids) during 2018 or 2019, or independently achieved NCQA PCMH Level 3 recognition during 2018 or 2019
  - Year 3: Practice joined CTC prior to January 1, 2018 or independently achieved NCQA PCMH Level 3 recognition prior to January 1, 2018
- Date Survey Completed

#### Behavioral Health Integration and Depression Screening

Please answer the following questions about what your practice site is doing to integrate behavioral health and to capture depression screening and follow-up electronically.

- Has the practice received the NCQA Behavioral Health Distinction, or is the practice receiving facilitated assistance from a formal program designed to assist primary care practices in achieving the NCQA Behavioral Health Distinction?<sup>1</sup>
  - o Yes
  - o No
- Does the practice currently, or did the practice participate in and successfully complete the CTC Integration Behavioral Health Program?
  - Yes
  - o No
- If you answered "no" to the previous two questions, has the practice completed a
  behavioral health integration self-assessment tool AND developed an action plan for
  improving its level of integration? Self-assessment tools include, but are not limited to:
   Organizational Assessment Toolkit for Primary and Behavioral Health Care Integration,
  the PCBH Implementation Kit, and the Maine Health Access Foundation Site Self-Assessment.
  - o Yes
  - o No
- In what format is this practice site capturing the results of a depression screen electronically (*check all that are applicable*)?
  - o A PHQ-9 total score
  - o A PHQ-9 score that includes a numeric value for every answer
  - o Positive/negative check box or similar yes/no method
  - o A score using another assessment tool
    - If yes, what is the assessment tool and for what population(s) is it employed?
  - Free text
  - Scanned copy of completed survey
  - Not applicable
  - Other (please specify)
- In what format is this practice site capturing follow-up activity performed by someone in your practice or ACO following a positive screen (*check all that are applicable*)?
  - o A drop-down menu or check boxes with specified follow-up actions
  - o A check-off box indicating follow-up was performed (without detail)
  - o Order or referral stored electronically in the EHR

<sup>&</sup>lt;sup>1</sup> Definition of "Formal program." A formal program consists of a structured training or support program for primary care providers and/or behavioral health providers with a pre-defined curriculum and technical assistance and designed to systematically build the skills within the practice with a goal of pursuing and attaining NCQA Behavioral Health Distinction.

- o Free text
- o Not applicable
- o Other (please specify)
- In what format is this practice site capturing follow-up activity performed **outside your group** (typically by a behavioral health provider) [*check all that are applicable*]?
  - o Scanned consult note into media
  - o Closed order or referral
  - o Free text
  - o Not applicable
  - o Other (please specify)
- Please provide any additional information about what this practice site is doing to capture depression screening and follow-up electronically.

## Adult Measures

## **Colorectal Cancer Screening**

Description:	The percentage of active patients 50 to 75 years of age who had an
	appropriate screening for colorectal cancer.
Age Criteria:	Eligible population is determined as patients 51 to 75 years of age at the end of the measurement period. (Description states 50 since someone could be 50 throughout the measurement year and not turn 51 until the last day of the measurement period).
Numerator	Active patients 51 to 75 at the end of the measurement period who
Statement:	received an acceptable colorectal screening during the identified lookback period (See below).
Denominator	Active patients 51-75 at the end of the measurement period.
Statement:	
Acceptable	Either of the following at any time in the member's history
Exclusions:	through the end of the measurement period:
LACIUSIOIIS.	
	Colorectal cancer
	Total colectomy
	Patients 66 to 75 as of December 31st of the measurement year
	with frailty (Frailty Value Set) AND advanced illness during the
	measurement year. To identify members with advanced illness,
	any of the following during the measurement year or the year
	prior, meet the criteria:
	At least two outpatient visits (Outpatient Value Set),
	observation visits (Observation Value Set), ED visits (ED
	Value Set) or nonacute inpatient encounters (Nonacute
	, , , , , , , , , , , , , , , , , , , ,
	Inpatient Value Set) on different dates of service with an
	advanced illness diagnosis (Advanced Illness Value Set).
	Visit type need not be the same for the two visits.
	<ul> <li>At least one acute inpatient encounter (Acute Inpatient</li> </ul>
	Value Set) with an advanced illness diagnosis
	(Advanced Illness Value Set)
	o À dispensed dementia medication (Dementia
	Medications List)
Look Back Period:	Varies based on test performed:
20011 20011 1 0110 011	Fecal occult blood test during the measurement year (FOBT)
	Value Set)
	,
	Flexible sigmoidoscopy during the measurement year or the  four years prior to the measurement year (Flexible)
	four years prior to the measurement year (Flexible
	Sigmoidoscopy Value Set)
	Colonoscopy during the measurement year or the nine years
	prior to the measurement year (Colonoscopy Value Set)
	CT colonography during the measurement year or the four
	years prior to the measurement year (CT Colonography Value
	Set)
	,

#### FIT-DNA (Cologuard) test during measurement year or the two years prior to the measurement year (FIT-DNA Value Set)

## Medical Record Documentation:

If a copy of the actual procedure/test/lab result is not present, documentation in the medical record must include a note indicating the date when the colorectal screening was performed. A result is not required if the documentation is clearly part of the "medical history" section of the record. If that is not clear, the result finding must be present (this ensures the screening was performed and not merely ordered).

A pathology report that indicates the type of screening meets the criteria.

For pathology reports that do not indicate the type of screening and for incomplete procedures:

- Evidence that the scope advanced beyond the splenic flexure meets criteria for a completed colonoscopy.
- Evidence that the scope advanced into the sigmoid colon meets criteria for a completed flexible sigmoidoscopy.

There are two types of FOBT tests: guaiac (gFOBT) and immunochemical (FIT). Depending on the type of FOBT test, a certain number of samples are required for numerator compliance. Follow the instructions below to determine member compliance.

- If the medical record does not indicate the type of test and there is no indication of how many samples were returned, assume the required number was returned. The member meets the screening criteria for inclusion in the numerator.
- If the medical record does not indicate the type of test and the number of returned samples is specified, the member meets the screening criteria only if the number of samples specified is greater than or equal to three samples. If there are fewer than three samples, the member does not meet the screening criteria for inclusion.
- FIT tests may require fewer than three samples. If the medical record indicates that an FIT was done, the member meets the screening criteria, regardless of how many samples were returned.
- If the medical record indicates that a gFOBT was done, follow the scenarios below.
  - If the medical record does not indicate the number of returned samples, assume the required number was returned. The member meets the screening criteria for inclusion in the numerator.

	<ul> <li>If the medical record indicates that three or more samples were returned, the member meets the screening criteria for inclusion in the numerator.</li> <li>If the medical record indicates that fewer than three samples were returned, the member does not meet the screening criteria.</li> </ul>
	Do not count digital rectal exams (DRE), FOBT tests performed in an office setting or performed on a sample collected via DRE.
Source:	HEDIS®

- Please indicate for what performance period you are reporting data for "Colorectal Cancer Screening."
  - o October 1, 2019 September 30, 2020
  - o Other (Please Specify)
- Enter the numerator for "Colorectal Cancer Screening."
- Enter the denominator for "Colorectal Cancer Screening."

## Comprehensive Diabetes Care - Eye Exam

Description:	The percentage of active diabetic patients between 18 and 75 years of age with up to date screening or monitoring for diabetic retinal disease
Age Criteria:	Eligible population is determined as 18 or 75 at the end of the measurement period.  Example:  Measurement period end date 12/31/2019  Patient age between 18 as of 12/31/2019 to 75 as of 12/31/2019
Numerator	Active patients with diabetes between 18 and 75 years of age at the end
Statement:	<ul> <li>of the measurement period who had any of the following:</li> <li>A retinal or dilated eye exam by an eye care professional (optometrist or ophthalmologist) in the measurement year</li> <li>A negative retinal or dilated eye exam (negative for retinopathy) by an eye care professional in the year prior to the measurement year.</li> <li>Bilateral eye enucleation anytime during the member's history through the end of the measurement year</li> <li>Please note, documentation in the chart must include one of the following:</li> </ul>
	<ul> <li>A note or letter prepared by an ophthalmologist, optometrist, PCP or other health care professional indicating that an ophthalmoscopic exam was completed by an eye care professional, the date when the procedure was performed and the results.</li> <li>A chart or photograph indicating the date when the fundus photography was performed and evidence that an eye care professional reviewed the results. Alternatively, results may be read by a qualified reading center that operates under the direction of a medical director who is a retinal specialist.</li> <li>Evidence that the member had bilateral eye enucleation or acquired absence of both eyes. Look as far back as possible in the patient's history through end of the measurement year.</li> <li>Documentation of a negative retinal or dilated exam by an eye care professional in the year prior to the measurement year, where results indicate retinopathy was not present (e.g. documentation of normal findings).</li> </ul>
Denominator Statement:	Active patients with diabetes between 18 and 75 years of age at the end of the measurement period with documentation of diabetes during the measurement year or the year prior. Patients with diabetes are identified in the following ways:

1. Encounter-based - Members who met any of the following criteria during the measurement year or the year prior to the measurement year: a. At least two outpatient visits (Outpatient Value Set, Telephone Visits Value Set, Online Assessments Value Set), on different dates of service, with a diagnosis of diabetes (Diabetes Value Set). Only one of the two visits may be a telehealth visit, a telephone visit or an online assessment. Identify telehealth visits by the presence of a telehealth modifier (Telehealth Modifier Value Set) or the presence of a telehealth POS code (Telehealth POS Value Set) associated with the outpatient visit. Use the code combinations below to identify telephone visits and online assessments: o A telephone visit (Telephone Visits Value Set) with a diagnosis of diabetes (Diabetes Value Set) o An online assessment (Online Assessments Value Set) with any diagnosis of diabetes (Diabetes Value Set) 2. Pharmacy Data: Patients who were dispensed insulin or hypoglycemics/antihyperglycemics on an ambulatory basis during the measurement year or year prior (Diabetes Medications List). Note that Glucophage/metformin as a solo agent is NOT included because it is used to treat conditions other than diabetes; patients with diabetes on these medications are identified through diagnosis codes only. Acceptable Patients who do not have a diagnosis of diabetes (Diabetes Value **Exclusions:** Set) in any setting during the measurement year or year prior AND who had a diagnosis included in the Diabetes Exclusions Value Set during the measurement year or year prior. (Historically, these exclusions were limited to gestational and steroid induced diabetes, but the exclusion value set includes additional conditions focused heavily on diabetes caused by an underlying condition). 2. Patients 66 to 75 as of December 31st of the measurement year with frailty (Frailty Value Set) AND advanced illness during the measurement year. To identify members with advanced illness, any of the following during the measurement year or the year prior, meet the criteria: a. At least two outpatient visits (Outpatient Value Set), observation visits (Observation Value Set), ED visits (ED Value Set) or nonacute inpatient encounters (Nonacute Inpatient Value Set) on different dates of service with an advanced illness diagnosis (Advanced Illness Value Set).

Visit type need not be the same for the two visits.

	<ul> <li>b. At least one acute inpatient encounter (Acute Inpatient Value Set) with an advanced illness diagnosis (Advanced Illness Value Set)</li> <li>c. A dispensed dementia medication (Dementia Medications List)</li> <li>3. Patients in hospice</li> </ul>
Look Back	24 months, if negative retinopathy, 12 if positive or unknown
Period:	
Source:	HEDIS®

- Please indicate for what performance period you are reporting data for "Comprehensive Diabetes Care Eye Exam."
  - o October 1, 2019 September 30, 2020
  - o Other (Please Specify)
- Enter the numerator for "Comprehensive Diabetes Care Eye Exam."
- Enter the denominator for "Comprehensive Diabetes Care Eye Exam."

## Comprehensive Diabetes Care: HbA1c Control (<8)

Description:	The percentage of active diabetic patients between 18 and 75 years of age whose most recent HbA1C value was less than 8
Age Criteria:	Eligible population is determined as 18 or 75 at the end of the measurement period.  Example: Measurement period end date 12/31/2019 Patient age between 18 as of 12/31/2019 to 75 as of 12/31/2019
Numerator	Active diabetic patients between 18 and 75 years of age at the end of the
Statement:	measurement period whose most recent HbA1C value in the measurement year was less than 8
Denominator	Active patients with diabetes between 18 and 75 years of age at the end
Statement:	of the measurement period with documentation of diabetes during the measurement year or the year prior. Patients with diabetes are identified in the following ways:
	<ul> <li>2. Encounter-based - Members who met any of the following criteria during the measurement year or the year prior to the measurement year: <ul> <li>a. At least two outpatient visits (Outpatient Value Set, Telephone Visits Value Set, Online Assessments Value Set), on different dates of service, with a diagnosis of diabetes (Diabetes Value Set). Only one of the two visits may be a telehealth visit, a telephone visit or an online assessment. Identify telehealth visits by the presence of a telehealth modifier (Telehealth Modifier Value Set) or the presence of a telehealth POS code (Telehealth POS Value Set) associated with the outpatient visit. Use the code combinations below to identify telephone visits and online assessments:</li> <li>A telephone visit (Telephone Visits Value Set) with a diagnosis of diabetes (Diabetes Value Set) with any diagnosis of diabetes (Diabetes Value Set) with any diagnosis of diabetes (Diabetes Value Set)</li> </ul> </li> </ul>
	3. Pharmacy Data: Patients who were dispensed insulin or hypoglycemics/antihyperglycemics on an ambulatory basis during the measurement year or year prior (Diabetes Medications List). Note that Glucophage/metformin as a solo agent is NOT included because it is used to treat conditions other than diabetes; patients with diabetes on these medications are identified through diagnosis codes only.
Acceptable Exclusions:	Patients who do not have a diagnosis of diabetes (Diabetes Value Set) in any setting during the measurement year or year prior AND who had a diagnosis included in the Diabetes Exclusions

	77 1 0 1 1 1 1
	Value Set during the measurement year or year prior.
	(Historically, these exclusions were limited to gestational and
	steroid induced diabetes, but the exclusion value set includes
	additional conditions focused heavily on diabetes caused by an
	underlying condition).
	2. Patients 66 and older as of December 31st of the measurement
	year with frailty (Frailty Value Set) AND advanced illness during
	the measurement year. To identify members with advanced
	illness, any of the following during the measurement year or the
	year prior, meet the criteria:
	a. At least two outpatient visits (Outpatient Value Set),
	observation visits (Observation Value Set), ED visits (ED
	Value Set) or nonacute inpatient encounters (Nonacute
	Inpatient Value Set) on different dates of service with an
	advanced illness diagnosis (Advanced Illness Value Set).
	Visit type need not be the same for the two visits.
	b. At least one acute inpatient encounter (Acute Inpatient
	Value Set) with an advanced illness diagnosis (Advanced
	Illness Value Set)
	c. A dispensed dementia medication (Dementia Medications
	List)
	3. Patients in hospice
Diabetics	If no A1C reading was rendered during the measurement year, patient
	counts as non-adherent.
Documented:	
Look Back	12 months
Period:	
Source:	HEDIS®

- Please indicate for what performance period you are reporting data for "Comprehensive Diabetes Care HbA1c Control (<8)."
  - o October 1, 2019 September 30, 2020
  - Other (Please Specify)
- Enter the numerator for "Comprehensive Diabetes Care HbA1c Control (<8)."
- Enter the denominator for "Comprehensive Diabetes Care HbA1c Control (<8)."

## **Controlling High Blood Pressure**

Description:	The percentage of active patients between 18 and 85 years who had a diagnosis of hypertension and whose BP was adequately controlled during the measurement year based on the following criteria:
	<ul> <li>Patients 18-85 years of age whose BP was &lt;140/90 mm Hg</li> </ul>
Age Criteria:	Eligible population is determined as 18 or 85 at the end of the measurement period.
	Example:
	Measurement period end date 12/31/2019
	Patient age between 18 as of 12/31/2019 to 85 as of 12/31/2019
Numerator Statement:	Active hypertensive patients between 18 and 85 years of age at the end of the measurement period whose BP was adequately controlled during the measurement year based on the following criteria:
	Patients 18-85 years of age whose most recent BP reading during the measurement year, on or after the second diagnosis of hypertension, (hypertension diagnosis may be established prior to the measurement year if patient has already had two dates of service with a hypertension diagnosis) was <140/90 mm Hg
Denominator	Active hypertensive patients between 18 and 85 years of age at the end of
Statement:	the measurement period. Active hypertension patients are identified as patients who had at least two visits on different dates of service with a diagnosis of hypertension during the measurement year or year prior to the measurement year (count services that occur over both years). Only one of the two visits may be a telephone visit, an online assessment or telehealth visit. Any of the following code combinations meet criteria:  • Outpatient visit (Outpatient Without UBREV Value Set) with or without a telehealth modifier with any diagnosis of hypertension (Essential Hypertension Value Set)  • A telephone visit (Telephone Visits Value Set) with any diagnosis of hypertension (Essential Hypertension Value Set)  • An online assessment (Online Assessment Value Set) with any diagnosis of hypertension (Essential Hypertension Value Set)
Acceptable Exclusions:	<ol> <li>Patients 81 years of age and older as of December 31st of the measurement year with frailty (Frailty Value Set) during the measurement year.</li> <li>Patients 66-85 as of December 31st of the measurement year with frailty (Frailty Value Set) AND advanced illness during the measurement year. To identify members with advanced illness, any of the following during the measurement year or the year prior, meet the criteria:         <ul> <li>At least two outpatient visits (Outpatient Value Set),</li> </ul> </li> </ol>
	observation visits (Observation Value Set), ED visits (ED Value Set) or nonacute inpatient encounters (Nonacute

	Inpatient Value Set) on different dates of service with an
	advanced illness diagnosis (Advanced Illness Value Set).
	Visit type need not be the same for the two visits.
	b. At least one acute inpatient encounter (Acute Inpatient
	Value Set) with an advanced illness diagnosis (Advanced
	Illness Value Set)
	c. A dispensed dementia medication (Dementia Medications List)
	3. Patients with ESRD (ESRD Value Set: ESRD Obsolete Value Set)
	or kidney transplant (Kidney Transplant Value Set) on or prior to
	December 31 of the measurement year. Documentation in the
	medical record must include a dated note indicating evidence of
	ESRD, kidney transplant, or dialysis. (optional)
	4. Patients with a diagnosis of pregnancy (Pregnancy Value Set) during the measurement year (optional)
	5. Patients who had a non-acute inpatient admission during the
	measurement year. (This exclusion is much more feasible for a
	health plan to apply than a practice). To identify non-acute
	inpatient admissions:
	a. Identify all acute and non-acute inpatient stays (Inpatient
	Stay Value Set).  b. Confirm the stay was for non-acute care based on the
	presence of a non-acute code (Non-acute Inpatient Stay
	Value Set) on the claim.
	c. Identify the discharge date for the stay.
	6. Patients in hospice
BP Documentations	The most recent BP reading during the measurement year on or after the
Documentation:	second diagnosis of hypertension (hypertension diagnosis may be established prior to the measurement year if patient has already had two
	dates of service with a hypertension diagnosis). If multiple BP
	measurements occur on the same date, or are noted in the chart on the
	same date, use the lowest systolic and lowest diastolic BP reading. If no
	BP reading is recorded during the measurement year, assume that the
	patient is "not controlled." BP readings from remote monitoring devices that are digitally stored and transmitted to the provider may be included.
	There must be documentation in the medical record that clearly states the
	reading was taken by an electronic device, and results were digitally
	stored and transmitted to the provider, and interpreted by the provider.
	<b>Note:</b> Member-reported results to the provider from a remote monitoring device
	are not acceptable.
	Coding tip: HEDIS does allow ambulatory BP monitoring (CPT
	93784,93788 and 93790) and analysis of electronic data (99091) to qualify as measurement methodologies.
Look Back	12 months
Period:	
Source:	HEDIS®

- Please indicate for what performance period you are reporting data for "Controlling High Blood Pressure."
  - o October 1, 2019 September 30, 2020
  - o Other (Please Specify)
- Enter the numerator for "Controlling High Blood Pressure."
- Enter the denominator for "Controlling High Blood Pressure."

#### Pediatric Measures

#### **Adolescent Well Care Visits**

Description:	The percentage of active patients 12-21 years of age with a documented
	well child encounter during the measurement year
Age Criteria:	Active patients 12-21 years of age at the end of the measurement year.
Numerator	Active patients 12-21 years of age at the end of the measurement year
Statement:	with a note indicating a visit to a PCP or OBGYN, the date of well visit,
	and evidence of all of the following:
	A health and developmental history (physical and mental)
	A physical exam
	Health education/anticipatory guidance
	, 1 3 0
	If standard preventive visit templates consistently incorporate the above
	information, practices may simply use encounter information to verify
	compliance.
Denominator	Active patients 12-21 years of age at the end of the measurement year
Statement:	
Acceptable	None
<b>Exclusions:</b>	
Codes to Identify	CPT: 99383-99385; 99393-99395
Adolescent Well-	ICD-10: Z00.00, Z00.01, Z00.121, Z00.129, Z00.5, Z00.8, Z02.0, Z02.1,
Care Visits	Z02.2, Z02.3, Z02.4, Z02.5, Z02.6, Z02.71, Z02.79, Z02.81, Z02.82, Z02.83,
	Z02.89, Z02.9
Look Back	12 months
Period:	
Source:	HEDIS®

- Please indicate for what performance period you are reporting data for "Adolescent Well Care Visits."
  - o October 1, 2019 September 30, 2020
  - o Other (Please Specify)
- Enter the numerator for "Adolescent Well Care Visits."
- Enter the denominator for "Adolescent Well Care Visits."

## **Developmental Screening in the First Three Years of Life**

Description:  Age Criteria:	The percentage of active patients screened for risk of developmental, behavioral and social delays using a standardized screening tool in the first three years of life. This is a measure of screening in the first three years of life that includes three, age-specific indicators assessing whether children are screened by 12 months of age, by 24 months of age and by 36 months of age.  Children who turn 1, 2, or 3 years of age during the measurement year.
Numerator	The numerator identifies children who were screened for risk of
Statement:	developmental, behavioral and social delays using a standardized tool. National recommendations call for children to be screened at the 9, 18, and 24- OR 30-month well visits to ensure periodic screening in the first, second, and third years of life. The measure is based on three, agespecific indicators.
	Numerators 1-3 are for your understanding of the measures. Only
	Numerator 4 is required to report to PCMH-Kids.
	Numerator 1: Children in Denominator 1 who had screening for risk of developmental, behavioral and social delays using a standardized screening tool that was documented by their first birthday
	<ul> <li>Numerator 2: Children in Denominator 2 who had screening for risk of developmental, behavioral and social delays using a standardized screening tool that was documented after their first and before or on their second birthday</li> </ul>
	Numerator 3: Children in Denominator 3 who had screening for risk of developmental, behavioral and social delays using a standardized screening tool that was documented after their second and before or on their third birthday
	• Numerator 4: Children in Denominator 4 who had screening for risk of developmental, behavioral and social delays using a standardized screening tool that was documented by their first, second or third birthday, i.e., the sum of numerators 1, 2, and 3.
	<ul> <li>Documentation in the medical record must include all of the following:</li> <li>A note indicating the date on which the test was performed, and</li> <li>The standardized tool used (see below), and</li> <li>Evidence of a screening result or screening score</li> </ul>
	<ol> <li>Tools must meet the following criteria:</li> <li>Developmental domains: The following domains must be included in the standardized developmental screening tool: motor, language, cognitive, and social-emotional.</li> <li>Established Reliability: Reliability scores of approximately 0.70 or above</li> </ol>

3. Established Findings Regarding the Validity: Validity scores for the tool must be approximately 0.70 or above. Measures of validity must be conducted on a significant number of children and using an appropriate standardized developmental or socialemotional assessment instrument(s). 4. Established Sensitivity/Specificity: Sensitivity and specificity scores of approximately 0.70 or above Current recommended tools that meet these criteria: 1. Ages and Stages Questionnaire (ASQ) - 2 months - 5 years 2. Ages and Stages Questionnaire - 3rd Edition (ASQ-3) 3. Battelle Developmental Inventory Screening Tool (BDI-ST) – Birth - 95 months 4. Bayley Infant Neuro-developmental Screen (BINS) - 3 months - 2 vears 5. Brigance Screens-II - Birth - 90 months 6. Child Development Inventory (CDI) - 18 months-6 years 7. Infant Development Inventory - Birth - 18 months 8. Parents' Evaluation of Developmental Status (PEDS) - Birth - 8 vears 9. Parent's Evaluation of Developmental Status - Developmental Milestones (PEDS-DM) 10. Survey of Wellbeing of Young Children (SWYC) Tools NOT included in this measure: It is important to note that standardized tools specifically focused on one domain of development [e.g. child's socio-emotional development (ASQ-SE) or autism (M-CHAT)] are not included in the list above as this measure is anchored to recommendations focused on global developmental screening using tools that focus on identifying risk for developmental, behavioral and social delays. Denominator Active patients who have been seen by the primary care clinician at the **Statement:** PCMH in the previous 12 months who meet the following eligibility requirement based on child's age at end of measurement year **Denominator 1**: Active Patients who turn 1 during measurement **Denominator 2**: Active Patients who turn 2 during measurement **Denominator 3**: Active Patients who turn 3 during measurement **Denominator 4**: All Active Patients who turn 1, 2, or 3 during the measurement year, i.e., the sum of denominators 1, 2, and 3 Acceptable None **Exclusions:** 

Look Back	Screenings must be completed prior to the patient's birthdate. In order to
Period:	account for patients with birthdates at the beginning of the measurement
	year, reports should account for these encounters accordingly and place a
	lookback period on the patient's DOB rather than the measurement
	period. In order to account for age appropriate screenings, this look back
	should not exceed a 6 month lookback from the DOB in order to avoid
	erroneously counting developmental screenings used for prior years of
	age.
	Example:
	Patient 1 DOB: 1/15/2019
	Patient 2 DOB: 5/31/2019
	Measurement period for both Patient 1 and 2: 1/1/2019 – 12/31/2019
	Lookback period for Patient 1: 7/15/2018 -1/14/2019
	Lookback period for Patient 2: 11/15/2018 - 5/30/2019
Source:	Oregon Pediatric Improvement Partnership at Oregon Health and
	Science University (OHSU)

- Please indicate for what performance period you are reporting data for "Developmental Screening in the First Three Years of Life."
  - o October 1, 2019 September 30, 2020
  - Other (Please Specify)
- Enter the numerator for "Developmental Screening in the First Three Years of Life."
- Enter the denominator for "Developmental Screening in the First Three Years of Life."

#### **Lead Screening in Children**

Description:	The percentage of active patients two years of age who had one or more capillary or venous lead blood test for lead poisoning by their second birthday.
Age Criteria:	Active patients who turn two years of age during the measurement year.
Numerator	Active patients who turn two years of age during the measurement year
<b>Statement:</b>	with at least one lead capillary or venous blood test (Lead Tests Value
	Set) on or before the child's second birthday. Documentation must
	include both of the following:
	A note indicating the date the test was performed.
	The result or finding.
Denominator	Active patients who turn two years of age during the measurement year
Statement:	
Acceptable	None
<b>Exclusions:</b>	
Look Back	12 months
Period:	
Source:	HEDIS® (modified by OHIC to obtain data through KIDSNET. OHIC is
	working with RIDOH and KIDSNET to obtain data for this measure on
	behalf of practices. HEDIS limits the population to Medicaid-only.
	KIDSNET has adapted the measure specification to apply to all children,
	regardless of insurance type, and defines the active patient using its
	attribution methodology rather than that of the practice site. KIDSNET
	attributes patients based on the provider who delivered the last
	vaccination.)

- Does this practice site give RIDOH permission to release practice-level lead screening data to OHIC for the purposes of assessing whether the practice met the performance improvement requirement of OHIC's PCMH definition?
  - o Yes
  - o No
  - o [If yes] What is your Rhode Island KIDSNET Practice ID?
  - o [*If no*] Please indicate for what performance period you are reporting data for "Lead Screening in Children."
    - October 1, 2019 September 30, 2020
    - Other (Please Specify)
  - o [If no] Enter the numerator for "Lead Screening in Children."
  - o [If no] Enter the denominator for "Lead Screening in Children."

## Weight Assessment and Counseling for Nutrition and Physical Activity

Description:	Percentage of active patients 3-17 years of age who had an outpatient visit in the last twelve months with a primary care clinician of the PCMH who had evidence of the following during the measurement year:  • Body mass index (BMI) percentile documentation,  • Counseling for nutrition, AND  • Counseling for physical activity
Age Criteria:	Eligible population is determined as 3-17 at the end of the measurement year.  Example:  Measurement period end date 12/31/2019  Patient age between 3 as of 12/31/2019 to 17 as of 12/31/2019
Numerator	Patients in the denominator who had evidence of a body mass index
Statement:	(BMI) percentile documentation, counseling for nutrition, AND
	counseling for physical activity during the measurement year
	BMI percentile: documentation must include height, weight, and BMI percentile during the measurement year. The height, weight, and BMI must be from the same data source.  Either of the following meets criteria for BMI percentile:  BMI percentile, or  Ranges and thresholds do not meet criteria for this indicator. A distinct BMI percentile is required for numerator compliance. Documentation of >99% or <1% meet criteria because a distinct BMI percentile is evident (i.e., 100% or 0%).  Practices may utilize the BMI Percentile Value Set as a mechanism to achieve this component of the measure.  Counseling for nutrition: documentation of counseling for nutrition or referral for nutrition education during the measurement year. Documentation must include a note indicating the date and at least one of the following:  Discussion of current nutrition behaviors (e.g. eating habits, dieting behaviors)  Checklist indicating nutrition was addressed  Counseling or referral for nutrition education  Patient received educational materials on nutrition during a face to face visit  Anticipatory guidance for nutrition  Weight or obesity counseling  Practices may utilize the Nutrition Counseling Value Set as a mechanism to achieve this component of the measure, but still must meet the above documentation requirements.

	Counseling for physical activity, documentation of counseling for
	Counseling for physical activity: documentation of counseling for      physical activity: an referred for physical activity during the
	physical activity or referral for physical activity during the
	measurement year. Documentation must include a note
	indicating the date and at least one of the following:
	<ul> <li>Discussion of current physical activity behaviors (e.g.,</li> </ul>
	exercise routine, participation in sports activities, exam for
	sports participation)
	<ul> <li>Checklist indicating physical activity was addressed</li> </ul>
	<ul> <li>Counseling or referral for physical activity</li> </ul>
	<ul> <li>Patient received education materials on physical activity</li> </ul>
	during face to face visit
	<ul> <li>Anticipatory guidance for physical activity</li> </ul>
	<ul> <li>Weight or obesity counseling</li> </ul>
	Practices may utilize the Physical Activity Counseling Value Set as a
	mechanism to achieve this component of the measure, but still must meet
	the above documentation requirements.
	The state of the s
	Services rendered for obesity or eating disorders may be used to meet
	criteria for the Counseling for Nutrition and Counseling for Physical
	Activity indicators if the specified documentation is present.
Denominator	All active patients 3-17 at the end of the measurement year with a
Statement:	documented encounter during the measurement year
Acceptable	
Exclusions:	
Exclusions.	during the measurement year
	Patients in hospice
Look Back	12 months
Period:	
Source:	HEDIS® (modified by OHIC to become an "all-or-nothing" measure,
	including the three sub-measures)

- Please indicate for what performance period you are reporting data for "Weight Assessment and Counseling for Nutrition and Physical Activity."
  - o October 1, 2019 September 30, 2020
  - Other (Please Specify)
- Enter the numerator for "Weight Assessment and Counseling for Nutrition and Physical Activity."
- Enter the denominator for "Weight Assessment and Counseling for Nutrition and Physical Activity."