



#### RESOURCE AND PATIENT MANAGEMENT SYSTEM

## **Clinical Reporting System**

(BGP)

# CRS Clinical Performance Measure Logic Manual for FY 2012 Clinical Measures

Version 12.0 December 2011

Office of Information Technology (OIT)
Division of Information Resource Management
Albuquerque, New Mexico

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### **Preface**

The Government Performance and Results Act (GPRA) requires federal agencies to report annually on how the agency measured up against the performance targets set in its annual Plan. The Indian Health Service (IHS) GPRA report includes measures for clinical prevention and treatment, quality of care, infrastructure, and administrative efficiency functions.

The IHS Clinical Reporting System (CRS) is a Resource and Patient Management System (RPMS) software application designed for national reporting as well as Area Office and local monitoring of clinical GPRA and developmental measures. CRS was first released for Fiscal Year (FY) 2002 performance measures (as GPRA+) and is based on a design by the Aberdeen Area (GPRA2000).

This manual contains the FY 2012 clinical performance measure definitions and logic for the CRS 2012 Version 12.0 software. CRS is the reporting tool used by the IHS Office of Planning and Evaluation to collect and report clinical performance results annually to the Department of Health and Human Services and to Congress.

Each year, an updated version of CRS software is released to reflect changes in the logic descriptions of the different denominators and numerators. Additional performance measures may also be added. Local facilities can run reports as often as they want to and can also use CRS to transmit data to their Area. The Area Office can use CRS to produce an aggregated Area report for either annual GPRA or Area Director Performance reports.

CRS produces reports on demand from local RPMS databases for both GPRA and developmental clinical measures that are based on RPMS data, thus eliminating the need for manual chart audits for evaluating and reporting clinical measures.

To produce reports with comparable data across every facility, the GPRA measures definitions was "translated" into programming code with the assistance of clinical subject matter experts. CRS uses predefined taxonomies to find data items in the RPMS Patient Care Component (PCC) to determine if a patient meets the performance measure criteria. Taxonomies contain groups of codes (e.g., diagnoses or procedures) or site-specific terms. Each performance measure has one or more defined denominators and numerators.

Administrative and clinical users can produce reports for selected measures at any time to:

• Identify potential data issues in their RPMS, i.e., missing or incorrect data.

- Monitor their site's performance against past national performance and upcoming agency goals.
- Identify specific areas where the facility is not meeting the measure in order to initiate business process or other changes.
- Quickly measure impact of process changes on performance measures.
- Identify areas meeting or exceeding measures to provide lessons learned.

#### Users of the RPMS CRS include:

- Area Office and site quality improvement staff
- Compliance Officers
- GPRA coordinators
- Clinical staff, such as physicians, nurses, nurse practitioners, and other providers
- Area Office directors
- Any staff involved with quality assurance initiatives
- Staff who run various CRS reports

## 1.0 Introduction

This manual provides information on the performance measure logic used by the Clinical Reporting System (CRS) Version 12.0 Selected Measures (Local) Report (Fiscal Year [FY] 2012 Clinical Performance Measures). For information on system setup, available reports and steps for running the reports, and performing Area Office functions, refer to the CRS Version 12.0 User Manual.

## 2.0 Performance Measure Logic

This section provides the following information for each performance measure topic:

- For Government Performance and Results Act (GPRA) measures, the measure description is provided as stated in the Indian Health Service (IHS) Annual Performance Report to Congress
- Definitions of all denominators and numerators for each performance measure topic
- Detailed description of the logic for the denominator and numerator, including specific codes, fields, taxonomies, and/or values searched
- Key changes to logic from the previous year, if any
- Description of which patients and information are contained on the patient list
- Performance measure source and past IHS performance, if any, and IHS or Healthy People (HP) 2020 targets for the performance measure
- Report examples
- Patient list examples

**Note:** All report examples and patient list examples used in this section were produced from "scrubbed" demonstration databases and do not represent individual patient data.

### 2.1 Performance Measure Logic Basics

#### 2.1.1 CRS Denominator Definitions

Each performance measure topic has one or more define denominators and numerators. The denominator is the total population that is being reviewed for a specific measure. For the National GPRA & Program Assessment Rating Tool (PART) Report, only one denominator for each topic is reported. These denominators are pre-defined, based on the Active Clinical Population definition. For the Selected Measures reports for local use (CRS Version 12.0 User Manual, Section 5.12), multiple denominators may be reported to provide a complete picture of clinical performance. There are also additional options available to further refine denominator definitions.

#### 2.1.1.1 Denominator Definitions for National GPRA Reporting

The Active Clinical population is the denominator definition used as the basis for *most* GPRA measures. This denominator was developed in FY 2003 specifically for clinical performance measures because it is more representative of the active clinical population.

Note: There are facilities that do not offer direct care. Patients in these facilities receive only Contract Health Services (CHS) and therefore do not meet the requirements of the Active Clinical population. A new site parameter, Contract Health Site Only, was added for these facilities in FY2006.

Prior to FY 2003, the GPRA User Population denominator definition was used for national reporting, similar to the agency's IHS User Population definition.

The *Active Clinical* population for the National GPRA & PART Report is defined by the following criteria:

- Patients with the name of "DEMO, PATIENT" or who are included in the Resource and Patient Management System (RPMS) Demo/Test Patient Search Template (DPST option located in the PCC Management Reports, Other section) will be automatically excluded from the denominator.
- Patient must have two visits to medical clinics in the past three years. At least one visit must be to one of the following core medical clinics:

01	General	24	Well Child
06	Diabetic	28	Family Practice
10	GYN	57	EPSDT
12	Immunization	70	Women's Health
13	Internal Medicine	80	Urgent Care
20	Pediatrics	89	Evening

The second visit can be either to one of the core medical clinics in the previous list or to one of the following additional medical clinics:

02	Cardiac	37	Neurology
03	Chest And TB	38	Rheumatology
05	Dermatology	49	Nephrology
07	ENT	50	Chronic Disease
08	Family Planning	69	Endocrinology
16	Obstetrics	75	Urology
19	Orthopedic	81	Men's Health Screening
23	Surgical	85	Teen Clinic
25	Other	88	Sports Medicine
26	High Risk	B8	Gastroenterology - Hepatology
27	General Preventive	В9	Oncology - Hematology
31	Hypertension	C3	Colposcopy
32	Postpartum		

- Patient must be alive on the last day of the report period.
- Patient must be American Indian/Alaska Native (AI/AN) (defined as Beneficiary 01). This data item is entered and updated during the patient registration process.
- Patient must reside in a community included in the site's "official" GPRA
  community taxonomy, defined as all communities of residence in the CHS
  catchment area specified in the community taxonomy specified by the user.

The *Active Clinical CHS Population* for National GPRA & PART Reports is defined as follows:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS Demo/Test Patient Search Template (DPST option located in the Patient Care Component [PCC] Management Reports, Other section) will be automatically excluded from the denominator.
- Patient must have two CHS visits in the three years prior to the end of the report period.
- Patient must be alive on the last day of the report period.
- Patient must be AI/AN (defined as Beneficiary 01). This item is entered and updated during the patient registration process.
- User must reside in a community included in the site's official GPRA community taxonomy, defined as all communities of residence in the CHS catchment area specified in the community taxonomy that is specified by the user.

The Active Clinical Behavioral Health Population for National GPRA & PART Reports is defined as follows:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS Demo/Test Patient Search Template (DPST option located in the Patient Care Component [PCC] Management Reports, Other section) will be automatically excluded from the denominator.
- Patient must have two behavioral health visits in the three years prior to the end of the report period.
- Patient must be alive on the last day of the report period.
- Patient must be AI/AN (defined as Beneficiary 01). This item is entered and updated during the patient registration process.
- User must reside in a community included in the site's official GPRA community taxonomy, defined as all communities of residence in the Urban Outreach & Referral catchment area specified in the community taxonomy that is specified by the user.

The *GPRA User Population* for the National GPRA & PART Report is defined by the following criteria:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS
  Demo/Test Patient Search Template (DPST option located in the PCC
  Management Reports, Other section) will be automatically excluded from the
  denominator.
- Patient must have been seen at least once in the three years prior to the end of the time period, regardless of the clinic type.
- Patient must be alive on the last day of the report period.
- Patient must be AI/AN (defined as Beneficiary 01). This data item is entered and updated during the patient registration process.
- Patient must reside in a community included in the site's "official" GPRA
  community taxonomy, defined as all communities of residence in the CHS
  catchment area specified in the community taxonomy specified by the user.

**Note:** The GPRA User Population definition is similar, but not identical, to the definition used by IHS headquarters (HQ) for annual user population statistics. GPRA "visits" are not required to be workload reportable as defined by IHS HQ.

#### 2.1.1.2 Denominator Definitions for Selected Measures Reports

In addition to the National GPRA & PART Report, CRS provides Selected Measures reports intended for local facility use for specific public health and/or performance improvement initiatives (CRS Version 12.0 User Manual, Section 5.12). Multiple denominators and numerators will be reported for each measure (e.g., *both* Active Clinical and GPRA User Population). Users have additional options to define the denominators as explained below.

The Active Clinical Population for Selected Measures (Local) Reports is defined as follows:

- Patients with name "DEMO, PATIENT" or who are included in the RPMS
  Demo/Test Patient Search Template (DPST option located in the PCC
  Management Reports, Other section) will be automatically excluded from the
  denominator.
- Patient must have two visits to medical clinics in the past three years. At least one visit must be to one of the following core medical clinics:

01	General	24	Well Child
06	Diabetic	28	Family Practice
10	GYN	57	EPSDT
12	Immunization	70	Women's Health
13	Internal Medicine	80 Urgent Care	
20	Pediatrics	89	Evening

The second visit can be either to one of the core medical clinics in the previous list or to one of the following additional medical clinics:

02	Cardiac	37	Neurology
03	Chest And TB	38	Rheumatology
05	Dermatology	49	Nephrology
07	ENT	50	Chronic Disease
08	Family Planning	69	Endocrinology
16	Obstetrics	75	Urology
19	Orthopedic	81	Men's Health Screening
23	Surgical	85	Teen Clinic
25	Other	88	Sports Medicine
26	High Risk	B8	Gastroenterology - Hepatology
27	General Preventive	В9	Oncology - Hematology
31	Hypertension	C3	Colposcopy

32 Postpartum

- Patient must be alive on the last day of the Report period.
- User defines population type: AI/AN patients only, non AI/AN, or both. This data item is entered and updated during the patient registration process.
- User defines general population: single community; group of multiple communities (community taxonomy); user-defined list of patient (patient panel); or all patients regardless of community of residence.

The *Active Clinical CHS Population* for Selected Measures (Local) Reports is defined as follows:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS Demo/Test Patient Search Template (DPST option located in the PCC Management Reports, Other section) will be automatically excluded from the denominator.
- Patient must have two CHS visits in the three years prior to the end of the Report Period.
- Patient must be alive on the last day of the report period.
- User defines population type: AI/AN patients only, non AI/AN, or both.
- User defines general population: single community; group of multiple communities (community taxonomy); user-defined list of patient (patient panel); or all patients regardless of community of residence.

The *Active Clinical Behavioral Health Population* for Selected Measures (Local) Reports is defined as follows:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS
  Demo/Test Patient Search Template (DPST option located in the PCC
  Management Reports, Other section) will be automatically excluded from the
  denominator.
- Patient must have two behavioral health visits in the three years prior to the end of the Report Period.
- Patient must be alive on the last day of the report period.
- User defines population type: AI/AN patients only, non AI/AN, or both.
- User defines general population: single community; group of multiple communities (community taxonomy); user-defined list of patient (patient panel); or all patients regardless of community of residence.

The *User Population* for Selected Measures (Local) reports is defined as follows:

- Patients with the name of "DEMO, PATIENT" or who are included in the RPMS
  Demo/Test Patient Search Template (DPST option located in the PCC
  Management Reports, Other section) will be excluded from the denominator
  automatically.
- Patient must have been seen at least once in the three years prior to the end of the time period, regardless of the clinic type.
- Patient must be alive on the last day of the report period.
- User defines population type: AI/AN patients only, non AI/AN, or both.
- User defines general population: single community, group of multiple communities (community taxonomy); user-defined list of patient (patient panel); or all patients regardless of community of residence.

#### 2.1.2 Performance Measure Logic Example

Cancer Screening: Pap Smear Rates: During FY 2012, achieve the tentative target rate of 59.5% for the proportion of female patients ages 21 through 64 without a documented history of hysterectomy who have had a Pap screen within the previous three years.

For CRS, the GPRA measure definition is defined as:

- Denominator (total number of patients evaluated): Active Clinical female patients ages 21 through 64, excluding those with documented history of hysterectomy. (The clinical owner of the measure has determined based on current medical guidelines that "eligible" women are defined as ages 21–64.)
- Numerator (those from the denominator who meet the criteria for the measure): patients with documented Pap smear in past three years or refusal in past year.

For the programmer, the Pap Smear measure is described in terms of the following logic:

- 1. Begin with the Active Clinical population definition.
  - Exclude any patients with the name of "DEMO, PATIENT."
  - Exclude any patient records that are included in the RPMS Demo/Test Patient Search Template.
  - Exclude any patients with a date of death in the Patient Registration file.
  - Exclude any patients who do *not* have value 01 (AI/AN) in the Beneficiary field in Patient Registration file.
  - Exclude any patients whose Community of Residence is not included in the site's defined GPRA Community Taxonomy for this report.

- For the remaining patients, search visit files for the three years prior to the selected report end date; exclude any patients whose visits do not meet the "two medical clinics" definition; *or*, for facilities with the CHS-Only site parameter set to Yes, exclude any patients who do not have two CHS visits in the past three years.
- 2. From these patients, identify the subset that are female and that are at least age 21 on the first day of the current report period and less than age 65 on the last day of the report period.
- 3. Exclude patients with documented hysterectomy by searching the V Procedure file for procedure codes 68.4–68.8 or V CPT for CPT codes 51925, 56308 (old code), 57540, 57545, 57550, 57555, 57556, 58150, 58152, 58200–58294, 58548, 58550–58554, 58570–58573, 58951, 58953–58954, 58956, 59135 or V POV 618.5, V88.01, V88.03 or Women's Health procedure called Hysterectomy any time before the end of the report period.
- 4. For these patients (the denominator), check for a Pap smear in the past three years in the following order:
  - a. Check V Lab for a lab test called Pap Smear and for any site-populated pap smear lab test documented in the BGP PAP SMEAR TAX taxonomy; *or*
  - b. Check V Lab for any LOINC code listed in the predefined BGP PAP LOINC CODES taxonomy (see the *CRS Technical Manual* for specific codes); *or*
  - c. Check the Purpose of Visit file (V POV) for: a diagnosis of: V67.01 Follow-up Vaginal Pap Smear, V76.2-Screen Mal Neop-Cervix, V72.32 Encounter for Pap Cervical Smear to Confirm Findings of Recent Normal Smear Following Initial Abnormal Smear, V72.3 Gynecological Examination, Pap Cervical Smear as Part of General Gynecological Exam, Pelvic Exam (annual) (periodic) (old code, to be counted for visits prior to 10/1/04 only), V76.47 Vaginal Pap Smear for Post-Hysterectomy Patients, 795.0\*, 795.10–16, 795.19; or
  - d. Check V Procedures for a procedure of 91.46; or
  - e. Check V CPT for the following CPT codes: 88141–88167, 88174–88175, G0123, G0124, G0141, G0143–G0145, G0147, G0148, P3000, P3001, Q0091; *or*
  - f. Check the Women's Health Tracking package for documentation of a procedure called Pap Smear and where the result does *not* have "ERROR/DISREGARD".

If a visit with any of the specified codes is found, the patient is considered to have met the measure, and the program checks the next patient.

#### 2.1.3 Age Ranges

Unless otherwise noted, for the purposes of CRS reports, the age of a patient is calculated at the beginning of the report period. For example, for a report period of July 1, 2011 through June 30, 2012, Jane Doe is defined as age 74 if her birth date is June 10, 1937, even though she becomes age 75 during the report period.

#### 2.1.4 Standard Health Care Codes

#### 2.1.4.1 Current Procedural Terminology Codes

One of several code sets used by the healthcare industry to standardize data, and allow for comparison and analysis. Current Procedural Terminology (CPT) was developed and is updated annually by the American Medical Association, and is widely used in producing bills for services rendered to patients. CPTs include codes for diagnostic and therapeutic procedures, and specify information that differentiates the codes based on cost. CPT codes are the most widely accepted nomenclature in the United States for reporting physician procedures and services for federal and private insurance third-party reimbursement. CRS searches for CPT and other codes as specified in the logic definition to determine if a patient meets a denominator or numerator definition.

#### 2.1.4.2 International Classification of Disease Codes

One of several code sets used by the healthcare industry to standardize data. The International Classification of Disease (ICD) is an international diagnostic coding scheme. In addition to diseases, ICD also includes several families of terms for medical-specialty diagnoses, health status, disablements, procedure, and reasons for contact with healthcare providers. IHS currently uses ICD, Ninth Revision (ICD-9) for coding. CRS searches for ICD and other codes as specified in the logic definition, to determine if a patient meets a denominator or numerator definition.

#### 2.1.4.3 Logical Observation Identifiers Names and Codes

Logical Observation Identifiers Names and Codes (LOINC®). A standard coding system originally initiated for laboratory values, the system is being extended to include nonlaboratory observations (electrocardiograms, vital signs, etc.). Standard code sets are used to define individual tests and mitigate variations in local terminologies for laboratory and other healthcare procedures, for example, Glucose or Glucose Test. IHS began integrating LOINC values into RPMS in several pilot sites in 2002.

Refer to the CRS Version 12.0 Technical Guide for a list of specific LOINC codes included in each LOINC taxonomy.

### 2.2 Diabetes Related Measure Topics

#### 2.2.1 Diabetes Prevalence

#### **Denominators**

All *User Population patients*. Broken down by gender and age groups (<15, 15–19, 20–24, 25–34, 35–44, 45–54, 55–64, >64 yrs).

#### **Numerators**

Anyone diagnosed with Diabetes at any time before the end of the Report Period.

Anyone diagnosed with Diabetes during the Report Period.

#### **Logic Description**

Age is calculated at the beginning of the Report Period.

Diabetes definition: At least one diagnosis of 250.00–250.93 recorded in V POV file.

#### **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of diabetic patients with most recent diagnosis.

#### **Measure Source**

HP 2010 5-2, 5-3

#### **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	12.8%
IHS FY 2010 Performance	12.0%
IHS FY 2009 Performance	12.0%
IHS FY 2008 Performance	12.0%
IHS FY 2007 Performance	11.0%
IHS FY 2006 Performance	11.0%
IHS FY 2005 Performance	11.0%
IHS FY 2004 Performance	10.0%

November 25, 2012 Page 1 \*\*\* IHS 2012 Selected Measures with Community Specified Report \*\*\* DEMO INDIAN HOSPITAL Report Period: Jan 01, 2012 to Dec 31, 2012 Previous Year Period: Jan 01, 2011 to Dec 31, 2011 Baseline Period: Jan 01, 2000 to Dec 31, 2000 Diabetes Prevalence Denominator(s): - All User Population patients. Broken down by gender and by age groups: < 15, 15-19, 20-24, 25-34, 35-44, 45-54, 55-64, > 64. Numerator(s): - Anyone diagnosed with Diabetes at any time before the end of the Report - Anyone diagnosed with Diabetes during the Report Period. Logic: Age is calculated at the beginning of the Report Period. Diabetes diagnosis is defined as at least one diagnosis 250.00-250.93 recorded in the V POV file. Performance Measure Description: Continue tracking (i.e., data collection and analyses) Area age-specific diabetes prevalence rates to identify trends in the age-specific prevalence of diabetes (as a surrogate marker for diabetes incidence) for the AI/AN population. Past Performance and/or Target: IHS Performance: FY 2011 - 12.8%, FY 2010 - 12%, FY 2009 - 12%, FY 2008 - 12%, FY 2007 - 11%, FY 2006 - 11%, FY 2005 - 11%, FY 2004 - 10% Source: HP 2010 5-2, 5-3 Diabetes Prevalence REPORT % PREV YR % CHG from BASE % CHG from PREV YR % PERIOD BASE % PERIOD PERIOD 2,896 2,456 2,346 # User Pop # w/ any DM DX 265 9.2 243 9.9 -0.7 197 8.4 +0.8 # w/ DM DX w/in past year 173 6.0 146 5.9 +0.0 100 4.3 +1.7 1,368 1,109 # Male User Pop 1,152 # w/ any DM DX 114 8.3 104 9.0 -0.7 72 6.5 +1.8 # w/DM DX w/in past year 82 6.0 78 6.8 -0.8 48 4.3 +1.7 # Female User Pop 1,528 1,304 1,237 # w/ any DM DX 151 9.9 139 10.7 125 10.1 -0.8 -0.2 # w/ DM DX w/in past year 91 6.0 68 5.2 +0.7 52 4.2 +1.8

Figure 2-1: Sample Summary Report, Diabetes Prevalence Topic

DU November 25, 2012 Page 2  *** IHS 2012 Selected Measures with Community Specified Report ***									
DEMO INDIAN HOSPITAL									
	Report Period: Jan 01, 2012 to Dec 31, 2012 Previous Year Period: Jan 01, 2011 to Dec 31, 2011								
		iod: J							
Diabetes Prevalence (co	on't)								
		TOTAL U	-	-	bution				
	<15	15-19	_				55-64	>64 yrs	
CURRENT REPORT PERIOD									
	730	239	261	403	389	390	262	222	
Total # User Pop # w/ DM DX ever	1	3	7	37	57	67	50	43	
% w/ DM DX ever	0.1	1.3	2.7	9.2	14.7	17.2		19.4	
# w/DM DX in past yr	0	2	3	14	42	49	34	29	
% w/DM DX in past yr	0.0	0.8	1.1	3.5	10.8	12.6			
PREVIOUS YEAR PERIOD									
Total # User Pop	711	227	243	352	316	277	181	149	
		4	9	31	54	57			
# w/ DM DX ever % w/ DM DX ever	0.4	1.8	3.7	8.8	54 17.1	20.6	24.9	26.8	
# w/DM DX in past yr	1	3	3	9	33	37	31	29	
# w/DM DX in past yr % w/DM DX in past yr	0.1	1.3	1.2	2.6	10.4	13.4	17.1	19.5	
CHANGE FROM PREV YR %									
w/ DM DX ever	-0.3	-0.5	-1.0	+0.4	-2.4	-3.4	-5.8	-7.5	
w/DM DX in past yr	-0.1	-0.5	-0.1	+0.9	+0.4	-0.8	-4.1	-6.4	
BASELINE REPORT PERIOD									
Total # User Pop	787	208	217	329	293	228	141	143	
# w/ DM DX ever	2	4	12	20	38	46	31	44	
# w/ DM DX ever % w/ DM DX ever	0.3	1.9	5.5	6.1	13.0	20.2	22.0	30.8	
# w/DM DX in past yr	2	1	3	7	18	21	20	28	
# w/DM DX in past yr % w/DM DX in past yr	0.3	0.5	1.4	2.1	6.1	9.2	14.2	19.6	
CHANGE FROM BASE YR %									
w/ DM DX ever	-0.1	-0.7	-2.8	+3.1	+1.7	-3.0	-2.9	-11.4	
w/DM DX in past yr	-0.3	+0.4	-0.2	+1.3	+4.7	+3.4	-1.2	-6.5	

Figure 2-2: Sample Age Breakdown Page, Diabetes Prevalence Topic

```
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*** IHS 2012 Clinical Performance Measure Patient List ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Entire Patient List

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
```

```
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Diabetes Prevalence: List of diabetic patients with most recent diagnosis
                          COMMUNITY
                                       SEX AGE
PATIENT NAME
                     HRN
DENOMINATOR
                           NUMERATOR
PATIENT1, DEBORAH 000001 COMMUNITY #1 F 20
                          02/01/12 POV 250.00
PATIENT2, TARA 000002 COMMUNITY #1 F 21
                          05/24/12 POV 250.00
PATIENT3, BOBBIE 000003 COMMUNITY #1 F 28
                          03/30/12 POV 250.00
PATIENT4, WINONA 000004 COMMUNITY #1 F 37
                          04/30/12 POV 250.00
PATIENTS, NADINE
UP
                     000005 COMMUNITY #1 F 44
                          03/19/12 POV 250.00
PATIENT6, RUTH
                     000006 COMMUNITY #1 F 44
                     03/19/12 POV 250.00
```

Figure 2-3: Sample Patient List, Diabetes Prevalence, Patients with Diabetes Diagnosis

#### 2.2.2 Diabetes Comprehensive Care

#### **Denominators**

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two diabetes mellitus- (DM-) related visits ever.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two diabetes mellitus- (DM-) related visits ever, without a documented history of bilateral foot amputation or two separate unilateral foot amputations.

#### **Numerators**

Patients with hemoglobin A1c documented during the Report Period, regardless of result

Patients with blood pressure (BP) documented during the Report Period

Patients with controlled BP, defined as < 130/80, i.e., the mean systolic value is less than 130 and the mean diastolic value is less than 80

Patients with low-density lipoprotein (LDL) completed during the Report Period, regardless of result

Patients with nephropathy assessment, defined as an estimated GFR with result and a quantitative urinary protein assessment during the Report Period *or* with the evidence of diagnosis and/or treatment of end-stage renal disease (ESRD) at any time before the end of the Report Period

Patients receiving a qualified retinal evaluation during the Report Period

**Note**: This numerator does *not* include refusals.

Patients with diabetic foot exam during the Report Period

**Note**: This numerator does *not* include refusals.

Patients with comprehensive diabetes care (Documented A1c and Blood Pressure and LDL and Nephropathy Assessment and Retinal Exam and Diabetic Foot Exam).

#### **Logic Description**

*Diabetes definition*: First Purpose of Visit (POV) 250.00–250.93 recorded in the POV file prior to the Report Period.

*A1c definition*: Searches for most recent A1c test with a result during the Report Period. If none found, CRS searches for the most recent A1c test without a result. A1c defined as: CPT 83036, 83037, 3044F–3046F, 3047F (old code); LOINC taxonomy; or site-populated taxonomy DM AUDIT HGB A1C TAX.

*BP documented definition*: Having a minimum of two BPs documented on non-Emergency Room (ER) visits during the Report Period.

CRS uses mean of last three BPs documented on non-ER visits during the Report Period. If three BPs are not available, uses mean of last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) BPs and dividing by three (or two).

If CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on a non-ER visit during the Report Period.

Controlled BP definition: CRS uses a mean, as described above. If the mean systolic and diastolic values do not both meet the criteria for controlled, then the value is considered not controlled.

BP documented and Controlled BP: If CRS is not able to calculate a mean BP from BP measurements, it will search for the most recent of any of the following CPT codes documented on non-ER visits during the Report Period: BP Documented: 0001F or 2000F; *or* Systolic 3074F, 3075F, or 3077F *with* Diastolic: 3078F, 3079F, or 3080F. The systolic and diastolic values do not have to be recorded on the same day. If there are multiple values on the same day, the CPT indicating the lowest value will be used. The following combination represents BP < 130/80 and will be included in the Controlled BP numerator: CPT 3074F *and* 3078F. All other combinations *will not* be included in the Controlled BP numerator.

*LDL definition*: Finds last test done during the Report Period; defined as CPT 80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F; LOINC taxonomy; site-populated taxonomy DM AUDIT LDL CHOLESTEROL TAX.

Nephropathy assessment definition: (1) Estimated GFR with result during the report period, defined as any of the following: (A) Site-populated taxonomy BGP GPRA ESTIMATED GFR TAX or (B) LOINC taxonomy, and (2) Quantitative Urinary Protein Assessment during the Report Period, defined as any of the following: (A) CPT 82042, 82043, or 84156; (B) LOINC taxonomy; or (C) site-populated taxonomy BGP QUANT URINE PROTEIN.

**Note**: Be sure and check with your laboratory supervisor that the names you add to your taxonomy reflect quantitative test values.

(3) ESRD diagnosis/treatment defined as any of the following ever: (A) V CPT 36145, 36147, 36800, 36810, 36815, 36818, 36819, 36820, 36821, 36831–36833, 50300, 50320, 50340, 50360, 50365, 50370, 50380, 90951–90970 or old codes 90918–90925, 90935, 90937, 90939 (old code), 90940, 90945, 90947, 90989, 90993, 90997, 90999, 99512, G0257, G0308–G0327 (old codes), G0392 (old code), G0393 (old code), or S9339; (B) V POV 585.5, 585.6, V42.0, V45.1 (old code), V45.11, V45.12, or V56.\*; (C) V Procedure 38.95, 39.27, 39.42, 39.43, 39.53, 39.93–39.95, 54.98, or 55.6\*.

Qualified retinal evaluation definition: (1) diabetic retinal exam or (2) other eye exam. The following methods are qualifying for this measure: (1) dilated retinal evaluation by an optometrist or ophthalmologist, or (2) seven standard fields stereoscopic photos (ETDRS) evaluated by an optometrist or ophthalmologist, or (3) any photographic method formally validated to seven standard fields (ETDRS).

- Diabetic Retinal Exam: Any of the following during the Report Period: (1) Exam Code 03 Diabetic Eye Exam (dilated retinal examination or validated photographic equivalent), (2) CPT 2022F Dilated retinal eye exam; 2024F Seven standard field stereoscopic photos with interpretation by an ophthalmologist or optometrist; 2026F Eye imaging validated to match the diagnosis from seven standard field stereoscopic photos; S0620 Routine ophthalmological examination including refraction; new patient; S0621 Routine ophthalmological examination including refraction; established patient; S3000 Diabetic indicator; retinal eye exam, dilated, bilateral.
- Other Eye Exam: (1) Non-DNKA (did not keep appointment) visits to ophthalmology, optometry or validated teleophthalmology retinal evaluation clinics or (2) non-DNKA visits to an optometrist or ophthalmologist. Searches for the following codes in the following order: Clinic Codes A2, 17, 18, 64; Provider Code 24, 79, 08; CPT 67028, 67038 (old code), 67039, 67040, 92002, 92004, 92012, 92014; POV V72.0; Procedure 95.02.

Diabetic foot exam definition: (1) Exam Code 28 Diabetic Foot Exam, Complete; (2) non-DNKA visit with a podiatrist (Provider Codes 33, 84 or 25), (3) non-DNKA visit to Podiatry Clinic (Clinic Code 65), or (4) CPT 2028F.

Bilateral foot amputation definition: CPT 27290.50-27295.50, 27590.50-27592.50, 27598.50, 27880.50-27882.50 (.50 modifier indicates bilateral)

*Unilateral foot amputation definition:* Must have 2 separate occurrences for either CPT or Procedure codes on 2 different dates of service: (1) CPT 27290-27295, 27590-27592, 27598, 27880-27882, or (2) ICD Procedure codes 84.10, 84.13-84.19.

#### **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

#### **Patient List Description**

List of diabetic patients with documented tests, if any.

#### **Measure Source**

Foot Exam: HP 2020 D-9

#### **Measure Past Performance and Long-Term Targets**

Target	Percent
HP 2020 goal for foot exam	74.8%

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\*\*\* IHS 2012 Selected Measures with Community Specified Report \*\*\*

DEMO INDIAN HOSPITAL Report Period: Jan 01, 2012 to Dec 31, 2012 Previous Year Period: Jan 01, 2011 to Dec 31, 2011 Baseline Period: Jan 01, 2000 to Dec 31, 2000								
Diabetes Comprehensiv	e Care							
	REPORT PERIOD				CHG from PREV YR %			CHG from BASE %
Active Diabetic Pts	144		97			87		
# w/Comp Diabetes Care # w/Alc done	10	6.9	1	1.0	+5.9	0	0.0	+6.9
w/ or w/o result	92	63.9	70	72.2	-8.3	52	59.8	+4.1
# w/ BPs documented # w/Controlled BP		83.3		80.4			85.1	
<130/80	28	19.4	20	20.6	-1.2	13	14.9	+4.5
# w/ LDL done # w/ est GFR & quant UP assmt or	78	54.2	46	47.4	+6.7	23	26.4	+27.7
w/ESRD	53	36.8	11	11.3	+25.5	6	6.9	+29.9
# w/Retinal Evaluation	n							
-No Refusals	54	37.5	39	40.2	-2.7	44	50.6	-13.1
Active Diabetic Pts w/o Hx of Bilateral								
Amputation	137		97			87		
# w/Diabetic Foot Exa -No Refusals		14.6	18	18.6	-4.0	16	18.4	-3.8

Figure 2-4: Sample Summary Report, Diabetes Comprehensive Care Topic

```
***** CONFIDENTIAL PATIENT INFORMATION, COVERED BY THE PRIVACY ACT *****
  DU
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                                                                    Page 11
         *** IHS 2011 Clinical Performance Measure Patient List ***
                            DEMO INDIAN HOSPITAL
                 Report Period: Jan 01, 2012 to Dec 31, 2012
                            Entire Patient List
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Diabetes Comprehensive Care: List of diabetic patients with documented tests,
if any.
PATIENT NAME
                     HRN COMMUNITY
                                          SEX AGE
DENOMINATOR
                           NUMERATOR
PATIENT1, DEBORAH 000001 COMMUNITY #1 F 45
                            Alc: 02/28/12 6.6; BPs: 133/82 UNC; LDL: 10/28/12 119;
EYE: 01/07/12 Cl 18
```

```
PATIENT2, TARA 000002 COMMUNITY #1 F 51

AD BP: <130/80: BPs: 118/61; ESRD: 03/03/12 90951; FOOT

AMPUTATION

PATIENT3, BOBBIE 000003 COMMUNITY #1 F 52

AD A1c: 04/09/12 6.5; BPs: 138/66 UNC; GFR: 04/09/12 &

QUANT UP: 03/31/12 QUANT URINE PROTEIN; EYE: 03/30/12 Cl 18; FOOT EXAM: 01/07/12 Cl 65
```

Figure 2-5: Sample Patient List, Diabetes Comprehensive Care

#### 2.2.3 Diabetes: Glycemic Control

#### **GPRA Measure Description, Poor Glycemic Control**

During FY 2012, achieve the target rate of 18.6% for the proportion of patients with diagnosed diabetes who have poor glycemic control (defined as A1c > 9.5).

#### **GPRA Measure Description, Ideal Glycemic Control**

During FY 2012, achieve the target rate of 32.7% for the proportion of patients with diagnosed diabetes who have ideal glycemic control (defined as A1c < 7).

#### **Denominators**

All *User Population patients* diagnosed with diabetes prior to the report period.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. (GPRA Denominator)

Active Adult Diabetic patients, defined by meeting the following criteria: (1) who are age 19 or older at the beginning of the Report Period, (2) whose first ever DM diagnosis occurred prior to the Report Period; (3) who had at least two DM related visits ever; (4) at least one encounter with DM POV in a primary clinic with a primary provider during the Report Period; and (5) never have had a creatinine value greater than five.

#### **Numerators**

Number of patients with a Hemoglobin A1c documented during the Report Period, regardless of result.

*Poor Control*: Total of Poor and Very Poor Control: Patients with A1c greater than (>) 9.5. (GPRA Numerator)

*Very Poor Control*: Patients with A1c equal to or greater than (=>) 12.

*Poor Control*: Patients with A1c greater than (>) 9.5 and less than (<) 12.

Fair Control: Patients with A1c equal to or greater than (=>) 8 and less than or equal to (<=) 9.5.

Good Control: Patients with A1c equal to or greater than (=>) 7 and less than (<) 8.

*Ideal Control*: Patients with A1c less than (<) 7. (GPRA Numerator)

Without Result: Patients with A1c documented but no value.

#### **Logic Description**

*Diabetes definition*: First Purpose of Visit 250.00–250.93 recorded in the V POV file prior to the Report Period.

Hemoglobin A1c definition: Searches for most recent A1c test with a result during the Report Period. If more than one A1c test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If an A1c test with a result is not found, CRS searches for the most recent A1c test without a result. Without result is defined as A1c documented but with no value.

CRS uses the following definitions:

	CPT Codes	LOINC Codes	Taxonomy
Creatinine (for Active Adult Diabetic denominator)	CPT codes are not included since they do not store the result, which is used in this topic	Yes	DM AUDIT CREATININE TAX
Hemoglobin A1c	83036, 83037, 3044F-3046F, 3047F (old code) <b>Note</b> : CPT 3044F represents A1c <7 and will be included in the Ideal Control numerator.	Yes	DM AUDIT HGB A1C TAX

In the CPT Codes column, specific LOINC codes used CRS are located in the *CRS Technical Manual*.

#### **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of diabetic patients with most recent A1c value, if any.

#### **Measure Source**

HEDIS; HP 2020 D-11, D-5

## **Measure Past Performance and Long-Term Targets Hemoglobin A1c Documented**

Performance	Percent
IHS FY 2011 Performance	83.0%
IHS FY 2010 Performance	82.0%
IHS FY 2009 Performance	80.0%
IHS FY 2008 Performance	79.0%
IHS FY 2007 Performance	79.0%
IHS FY 2006 Performance	79.0%
IHS FY 2005 Performance	78.0%
IHS FY 2004 Performance	77.0%
IHS FY 2003 Performance	75.0%
IHS FY 2002 Performance	73.0%
HP 2020 Goal	71.1%

#### **Poor Glycemic Control**

Performance	Percent
IHS FY 2011 Performance	19.1%
IHS FY 2010 Performance	18.0%
IHS FY 2009 Performance	18.0%
IHS FY 2008 Performance	17.0%
IHS FY 2007 Performance	16.0%
IHS FY 2006 Performance	16.0%
IHS FY 2005 Performance	15.0%
IHS FY 2004 Performance	17.0%
IHS FY 2003 Performance	17.0%
IHS FY 2002 Performance	18.0%

#### **Ideal Glycemic Control**

Performance	Percent
IHS FY 2011 Performance	31.9%
IHS FY 2010 Performance	32.0%
IHS FY 2009 Performance	31.0%

Performance	Percent
IHS FY 2008 Performance	32.0%
IHS FY 2007 Performance	31.0%
IHS FY 2006 Performance	31.0%
IHS FY 2005 Performance	30.0%
IHS FY 2004 Performance	27.0%
IHS FY 2003 Performance	28.0%
IHS FY 2002 Performance	25.0%
HP 2020 Goal	58.9%

DU *** IHS 2012	Selected	Measu	ember 25, res with	Commu	nity Speci:	fied Rep		Page 11 **	
Re	eport Per				ац Dec 31, 20	012			
Previo	ous Year	Period	: Jan 01	, 201	l to Dec 3	1, 2011			
Bas	seline Pe	riod:	Jan 01,	2000 1	to Dec 31,	2000			
Diabetes: Glycemic (	Control								
	REPORT PERIOD				CHG from PREV YR %				
User Pop w/ DM DX prior to report									
end date	229		202			180			
# w/Alc done w/									
or w/o result	95	41.5	72	35.6	+5.8	53	29.4	+12.0	
# w/A1c =>12	3	1.3	1	0.5	+0.8	3	1.7	-0.4	
# w/A1c >9.5									
and <12	15	6.6	3	1.5	+5.1	8	4.4	+2.1	
# w/A1c =>8									
and =<9.5	12	5.2	19	9.4	-4.2	10	5.6	-0.3	
# w/A1c=>7	1.2		1.6	7 0	2.2	7	2 0	. 1 0	
and <8 # w/A1c <7		5.7 16.6		7.9		23	3.9		
# W/Alc </td <td>30</td> <td>10.0</td> <td>33</td> <td>10.3</td> <td>+0.3</td> <td>43</td> <td>12.0</td> <td>+3.0</td> <td></td>	30	10.0	33	10.3	+0.3	43	12.0	+3.0	
w/o Result	14	6.1	0	0.0	+6.1	2	1.1	+5.0	
Active Diabetic Pts									
(GPRA)	144		97			87			
# w/Alc done w/									
or w/o result	92	63.9	70	72.2	-8.3	52	59.8	+4.1	
# w/Alc									
> 9.5 (GPRA)		12.5		4.1			12.6		
# w/A1c =>12	3	2.1	1	1.0	+1.1	3	3.4	-1.4	
# w/A1c >9.5 and < 12	1 -	10.4	2	3.1	+7.3	0	9.2	+1.2	
and < 12 # w/Alc =>8	15	10.4	3	3.1	+ / . 3	8	9.2	+1.2	
and =<9.5	12	8.3	19	19.6	-11.3	10	11.5	-3.2	
# w/A1c=>7	12	0.5	10	17.0	11.5	10		5.2	

CRS Clinical Performance Measure Logic Manual for FY 2012 Clinical Measures
December 2011

13	9.0	16	16.5	-7.5	7	8.0	+1.0
35	24.3	31	32.0	-7.7	22	25.3	-1.0
14	9.7	0	0.0	+9.7	2	2.3	+7.4
102		75			63		
71	69.6	61	81.3	-11.7	46	73.0	-3.4
3	2.9	1	1.3	+1.6	3	4.8	-1.8
13	12.7	2	2.7	+10.1	7	11.1	+1.6
11	10.8	18	24.0	-13.2	8	12.7	-1.9
10	9.8	12	16.0	-6.2	6	9.5	+0.3
28	27.5	28	37.3	-9.9	22	34.9	-7.5
6	5.9	0	0.0	+5.9	0	0.0	+5.9
	35 14 102 71 3 13 11 10 28	71 69.6 3 2.9 13 12.7 11 10.8 10 9.8 28 27.5	35 24.3 31 14 9.7 0  102 75  71 69.6 61 3 2.9 1 13 12.7 2 11 10.8 18 10 9.8 12 28 27.5 28	35 24.3 31 32.0 14 9.7 0 0.0  102 75  71 69.6 61 81.3 3 2.9 1 1.3  13 12.7 2 2.7  11 10.8 18 24.0  10 9.8 12 16.0 28 27.5 28 37.3	35 24.3 31 32.0 -7.7 14 9.7 0 0.0 +9.7  102 75  71 69.6 61 81.3 -11.7 3 2.9 1 1.3 +1.6  13 12.7 2 2.7 +10.1  11 10.8 18 24.0 -13.2  10 9.8 12 16.0 -6.2 28 27.5 28 37.3 -9.9	35       24.3       31       32.0       -7.7       22         14       9.7       0       0.0       +9.7       2         102       75       63         71       69.6       61       81.3       -11.7       46         3       2.9       1       1.3       +1.6       3         13       12.7       2       2.7       +10.1       7         11       10.8       18       24.0       -13.2       8         10       9.8       12       16.0       -6.2       6         28       27.5       28       37.3       -9.9       22	35       24.3       31       32.0       -7.7       22       25.3         14       9.7       0       0.0       +9.7       2       2.3         102       75       63         71       69.6       61       81.3       -11.7       46       73.0         3       2.9       1       1.3       +1.6       3       4.8         13       12.7       2       2.7       +10.1       7       11.1         11       10.8       18       24.0       -13.2       8       12.7         10       9.8       12       16.0       -6.2       6       9.5         28       27.5       28       37.3       -9.9       22       34.9

Figure 2-6: Sample Report, Diabetes: Glycemic Control Topic

**** CONFIDENTIAL PA	ATIENT INFORMATION, COVERED BY THE PRIVACY ACT *****  November 25, 2012 Page 16
	D11 Clinical Performance Measure Patient List ***  DEMO INDIAN HOSPITAL  Port Period: Jan 01, 2012 to Dec 31, 2012  Entire Patient List
PREG=Pregnant Female; CHD-Active Coronary H	re Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; Heart Disease; HR=High Risk Patient
PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR
PATIENT1, DEBORA UP, AD, AAD	000001 COMMUNITY #1  F 45 03/28/12 Alc: 6.6
PATIENT2, TARA UP, AD, AAD	
PATIENT3, BOBBIE UP, AD, AAD	000003 COMMUNITY #1 F 52 04/09/12 Alc: 6.5
PATIENT4, WINONA UP	000004 COMMUNITY #1 F 53
PATIENT5, NADINE UP, AD, AAD	000005 COMMUNITY #1 F 61 02/01/12 Alc: 6.5
PATIENT6, RUTH UP	000006 COMMUNITY #1 F 64

Figure 2-7: Sample Patient List, Diabetes: Glycemic Control

#### 2.2.4 Diabetes: Blood Pressure Control

#### **GPRA Measure Description**

During FY 2012, achieve the target rate of 38.7% for the proportion of patients with diagnosed diabetes who have achieved blood pressure control (defined as <130/80).

#### **Denominators**

All *User Population patients* diagnosed with diabetes prior to the Report Period.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. (GPRA Denominator)

Active Adult Diabetic patients, defined by meeting the following criteria: (1) who are age 19 or older at the beginning of the Report Period, (2) whose first ever DM diagnosis occurred prior to the Report Period; (3) who had at least two DM-related visits ever, (4) at least one encounter with DM POV in a primary clinic with a primary provider during the Report Period; and (5) never have had a creatinine value greater than five.

#### **Numerators**

Patients with BP documented during the report period.

Patients with controlled BP, defined as < 130/80, i.e., the mean systolic value is less than 130 and the mean diastolic value is less than 80 (GPRA Numerator).

Patients with BP that is not controlled.

#### **Logic Description**

*Diabetes definition*: First DM POV 250.00–250.93 recorded in the V POV file prior to the Report Period.

BP documented definition: CRS uses mean of last three BPs documented on non-ER visits during the Report Period. If three BPs are not available, uses mean of the last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) blood pressures and dividing by three (or two). If the systolic and diastolic values do not both meet the criteria for controlled, then the value is considered not controlled.

If CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F-3080F or POV V81.1 documented on a non-ER visit during the Report Period.

Controlled BP definition: CRS uses a mean, as described above where BP is <130/80. If the mean systolic and diastolic values do not both meet the criteria for controlled, then the value is considered not controlled.

BP documented and Controlled BP: If CRS is not able to calculate a mean BP from blood pressure measurements, it will search for the most recent of any of the following codes documented on non-ER visits during the Report Period: BP Documented: CPT 0001F or 2000F or POV V81.1; or Systolic: CPT 3074F, 3075F or 3077F with Diastolic: CPT 3078F, 3079F, or 3080F. The systolic and diastolic values do not have to be recorded on the same day. If there are multiple values on the same day, the CPT indicating the lowest value will be used. The following combination represents BP <130/80 and will be included in the Controlled BP numerator: CPT 3074F and 3078F. All other combinations will not be included in the Controlled BP numerator.

CRS uses the following definition:

	CPT Codes	LOINC Codes	Taxonomy
Creatinine (for Active Adult Diabetic denominator)	CPT codes are not included since they do not store the result, which is used in this topic	Yes	DM AUDIT CREATININE TAX

In the LOINC Codes column, specific LOINC codes by CRS are location in the CRS Technical Manual.

#### **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

#### **Patient List Description**

List of diabetic patients with BP value, if any.

#### Measure Source

HP 2020 D-7

## Measure Past Performance and Long-Term Targets Controlled BP

Performance	Percent
IHS FY 2011 Performance	37.8%
IHS FY 2010 Performance	38.0%

Performance	Percent
IHS FY 2009 Performance	37.0%
IHS FY 2008 Performance	38.0%
IHS FY 2007 Performance	39.0%
IHS FY 2006 Performance	37.0%
IHS FY 2005 Performance	37.0%
IHS FY 2004 Performance	35.0%
IHS FY 2003 Performance	37.0%
IHS FY 2002 Performance	36.1%
HP 2020 Goal	57.0%

#### **BP Assessed**

Performance	Percent
IHS FY 2011 Performance	87.9%
IHS FY 2010 Performance	89.0%
IHS FY 2009 Performance	88.0%
IHS FY 2008 Performance	89.0%
IHS FY 2005 Performance	89.0%

DU November 25, 2012 Page 13  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Diabetes: Blood Pres	REPORT	%			CHG from PREV YR %				
User Pop w/ DM DX pr. to report period			202			180			
# w/ BPs Documented # w/controlled BP	132	57.6	88	43.6	+14.1	84	46.7	+11.0	
< 130/80 # w/Not controlled BP					+2.1		10.0 36.7		
Active Diabetic Pts (GPRA)	144	23.7	97	32.,	. 12.0	87	30.7	. , , 0	
# w/ BPs Documented # w/Controlled BP	120	83.3	78	80.4	+2.9	74	85.1	-1.7	

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< 130/80 (GPRA)	28	19.4	20	20.6	-1.2	13	14.9	+4.5	
# w/Not controlled BP	92	63.9	58	59.8	+4.1	61	70.1	-6.2	
Active Adult Diabetic Patients	102		75			63			
# w/ BPs Documented # w/Controlled BP	82	80.4	61	81.3	-0.9	56	88.9	-8.5	
< 130/80	20	19.6	14	18.7	+0.9	8	12.7	+6.9	
# w/Not controlled BP	62	60.8	47	62.7	-1.9	48	76.2	-15.4	

Figure 2-8: Sample Report, Diabetes: Blood Pressure Control Topic

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*** IHS 2011 Clinical Performance Measure Patient List ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Entire Patient List							
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient  Diabetes: Blood Pressure Control: List of diabetic patients with BP value, if any.							
PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR						
PATIENT1, DEBORAH UP, AD, AAD	133/82 UNC						
PATIENT2, TARA	000002 COMMUNITY #1  F  51 3074F/3080F UNC						
UP,AD,AAD PATIENT3,BOBBIE UP,AD,AAD	000003 COMMUNITY #1 F 52 138/66 UNC						
PATIENT4, WINONA	000004 COMMUNITY #1 F 53						
PATIENT5, NADINE UP, AD, AAD	000005 COMMUNITY #1 F 61 159/86 UNC						
PATIENT6, RUTH UP	000006 COMMUNITY #1 F 64 139/74 UNC						

Figure 2-9: Sample Patient List, Diabetes: Blood Pressure Control

#### 2.2.5 Diabetes: LDL Assessment

#### **GPRA Measure Description**

During FY 2012, achieve the target rate of 70.3% for the proportion of patients with diagnosed diabetes who are assessed for dyslipidemia (LDL cholesterol).

#### **Denominators**

All *User Population patients* diagnosed with diabetes prior to the Report Period.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. (GPRA Denominator)

Active Adult Diabetic patients, defined by meeting the following criteria: (1) who are age 19 or older at the beginning of the Report Period, (2) whose first ever DM diagnosis occurred prior to the Report Period; (3) who had at least two DM related visits ever, (4) at least one encounter with DM POV in a primary clinic with a primary provider during the Report Period; and (5) never have had a creatinine value greater than five.

#### **Numerators**

Patients with LDL completed during the Report Period, regardless of result. (GPRA Numerator)

Patients with *LDL results* less than (<) 130.

- a. Patients with LDL results less than or equal to (<=) 100
- b. Patients with LDL results 101–129

#### **Logic Description**

*Diabetes definition*: First DM POV 250.00–250.93 recorded in the V POV file prior to the Report Period.

LDL definition: Searches for most recent LDL test with a result during the Report Period. If more than one LDL test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If an LDL test with a result is not found, CRS searches for the most recent LDL test without a result.

CRS uses the following to define the tests:

	CPT Codes	LOINC Codes	Taxonomy
Creatinine (for Active Adult Diabetic denominator)	CPT codes are not included since they do not store the result, which is used in this topic	Yes	DM AUDIT CREATININE TAX
LDL Done	80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F	Yes	DM AUDIT LDL CHOLESTEROL TAX
LDL <130	3048F, 3049F		Tests in above taxonomy with LDL <130
LDL =<100	3048F		Tests in above taxonomy with LDL =<100

In the LOINC Codes column, specific LOINC codes used by CRS are located in the CRS Technical Manual.

#### **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of diabetic patients with documented LDL cholesterol test, if any.

#### **Measure Source**

HP 2010 12-15

#### **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	68.7%
IHS FY 2010 Performance	67.0%
IHS FY 2009 Performance	65.0%
IHS FY 2008 Performance	63.0%
IHS FY 2007 Performance	61.0%
IHS FY 2006 Performance	60.0%
IHS FY 2005 Performance	53.0%

Performance	Percent
IHS FY 2004 Performance	53.0%
IHS FY 2003 Performance	47.5%
IHS FY 2002 Performance	43.7%

DU November 25, 2012 Page 15  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011									
Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Diabetes: LDL Assessment									
					CHG from PREV YR %			CHG from BASE %	
# w/ LDL done	84	36.7	48	23.8	+12.9	23	12.8	+23.9	
# w/LDL <130	58	25.3	40	19.8	+5.5	15	8.3	+17.0	
A. # w/LDL =<100	37	16.2	31			8	4.4	+11.7	
B. # w/LDL 101-129	17	7.4	9	4.5	+3.0	7	3.9	+3.5	
Active Diabetic Pts									
(GPRA)	144		97			87			
# w/ LDL done									
(GPRA)	78	54.2	46	47.4	+6.7	23	26.4	+27.7	
# w/LDL <130		37.5			-1.7	15	17.2	+20.3	
A. # w/LDL =<100	35	24.3	30	30.9				+15.1	
B. # w/LDL 101-129	15	10.4	8	8.2	+2.2	7	8.0	+2.4	
Active Adult Diabetic									
Patients	102		75			63			
# w/ LDL done	60	58.8	43	57.3	+1.5	21	33.3	+25.5	
# w/LDL <130		40.2		46.7	-6.5	13	20.6	+19.6	
A. # w/LDL =<100	26	25.5	26	34.7	-9.2	8	12.7	+12.8	
B. # w/LDL 101-129	B. # w/LDL 101-129 12 11.8 9 12.0 -0.2 5 7.9 +3.8								

Figure 2-10: Sample Report, Diabetes: LDL Assessment

```
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*** IHS 2011 Clinical Performance Measure Patient List ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Entire Patient List

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient
```

```
Diabetes: LDL Assessment: List of diabetic patients with documented LDL cholesterol test, if any

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

PATIENT1, DEBORAH 000001 COMMUNITY #1 F 45
UP, AD, AAD LDL: 03/28/12 119
PATIENT2, TARA 000002 COMMUNITY #1 F 51
UP, AD, AAD LDL: 02/20/12 86
PATIENT3, BOBBIE 000003 COMMUNITY #1 F 52
UP, AD, AAD
PATIENT4, WINONA 000004 COMMUNITY #1 F 53
UP
PATIENT5, NADINE 000005 COMMUNITY #1 F 61
UP, AD, AAD LDL: 02/06/12 CPT 3048F LDL<100
PATIENT6, RUTH 000006 COMMUNITY #1 F 64
UP LDL: 05/21/12 107
```

Figure 2-11: Sample Patient List, Diabetes: LDL Assessment

### 2.2.6 Diabetes: Nephropathy Assessment

### **GPRA Measure Description**

During FY 2012, achieve the target rate of 57.8% for the proportion of patients with diagnosed diabetes who are assessed for nephropathy.

#### **Denominators**

All *User Population patients* diagnosed with diabetes prior to the Report Period.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. (GPRA Denominator)

Active Adult Diabetic patients, defined by meeting the following criteria: (1) who are aged 19 or older at the beginning of the Report Period, (2) whose first ever DM diagnosis occurred prior to the Report Period; (3) who had at least two DM related visits ever, (4) at least one encounter with DM POV in a primary clinic with a primary provider during the Report Period; and (5) never have had a creatinine value greater than five.

### **Numerator**

Patients with nephropathy assessment, defined as an estimated GFR with result and a quantitative urinary protein assessment during the Report Period or with evidence of diagnosis and/or treatment of ESRD at any time before the end of the Report Period. (GPRA Numerator)

# **Logic Description**

*Diabetes definition*: First DM Purpose of Visit 250.00-250.93 recorded in the V POV file prior to the Report period.

Nephropathy assessment definition:

- Estimated GFR with result during the Report Period and Quantitative Urinary Protein Assessment during the Report Period, *or*
- ESRD diagnosis/treatment defined as any diagnosis ever.

CRS uses the following to define the tests/diagnoses:

	CPT Codes	LOINC Codes	Taxonomy
Creatinine (for Active Adult Diabetic Denominator)	CPT codes are not included since they do not store the result, which is used in this topic	Yes	DM AUDIT CREATININE TAX
Estimated GFR		Yes	BGP GPRA ESTIMATED GFR TAX
Quantitative Urinary Protein Assessment	CPT: 82042-82043, 84156	Yes	BGP QUANT URINE PROTEIN TAX Note: Be sure and check with your laboratory supervisor that the names you add to your taxonomy reflect quantitative test values

	CPT Codes	LOINC Codes	Taxonomy
End Stage Renal Disease	V CPT: 36145, 36147, 36800, 36810, 36815, 36818, 36819, 36820, 36821, 36831-36833, 50300, 50320, 50340, 50360, 50365, 50370, 50380, 90951-90970 or old codes 90918-90925, 90935, 90937, 90939 (old code), 90940, 90945, 90997, 90999, 99512, G0257, G0308-G0327 (old codes), G0392 (old code), G0393 (old code), or S9339 V POV: 585.5, 585.6, V42.0, V45.1 (old code), V45.11, V45.12, or V56* V Procedure: 38.95, 39.27, 39.42, 39.43, 39.53, 39.93-39.95, 54.98, or 55.6*		

In the LOINC Codes column, specific LOINC codes used by CRS are located in the CRS Technical Manual.

# **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of diabetic patients with nephropathy assessment, if any.

### **Measure Source**

HP 2010 5-11

# **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	56.5%
IHS FY 2010 Performance	55.0%
IHS FY 2009 Performance	50.0%
IHS FY 2008 Performance	50.0%
IHS FY 2007 Performance (new baseline established; revised standards of care resulted in revised measure definition)	40.0%

Performance	Percent
IHS FY 2006 Performance (measure definition was different from current definition)	55.0%
IHS FY 2005 Performance (measure definition was different from current definition)	47.0%
IHS FY 2004 Performance (measure definition was different from current definition)	42.0%
IHS FY 2003 Performance (measure definition was different from current definition)	37.5%
IHS FY 2002 Performance (measure definition was different from current definition)	35.0%

DU November 25, 2012 Page 17  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Diabetes: Nephropath	y Assess	ment							
					CHG from PREV YR %				
User Pop w/ DM DX pr to Report Period			202			180			
# w/ est GFR & quant UP assmt or w/ESRD	55	24.0	16	7.9	+16.1	8	4.4	+19.6	
Active Diabetic Pts (GPRA)	144		97			87			
# w/ est GFR & quant UP assmt or w/ESRD (GPRA)	53	36.8	11	11.3	+25.5	6	6.9	+29.9	
Active Adult Diabeti Patients	c 102		75			63			
# w/ est GFR & quant UP assmt or w/ESRD	41	40.2	6	8.0	+32.2	4	6.3	+33.8	

Figure 2-12: Sample Report, Diabetes: Nephropathy Assessment

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*** IHS 2011 Clinical Performance Measure Patient List ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012
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	Entire Patient List								
PREG=Pregnant Female; I	UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient								
Diabetes: Nephropathy A assessment, if any.	assessment: List of diabetic patients with nephropathy								
PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR								
PATIENT1, DEBORAH UP, AD, AAD	000001 COMMUNITY #1 F 45								
PATIENT2,TARA UP,AD,AAD	000002 COMMUNITY #1 F 51								
PATIENT3, BOBBIE	000003 COMMUNITY #1 F 52								
UP,AD,AAD PROTEIN	GFR: 09/09/12 & QUANT UP: 03/31/12 QUANT URINE								
PATIENT4, WINONA UP	000004 COMMUNITY #1 F 53								
PATIENT5, NADINE	000005 COMMUNITY #1 F 61								
UP,AD,AAD	ESRD: 03/03/12 CPT 90967								
PATIENT6,RUTH UP	000006 COMMUNITY #1 F 64								
PATIENT7, DANIELLE UP	000007 COMMUNITY #1 F 79 ESRD: 11/01/12 POV V56.8								

Figure 2-13: Sample Patient List, Diabetes: Nephropathy Assessment

# 2.2.7 Diabetic Retinopathy

### **GPRA Measure Description**

During FY 2012, achieve the target rate of 54.8% for the proportion of patients with diagnosed diabetes who receive an annual retinal examination.

#### **Denominators**

All *User Population patients* diagnosed with diabetes prior to the Report Period.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. (GPRA Denominator)

Active Adult Diabetic patients, defined by meeting the following criteria: (1) who are 19 or older at the beginning of the Report Period, (2) whose first ever DM diagnosis occurred prior to the Report Period; (3) who had at least two DM related visits ever, (4) at least one encounter with DM POV in a primary clinic with a primary provider during the Report Period; and (5) never have had a creatinine value greater than five.

#### **Numerators**

Patients receiving a qualified retinal evaluation during the Report Period.

**Note**: This numerator does not include refusals. (GPRA Numerator)

- a. Patients receiving diabetic retinal exam during the Report Period
- b. Patients receiving other eye exams during the Report Period
- c. Patients who refused a diabetic retinal exam during the Report Period.

### **Logic Description**

*Diabetes definition*: First DM Purpose of Visit 250.00–250.93 recorded in the V POV file prior to the Report Period.

Serum creatinine definition (used with Active Adult Diabetic denominator): Site-populated taxonomy DM AUDIT CREATININE TAX; or LOINC taxonomy (NOTE: CPT codes are not included since they do not store the result, which is used in this topic.).

Qualified retinal evaluation definition: (1) Diabetic retinal exam or (2) other eye exam, as shown below. The following methods are qualifying for this measure:

- Dilated retinal evaluation by an optometrist or ophthalmologist
- Seven standard fields stereoscopic photos (ETDRS) evaluated by an optometrist or ophthalmologist
- Any photographic method validated to seven standard fields (ETDRS).

CRS searches in the following order for:

### Diabetic Retinal Exam (any of the following during the report period)

Exam	CPT Codes	Other Codes
Diabetic Retinal Exam	2022F, 2024F, 2026F, S0620, S0621, S3000	VExam: 03 (dilated retinal examination or validated photographic equivalent)

# Other Eye Exam (any of the following during the report period)

Exam	CPT Codes	Other Codes
Non-Did Not Keep Appointment (DNKA) visit to ophthalmology or optometry or validated tele-ophthalmology retinal evaluation clinics		Clinic codes: A2, 17, 18, 64
Non-DNKA visit to an optometrist or ophthalmologist	67028, 67038 (old code), 67039, 67040, 92002, 92004, 92012, 92014	Provider Codes: 24, 79, 08 POV Code: V72.0 Procedure: 95.02
Refusal of a diabetic retinal exam. Refusals are only counted if the patient did not have a diabetic retinal exam or other eye exam. If a patient had both a diabetic retinal exam/other eye exam and a refusal, only the diabetic retinal exam/other eye exam will be counted.		Refusals Exam: 03

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of diabetic patients with qualified retinal evaluation or refusal, if any.

### **Measure Source**

HP 2020 D-10

# **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	54.8% (National rate)
IHS FY 2010 Performance	53.0% (National rate)
IHS FY 2009 Performance	51.0% (National rate)
IHS FY 2008 Performance	50.0% (National rate)
IHS FY 2007 Performance	49.0% (National rate)
IHS FY 2006 Performance	49.0% (National Rate) 52.0% (Designated Sites Rate)
IHS FY 2005 Performance	50.0% (National Rate) 50.0% (Designated Sites Rate)
IHS FY 2004 Performance	47.0% (National Rate) 55.0% (Designated Sites Rate)
IHS FY 2003 Performance	49.0%

Performance	Percent
IHS FY 2002 Performance	49.0%
HP 2020 Goal	58.7%

_	rt Per	l Measu DEMC iod: J	) INDIAN H Tan 01, 20	Commun OSPITA 12 to		012		age 19 **	
Basel	ine Pe	riod:	Jan 01,	2000	to Dec 31,	2000			
Diabetic Retinopathy	Diabetic Retinopathy								
	EPORT				CHG from PREV YR %				
User Pop w/ DM DX prio to report period			202			180			
# w/Retinal Evaluation -No Refusals A. # w/ DM Retinal		28.4	47	23.3	+5.1	54	30.0	-1.6	
exam B. # w/Other	7	3.1	6	3.0	+0.1	6	3.3	-0.3	
Eye Exams	58	25.3	41	20.3	+5.0	48	26.7	-1.3	
# w/Retinal Exam Refusal	3	1.3	0	0.0	+1.3	0	0.0	+1.3	
Active Diabetic Pts (GPRA)	117		95			87			
<pre># w/Retinal Evaluation -No Refusals (GPRA) A. # w/ DM Retinal</pre>		44.6	39	41.1	+3.6	44	50.6	-5.9	
exam B. # w/Other	7	5.8	6	6.3	-0.5	6	6.9	-1.1	
Eye Exams # w/Retinal Exam	47	38.8	33	34.7	+4.1	38	43.7	-4.8	
Refusal	3	2.5	0	0.0	+2.5	0	0.0	+2.5	
Active Diabetic Pts (GPRA)	144		97			87			
# w/Retinal Evaluation -No Refusals (GPRA)		37.5	39	40.2	-2.7	44	50.6	-13.1	
A. # w/ DM Retinal exam	7	4.9	6	6.2	-1.3	6	6.9	-2.0	
B. # w/Other Eye Exams	47	32.6	33	34.0	-1.4	38	43.7	-11.0	
<pre># w/Retinal Exam Refusal</pre>	3	2.1	0	0.0	+2.1	0	0.0	+2.1	
Active Adult Diabetic Patients	102		75			63			

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<pre># w/Retinal Evaluation -No Refusals A. # w/ DM Retinal</pre>	39	38.2	32	42.7	-4.4	39	61.9	-23.7	
exam  B. # w/Other	6	5.9	4	5.3	+0.5	6	9.5	-3.6	
Eye Exams # w/Retinal Exam	33	32.4	28	37.3	-5.0	33	52.4	-20.0	
Refusal	2	2.0	0	0.0	+2.0	0	0.0	+2.0	

Figure 2-14: Sample Report, Diabetic Retinopathy

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D0	November 25, 2012 Page 50				
	*** IHS 2011 Clinical Performance Measure Patient List ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Entire Patient List				
_	re Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic				
<u> </u>	IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;				
CHD-Active Coronary H	Meart Disease; HR=High Risk Patient				
	List of diabetic patients with qualified retinal				
evaluation or refusal	, if any.				
DARLENIE MANG	UDY COMMITTEE CEV ACE				
	HRN COMMUNITY SEX AGE				
DENOMINATOR	NUMERATOR				
PATIENT1.DEBORAH	000001 COMMUNITY #1 F 45				
UP,AD,AAD					
PATIENT2,TARA					
UP,AD,AAD					
PATIENT3,BOBBIE	000003 COMMUNITY #1 F 52				
UP,AD,AAD	06/30/12 Cl 18				
PATIENT4, WINONA	000004 COMMUNITY #1 F 53				
UP					
PATIENT5, NADINE	000005 COMMUNITY #1 F 61				
UP,AD,AAD	05/22/12 Refused				
PATIENT6, RUTH	000006 COMMUNITY #1 F 64				
UP					
PATIENT7, JONELLE	000007 COMMUNITY #1 F 69				
UP,AD,AAD	03/29/12 Diab Eye Ex				

Figure 2-15: Sample Patient List, Diabetic Retinopathy

### 2.2.8 ACEI/ARB Use in Diabetic Patients

### **Denominator**

Active Diabetic patients with HTN, defined as all Active Clinical patients diagnosed with diabetes and hypertension prior to the Report Period, AND at least two visits during the Report Period, AND 2 DM-related visits ever.

#### **Numerators**

Patients not receiving an ACEI or ARB medication during the Report Period.

a. Patients with contraindication/previous adverse reaction to ACEI/ARB therapy.

### **Logic Description**

*Diabetes definition*: First Purpose of Visit 250.00–250.93 recorded in the V POV file prior to the Report Period.

Hypertension definition: Diagnosis (POV or problem list) 401.\* prior to the Report Period, and at least one hypertension POV during the Report Period.

### **ACEI/ARB Numerator Logic**

Ace Inhibitor (ACEI) and ARB (Angiotensin Receptor Blocker) medication codes defined with medication taxonomy BGP PQA ACEI ARB MEDS.

ACEI medications are: Angiotensin Converting Enzyme Inhibitors (Benazepril, Captopril, Enalapril, Fosinopril, Lisinopril, Moexipril, Perindopril, Quinapril, Ramipril, Trandolopril).

Antihypertensive Combinations (Amlodipine-benazepril, Benazepril-hydrochlorothiazide, Captopril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Lisinopril-hydrochlorothiazide, Moexipril-hydrochlorothiazide, Quinapril-hydrochlorothiazide, Trandolapril-verapamil).

ARB medications: Angiotensin II Inhibitors (Candesartan, Eprosartan, Irbesartan, Losartan, Olmesartan, Telmisartan, Valsartan).

Antihypertensive Combinations (Aliskiren-valsartan, Amlodipine-valsartan, Amlodipine-hydrochlorothiazide-valsartan, Amlodipine-olmesartan, Candesartan-hydrochlorothiazide, Eprosartan-hydrochlorothiazide, Irbesartan-hydrochlorothiazide, Losartan-hydrochlorothiazide, Olmesartan-amlodipine-hydrochlorothiazide, Olmesartan-hydrochlorothiazide, Telmisartan-amlodipine, Telmisartan-hydrochlorothiazide, Valsartan-hydrochlorothiazide).

CRS uses the following codes to define contraindications to ACE inhibitors/ARBs.

Contraindication to ACE Inhibitors (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the Report Period with V22.0-V23.9, V72.42, 640.*-649.*, or 651.*-676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.
	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF-MK, or BF-N during the Report Period
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI (not medically indicated) refusal for any ACE inhibitor/ARB at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to ACE inhibitors/ARBs.

	ICD and Other Codes
Adverse Drug Reaction/	<b>POV:</b> 995.0-995.3 AND E942.6
Allergy to ACE Inhibitors/ARBs (any of the codes occurring	Entry in ART (Patient Allergies File): "ace inhibitor", "ACEI", "Angiotensin Receptor Blocker" or "ARB"
ever)	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "ace i*", "ACEI", "Angiotensin Receptor Blocker" or "ARB"

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of diabetic patients with hypertension, with ACEI/ARB medication, contraindication, or ADR, if any.

### **Measure Source**

PQA (Pharmacy Quality Alliance)

### **Measure Past Performance and Long-Term Targets**

None

DU November 25, 2012 Page 24  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000				
ACEI/ARB Use in Diab	etic Patients			
	REPORT % PERIOD	PREV YR % PERIOD	CHG from BASE PREV YR % PERIOD	% CHG from BASE %
Active Diabetic Pts w/ HTN	76	57	54	
# w/ no ACEI/ARB Rx or w/contra/				
ADR	21 27.6	9 15.8	+11.8 6	11.1 +16.5
A. # w/contra/ADR w/ % of Total	13 61.9	2 22.2	+39.7 2	33.3 +28.6

Figure 2-16: Sample Report, ACEI/ARB Use in Diabetic Patients

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient ACEI/ARB Use in Diabetic Patients: List of diabetic patients with hypertension, with ACEI/ARB medication, contraindication, or ADR, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, DEBORAH 000001 COMMUNITY #1 F 20 ACEI/ARB Contra pregnant PATIENT2, TARA 000002 COMMUNITY #1 F 44 06/05/12 PATIENT3, BOBBIE 000003 COMMUNITY #1 F 45 01/29/12 PATIENT4, WINONA 000004 COMMUNITY #1 F 57
AD ACEI/ARB Allergy: 05/0
PATIENT5, NADINE 000005 COMMUNITY #1 F 57
AD ACEI/ARB Allergy: 05/05/12 ADR POV 995.2 + E942.6 PATIENT6, RUTH 000006 COMMUNITY #1 F 61 AD 09/22/12 PATIENT7, JONELLE 000007 COMMUNITY #1 M 25 06/01/12 ACEI/ARB Contra POV 425.1

Figure 2-17: Sample Patient List: ACEI/ARB Use in Diabetic Patients

### 2.2.9 Diabetes: Access to Dental Services

#### **Denominator**

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever.

#### **Numerators**

Patients with documented dental visit during the Report Period.

**Note**: This numerator does *not* include refusals.

Patients with documented dental exam refusal during the Report Period.

#### **Logic Description**

*Diabetes definition:* First DM Purpose of Visit 250.00–250.93 recorded in the V POV file prior to the Report period.

Dental visit definition: For non-CHS dental visits, searches for V Dental ADA codes 0000 or 0190; VExam 30; or POV V72.2. For CHS dental visits, searches for any visit with an ADA code. CHS visit defined as Type code of C in Visit file.

*Refusal definition:* Non-CHS dental visit with refusal of ADA Code 0000 or 0190, or refusal of Exam 30. Refusals are only counted if the patient did not have a documented dental visit.

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of diabetic patients with documented dental visit or refusal, if any.

#### **Measure Source**

HP 2020 D-8

### **Measure Past Performance and Long-Term Targets:**

Past Performance	Percent
IHS FY 2005 Performance	39.0%
IHS FY 2004 Performance	37.0%
IHS FY 2003 Performance	36.0%
IHS FY 2002 Performance	36.0%
HP 2020 Goal	61.2%

### **Performance Improvement Tip**

If your facility's dental services are paid for with CHS funds, ensure the final payment for each purchase order is posted and the CHS to PCC link is set to the "on" position. Having the CHS to PCC link on will enable the CHS data to be passed from CHS/MIS to PCC, where CRS can find it and include it in your CRS reporting.

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					1 to Dec 31 to Dec 31,	-		
Diabetes: Access to	Dental S REPORT PERIOD			%	CHG from PREV YR %			CHG from BASE %
Active Diabetic Pts	144		97			87		
<pre># w/dental visit in   past yr-No Refusals # w/dental exam</pre>	17	11.8	20	20.6	-8.8	18	20.7	-8.9
refusal	4	2.8	0	0.0	+2.8	0	0.0	+2.8

Figure 2-18: Sample Report, Diabetes and Dental Access

Figure 2-19: Sample Patient List, Diabetes and Dental Access

# 2.3 Dental Measure Topics

### 2.3.1 Access to Dental Services

### **GPRA Measure Description**

During FY 2012, achieve the target rate of 26.9% for the proportion of patients who receive dental services.

#### **Denominators**

All patients in the *User Population*. Broken down by age groups (0–5, 6–21, 22–34, 35–44, 45–54, 55–74, >74). (GPRA Denominator)

#### **Numerators**

Patients with documented dental visit during the Report Period.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

Patients with documented dental exam refusal during the Report Period.

### **Logic Description**

Dental visit definition: For non-CHS dental visits, searches for V Dental ADA codes 0000 or 0190; VExam 30; or POV V72.2. For CHS dental visits, searches for any visit with an ADA code. CHS visit defined as Type code of C in Visit file.

*Refusal definition:* Non-CHS dental visit with refusal of ADA Code 0000 or 0190, or refusal of Exam 30. Refusals are only counted if the patient did not have a documented dental visit.

### **Key Logic Changes from CRS Version 11.1**

Updated age breakdowns to 0–5, 6–21, 22–34, 35–44, 45–54, 55–74, >74.

### **Patient List Description**

List of patients with documented dental visit or refusal and date.

### **Measure Source**

HP 2020 OH-7

Measure Past Performance and Long-Term Ta	argets:
---	---------

Performance	Percent
IHS FY 2011 Performance	26.9%
IHS FY 2010 Performance	25.0%
IHS FY 2009 Performance	25.0%
IHS FY 2008 Performance	25.0%
IHS FY 2007 Performance	25.0%
IHS FY 2006 Performance	23.0%
IHS FY 2005 Performance	24.0%
IHS FY 2004 Performance	24.0%
IHS FY 2003 Performance	25.0%
IHS FY 2002 Performance	24.9%
HP 2020 Goal	49.0%

### **Performance Improvement Tip**

If your facility's dental services are paid for with CHS funds, ensure the final payment for each purchase order is posted and the CHS to PCC link is set to the "on" position. Having the CHS to PCC link on will enable the CHS data to be passed from CHS/MIS to PCC, where CRS can find it and include it in your CRS reporting.

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                 Report Period: Jan 01, 2012 to Dec 31, 2012
              Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                 Baseline Period: Jan 01, 2000 to Dec 31, 2000
Access to Dental Services
                                             % CHG from BASE
                                                                     % CHG from
                     REPORT
                               % PREV YR
                     PERIOD
                                  PERIOD
                                                 PREV YR % PERIOD
                                                                        BASE %
# User Pop
(GPRA)
                      2,896
                                     2,456
                                                            2,346
# w/dental visit in
past yr-No Refusals
                        252
                                       201
 (GPRA)
                              8.7
                                             8.2
                                                     +0.5
                                                              207
                                                                    8.8
                                                                            -0.1
# w/dental exam
refusal
                              0.2
                                             0.0
                                                     +0.2
                                                                    0.0
                                                                            +0.2
```

Figure 2-20: Sample Report, Access to Dental Services

```
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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000
```

Access to Dental Services (con't)								
TOTAL USER POPULATION								
	0-5	6 01	_	Distri		EE 71	>74 yrs	
	0-5	0-21	22-34	35-44	45-54	55-74	>/4 yrs	
CURRENT REPORT PERIOD								
Total # User Pop		759	577	407	397	411	81	
<pre># w/dental visit in past Refusals (GPRA)</pre>		71	63	35	32	25	1	
<pre>Refusals (GPRA) % w/dental visit in past</pre>		. –	0.3	33	34	25	1	
Refusals (GPRA)		9.4	10.9	8.6	8.1	6.1	1.2	
<pre># w/dental exam refusal % w/dental exam refusal</pre>			0.0	0.0	0.5	3 0.7	0.0	
w/uencar exam rerusar	0.0	0.1	0.0	0.0	0.5	0.7	0.0	
PREVIOUS YEAR PERIOD								
Total # User Pop		704	517	332	284	292	56	
<pre># w/dental visit in past Refusals (GPRA)</pre>	yr-No 19	59	46	24	24	25	4	
% w/dental visit in past			40	24	24	45	4	
	5.3		8.9	7.2	8.5	8.6	7.1	
	_	_	•	•	_			
<pre># w/dental exam refusal % w/dental exam refusal</pre>					0.0	0.0	0.0	
w/ delical exam refusal	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
CHANGE FROM PREV YR %								
w/dental visit in past y								
Refusals (GPRA) w/dental exam refusal			+2.0 +0.0					
w/delical exam rerusal	+0.0	+U.I	+0.0	+0.0	+0.5	TU./	+0.0	
BASELINE REPORT PERIOD								
Total # User Pop		728	455	293	228	236	52	
<pre># w/dental visit in past Refusals (GPRA)</pre>	yr-No 17	70	39	31	27	20	3	
% w/dental visit in past			39	31	27	∠0	3	
Refusals (GPRA)	_	9.6	8.6	10.6	11.8	8.5	5.8	
<pre># w/dental exam refusal % w/dental exam refusal</pre>				0	0	0	0	
% w/uental exam refusal	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
CHANGE FROM BASE YR %								
w/dental visit in past y								
Refusals (GPRA) w/dental exam refusal		-0.3 +0.1		-2.0 +0.0				
w/dental exam refusal	+0.0	+0.1	+0.0	+0.0	+0.5	+0./	+0.0	

Figure 2-21: Sample Age Breakdown Report, Access to Dental Services

Figure 2-22: Sample Patient List, Access to Dental Services

### 2.3.2 Dental Sealants

### **GPRA Measure Description**

During FY 2012, achieve the target count of 276,893 sealants placed in American Indian and Alaska Native patients.

#### **Denominator**

No denominator. This measure is a total count only, not a percentage.

#### **Numerators**

For patients meeting the *User Population* definition, the total number of dental sealants during the report period.

**Note**: This numerator does not include refusals. (GPRA Numerator).

- a. Dental sealants in patients <12 yrs.
- b. Dental sealants in patients 12–18 yrs.
- c. Dental sealants in patients >18 yrs.

Age breakouts are based on Healthy People 2010 age groups for dental sealants.

### **Logic Description**

Age of the patient is calculated at the beginning of the report period.

Sealants definition: V Dental ADA Code 1351 or V CPT Code D1351. Only two sealants per tooth will be counted during the Report Period. Each tooth is identified by the data element Operative Site in RPMS. If both ADA and CPT codes are found on the same visit, only the ADA will be counted.

Refusal definition: Refusal of ADA Code 1351 or refusal of CPT Code D1351. Refusals are only counted if a patient did not have a sealant during the Report Period. If a patient had both a sealant and a refusal, only the sealant will be counted. If a patient has multiple refusals, only one refusal will be counted.

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients who received or refused dental sealants during Report Period.

#### **Measure Source**

HP 2010 21-8

### **Measure Past Performance and Long-Term Targets:**

Performance	# of Sealants
IHS FY 2011 Performance	276,893
IHS FY 2010 Performance	275,459
IHS FY 2009 Performance	257,067
IHS FY 2008 Performance	241,207
IHS FY 2007 Performance	245,449
IHS FY 2006 Performance	246,645
IHS FY 2005 Performance	249,882
IHS FY 2004 Performance	230,295
	287,158
IHS FY 2003 Performance	232,182
IHS FY 2002 Performance	227,945
IHS FY 2001 Performance	212,612

For the IHS FY 2004 Performance, # of Sealants, please note this was reported by the National Patient Information Reporting System (NPIRS).

### **Performance Improvement Tip**

If your facility's dental visits are paid for with CHS funds, ensure the final payment for each purchase order is posted and the CHS to PCC link is set to the "on" position. Having the CHS to PCC link on will enable the CHS data to be passed from CHS/MIS to PCC, where CRS can find it and include it in your CRS reporting.

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Dental Sealants					
	REPORT PERIOD	PREV YR PERIOD	CHG from PREV YR		CHG from BASE
Total # of Sealants Documented (GPRA)	50	61	-11	81	-31
A. # Dental Sealants documented pts <12 yrs	35	26	+9	40	-5
B. # Dental Sealants documented pts 12-18 yrs	13	34	-21	40	-27
C. # Dental Sealants pts >18 yrs	documented 2	1	+1	1	+1
# refusals	3	0	+3	0	+3

Figure 2-23: Sample Report, Dental Sealants

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Dental Sealants: List of patients who received or refused dental sealants during Report period. PATIENT NAME HRN COMMUNITY
DENOMINATOR NUMERATOR SEX AGE PATIENT20,GEORGE 000020 COMMUNITY #1 M 5 4 sealants: 03/28/12 ADA 1351 (1); 03/28/12 ADA 1351 (1); 03/28/12 ADA 1351 (1); 03/28/12 ADA 1351 (1) PATIENT21, CODY 000021 COMMUNITY #1 M 7
UP 0 sealants: 03/03/12 0 sealants: 03/03/12 Refused ADA 1351 PATIENT50, DAWN 000050 COMMUNITY #2 F 4 3 sealants: 04/15/12 ADA 1351 (1); 05/19/12 ADA 1351 (1); 05/19/12 ADA 1351 (1) PATIENT51,JOY 000051 COMMUNITY #2 F 6

```
UP 2 sealants: 03/17/12 ADA 1351 (2)
PATIENT52,DONALD 000052 COMMUNITY #2 M 8
UP 1 sealants: 02/02/12 CPT D1351 (1)
```

Figure 2-24: Sample Patient List, Dental Sealants

# 2.3.3 Topical Fluoride

### **GPRA Measure Description**

During FY 2012, achieve the target count of 161,461 AI/AN patients who receive at least one topical fluoride application.

#### **Denominator**

No denominator. This measure is a total count only, not a percentage.

#### **Numerators**

For patients meeting the *User Population* definition, the total number of patients with at least one topical fluoride treatment during the Report Period.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

For patients meeting the *User Population* definition, the total number of patients with a documented topical fluoride application refusal in past year.

For patients meeting the *User Population* definition, the total number of appropriate topical fluoride applications based on a maximum of four per patient per year.

For patients meeting the *User Population* definition, the total number of documented topical fluoride application refusals during past year.

### **Logic Description**

Topical fluoride application definition: (1) V Dental ADA Codes 1201 (old code), 1203, 1204, 1205 (old code), 1206, or 5986; (2) V CPT Codes D1201 (old code), D1203, D1204, D1205 (old code), D1206, or D5986; 3) V POV V07.31. A maximum of one application per patient per visit is allowed. A maximum of four topical fluoride applications are allowed per patient per year for the applications measure.

Refusal definition: Refusal of ADA Code 1201 (old code), 1203, 1204, 1205 (old code), 1206 or 5986, or refusal of CPT Code D1201 (old code), D1203, D1204, D1205 (old code), D1206, or D5986. Refusals are only counted if a patient did not have a topical fluoride application during the Report Period. If a patient had both an application and a refusal, only the application will be counted. If a patient has multiple refusals, only one refusal will be counted.

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients who received or refused at least one topical fluoride application during Report Period.

#### **Measure Source**

Not Available

### **Measure Past Performance and Long-Term Targets:**

Performance	Number of Patients
IHS FY 2011 Performance	161,461
IHS FY 2010 Performance	145,181
IHS FY 2009 Performance	136,794
IHS FY 2008 Performance	120,754
IHS FY 2007 Performance	107,934
IHS FY 2006 Performance	95,439
IHS FY 2005 Performance	85,318
IHS FY 2005 Performance	113,324

For the IHS FY 2005 Performance, Number of Patients (113,324) is the number of applications.

### **Performance Improvement Tip**

If your facility's dental visits are paid for with CHS funds, ensure the final payment for each purchase order is posted and the CHS to PCC link is set to the "on" position. Having the CHS to PCC link on will enable the CHS data to be passed from CHS/MIS to PCC, where CRS can find it and include it in your CRS reporting.

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Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011
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	Baseline Period:	Jan 01, 20	00 to Dec 31,	2000	
Topical Fluoride	:				
	REPORT PERIOD	PREV YR PERIOD	CHG from PREV YR		CHG from BASE
Total # of patie Least 1 Topical -No Refusals (G	Fluoride App	26	+19	15	+30
<pre># Patients w/Refusals</pre>	7	0	+7	0	+7
Total # of Topic Applications	al Fluoride 50	26	+24	15	+35
# Refusals	7	0	+7	0	+7

Figure 2-25: Sample Report, Topical Fluoride

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Topical Fluoride: List of patients who received or refused at least one topical fluoride application during Report period. HRN COMMUNITY PATIENT NAME SEX AGE DENOMINATOR NUMERATOR PATIENT20, GEORGE 000020 COMMUNITY #1 M 5 1 topical fluoride: 06/18/12 CPT D5986 PATIENT21,RYAN 000021 COMMUNITY #1 000021 PATIENT22, MICHAEL 000022 COMMUNITY #1 M 9 0 topical fluoride: 03/03/12 Refused ADA 1201 0 topical fluoride: 03/03/12 Refused CPT D1203 PATIENT23, MARTY 000023 COMMUNITY #1 M 15 2 topical fluoride: 01/07/12 ADA 1204; 08/27/12 ADA 1204

Figure 2-26: Sample Patient List, Topical Fluoride

# 2.4 Immunization Measure Topics

### 2.4.1 Influenza

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 63.4% for the proportion of non-institutionalized adults aged 65 years and older who receive an influenza immunization.

#### **Denominators**

All Active Clinical patients. Broken down by age groups.

- a. Active Clinical patients younger than age 18.
- b. Active Clinical patients ages 18-49.
- c. Active Clinical patients ages 18-49 and considered high risk for influenza.
- d. Active Clinical patients ages 50–64.
- e. Active Clinical patients ages 65 and older. (GPRA Denominator)

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever.

All User Population patients. Broken down by age groups.

- a. User Population patients younger than age 18.
- b. User Population patients ages 18-49.
- c. User Population patients ages 18-49 and considered high risk for influenza.
- d. User Population patients ages 50-64.
- e. User Population patients ages 65 and older.

#### **Numerators**

Patients with influenza vaccine documented during the report period or with a contraindication documented at any time before the end of the report period.

**Note**: The only refusals included in this numerator are not medically indicated (NMI) refusals. (GPRA Numerator)

a. Patients with a contraindication or a documented NMI refusal.

Patients with documented influenza refusal during the report period.

### **Logic Description**

Age of the patient is calculated at the beginning of the report period.

*Diabetes:* First DM POV 250.00–250.93 recorded in the V POV file prior to the Report period.

*Influenza definition:* Any of the following documented during the Report Period unless otherwise noted.

1. **Influenza immunization:** Any of the codes in the table below.

	CPT Codes	ICD and Other Codes
Influenza vaccine	90654-90662, 90724 (old code), G0008, G8108 (old code)	Immunization (CVX) Codes: 88-Influenza Virus Vaccine, NOS; 15 Inf Virus Vac SV; 16 Inf Virus Vac WV; 111 Inf Virus Vac Intranasal; 135 Inf High Dose Seasonal; 140 Inf Virus Vac SV Preservative Free; 141 Inf Virus Vac SV; 144 Inf Virus Vac SV Intradermal POV: V04.8 (old code), V04.81 NOT documented with 90663, 90664, 90666- 90668, 90470, G9141 or G9142, V06.6 NOT documented with 90663, 90664, 90666-90668, 90470, G9141 or G9142 ICD Procedure Code: 99.52

- 2. **Contraindication:** Any of the following documented at any time before the end of the Report Period, defined as: (A) Contraindication in the Immunization Package of "Egg Allergy" or "Anaphylaxis" or (B) PCC NMI Refusal.
- 3. **Refusal:** Any of the following during the Report Period: (A) Immunization codes 88, 111, 135, 140, 141, 144, 15, or 16, as documented in PCC Refusal File (i.e., REF), (B) CPT codes 90654-90662, 90724 (old code), G0008, G8108 (old code) as documented in PCC Refusal File (i.e., REF) or (C) Immunization Package as contraindication of "Patient Refusal."
- 4. **High Risk for Influenza:** Persons considered high risk for influenza are defined as those who have 2 or more visits in the past 3 years with a POV or Problem diagnosis of any of the following: HIV Infection (042, 042.0-044.9 (old codes)); Diabetes (250.00-250.93); Rheumatic Heart Disease (393.-398.99); Hypertensive Heart Disease (402.00-402.91); Hypertensive Heart/Renal Disease (404.00-404.93); Ischemic Heart Disease (410.00-414.9); Pulmonary Heart Disease (415.0-416.9); Other Endocardial Heart Disease (424.0-424.9); Cardiomyopathy (425.0-425.9); Congestive Heart Failure (428.0-428.9, 429.2); Chronic Bronchitis (491.0-491.9); Emphysema (492.0-492.8); Asthma (493.00-493.91); Bronchiectasis, CLD, COPD (494.0-496.); Pneumoconioses (500-505); Chronic Liver Disease (571.0-571.9); Nephrotic Syndrome (581.0-581.9); Renal Failure (585.6, 585.9); Transplant (996.80-996.89); Kidney Transplant (V42.0-V42.89); Chemotherapy (V58.1); Chemotherapy follow-up (V67.2).

### **Key Logic Changes from CRS Version 11.1**

- Added CVX code 144 to Influenza and refusal definitions.
- Added CPT codes 90654-90662, 90724 (old code), G0008, G8108 (old code) to Influenza refusal definition.

### **Patient List Description**

List of patients with Influenza code or refusal, if any.

#### **Measure Source**

HP 2020 IID-12.7

# Measure Past Performance and Long-Term Targets for Patients => 65 Vaccine Rate:

Performance	Percent
IHS FY 2011 Performance	63.4%
IHS FY 2010 Performance	62.0%
IHS FY 2009 Performance	59.0%
IHS FY 2008 Performance	62.0%
IHS FY 2007 Performance	59.0%
IHS FY 2006 Performance	58.0%
IHS FY 2005 Performance	59.0%
IHS FY 2004 Performance	54.0%
IHS FY 2003 Performance	51.0%
IHS FY 2002 Performance	51.4%
HP 2020 Goal	90.0%

# **Performance Improvement Tips**

- Providers should ask about and record off-site historical immunizations (IZ type, date received and location) on PCC forms. Data entry mnemonic: HIM
- Providers should document refusals; write "Refused" in Influenza Order box on PCC form. Data entry mnemonic: REF (Immunization, Value, Date Refused).

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Influenza REPORT PERIOD	% PR PE				BASE PERIOD		CHG from BASE %
Active Clinical Pts 1,554		1,209			1,101		
Total # w/Flu vaccine/ contra/NMI Refusal 194 A. # w/ Contraind/ NMI Ref	12.5	115	9.5	+3.0	49	4.5	+8.0
w/ % of Total IZ 12 # w/ Influenza	6.2	2	1.7	+4.4	0	0.0	+6.2

Refusal	17	1.1	6	0.5	+0.6	0	0.0	+1.1	
Active Clinical Pts <18 4	142		416			435			
		12.2	21	5.0	+7.2	6	1.4	+10.8	
A. # w/ Contraind/ NMI Re w/ % of Total IZ # w/ Influenza	ef 3	5.6	0	0.0	+5.6	0	0.0	+5.6	
Refusal	6	1.4	0	0.0	+1.4	0	0.0	+1.4	
Active Clinical Pts 18-49 7	752		572			486			
	48	6.4	33	5.8	+0.6	14	2.9	+3.5	
A. # w/ Contraind/ NMI Re	3	6.3	2	6.1	+0.2	0	0.0	+6.3	
# w/ Influenza Refusal	3	0.4	0	0.0	+0.4	0	0.0	+0.4	
Active Clinical Pts 18-49 high risk 1	.66		109			69			
Total # w/Flu vaccine/ contra/NMI Refusal A. # w/ Contraind/ NMI Re	25	15.1	25	22.9	-7.9	7	10.1	+4.9	
w/ % of Total IZ # w/ Influenza	3	12.0	1	4.0	+8.0	0	0.0	+12.0	
Refusal	3	1.8	0	0.0	+1.8	0	0.0	+1.8	
Active Clinical Patients ages 50-64 2	244		157			115			
Total # w/Flu vaccine/contra/ NMI Refusal A. # w/ Contraind/ NMI	59	24.2	37	23.6	+0.6	14	12.2	+12.0	
Ref w/ % of Total IZ	5	8.5	0	0.0	+8.5	0	0.0	+8.5	
# w/Influenza Refusal	3	1.2	5	3.2	-2.0	0	0.0	+1.2	

Figure 2-27: Sample Report, Adult Immunizations: Influenza

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```
UP,AC
PATIENT3,DEMETRIA 000003 COMMUNITY #1 F 35
UP,AC, HR 02/25/12 Imm 140
PATIENT4,JADE 000004 COMMUNITY #1 F 50
UP
PATIENT5,MARIE 000005 COMMUNITY #1 F 65
UP,AC,AD,HR 01/21/12 NMI Refusal
```

Figure 2-28: Sample Patient List, Adult Immunization: Influenza

### 2.4.2 Adult Immunizations

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 87.5% for the proportion of adult patients age 65 years and older who receive a pneumococcal immunization.

#### **Denominators**

Active Clinical patients ages 65 or older. (GPRA Denominator)

Active Clinical patients ages 18-64 and considered high risk for pneumococcal.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever.

Active Clinical patients ages 18-64.

All *User Population patients* ages 65 and older at beginning of Report Period.

*User Population patients* ages 18-64 and considered high risk for pneumococcal.

*User Population patients* ages 18-64.

#### **Numerators**

Patients with Pneumococcal vaccine or contraindication documented at any time before the end of the Report Period.

**Note**: The only refusals included in this numerator are NMI refusals. (GPRA Numerator)

a. Patients with a contraindication or a documented NMI refusal.

Patients with documented Pneumococcal refusal during the Report Period.

Patients with Pneumococcal vaccine or contraindication documented ever and, if patient is older than 65 years, either a dose of pneumovax after the age of 65 or a dose of pneumovax in the past 5 years.

**Note**: The only refusals included in this numerator are NMI refusals. (GPRA Numerator)

Patients who have received 1 dose of Tdap ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals. (GPRA Numerator)

Patients who have received 1 dose of Tdap/Td in the past 10 years, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals. (GPRA Numerator)

### **Logic Description**

Age of the patient is calculated at the beginning of the Report period.

*Diabetes definition:* First DM POV 250.00250.93 recorded in the V POV file prior to the Report period.

*Pneumococcal Immunization definition:* Any of the following documented any time before the end of the Report Period unless otherwise noted.

1. **Pneumoccocal immunization:** Any of the codes in the table below.

Immunization	CPT Codes	ICD and Other Codes
Pneumoccocal	90669, 90670, 90732,	Immunization (CVX) Codes:
Vaccine	G0009, G8115 (old	33 - Pneumococcal Polysaccaride Vaccine;
	code)	100 - Pneumococcal Conjugate Vaccine;
	,	109 – Pneumo NOS; 133 – Pneumo
		Conjugate
		<b>POV:</b> V06.6; V03.82
		V Procedure: 99.55

2. **Pneumoccocal Contraindication:** (A) Contraindication in the Immunization Package of "Anaphylaxis" or (B) PCC NMI Refusal.

- 3. **Pneumoccocal Refusal:** Any of the following during the Report Period: (A) Immunization codes 33, 100, 109, or 133, as documented in PCC Refusal File (i.e., REF), (B) CPT codes 90669, 90670, 90732, G0009, G8115 (old code) as documented in PCC Refusal File (i.e. REF) or (C) Immunization Package as contraindication of "Patient Refusal."
- 4. **High Risk for Pneumococcal:** Persons considered high risk for pneumococcal are defined as those who have 2 or more visits in the past 3 years with a POV or Problem diagnosis of any of the following: HIV Infection (042, 042.0-043.9 (old codes), 044.9 (old code)); Diabetes (250.00-250.93); Chronic alcoholism (303.90, 303.91); Congestive Heart Failure (428.0-428.9, 429.2); Emphysema (492.0-492.8); Asthma (493.00-493.91); Bronchiectasis, CLD, COPD (494.-496.); Pneumoconioses (501.-505.); Chronic Liver Disease (571.0-571.9); Nephrotic Syndrome (581.0-581.9); Renal Failure (585.6, 585.9); Injury to spleen (865.00-865.19); Transplant (996.80-996.89); Kidney Transplant (V42.0-V42.89); Chemotherapy (V58.1); Chemotherapy follow-up (V67.2).

*Tdap/Td Immunization definition:* Any of the following documented during the applicable time frame.

1. **Tdap/Td immunization:** Any of the codes in the table below.

Immunization	CPT Codes	ICD and Other Codes
Tdap Vaccine	90715	Immunization (CVX) Codes:  115 – Tetanus toxoid, reduced diphtheria toxoid, and acellular pertussis vaccine, adsorbed
Td Vaccine	90714, 90718	Immunization (CVX) Codes:  9 – Tetanus and diphtheria toxoids, adsorbed, for adult use  113 – Tetanus and diphtheria toxoids, adsorbed, preservative free, for adult use  POV: V06.5

2. **Tdap/Td Contraindication:** (A) Contraindication in the Immunization Package of "Anaphylaxis" or (B) PCC NMI Refusal.

### **Key Logic Changes from CRS Version 11.1**

Added CPT codes 90669, 90670, 90732, G0009, G8115 (old code) to Pneumococcal refusal definition.

#### **Patient List Description**

List of patients =>18 yrs or DM DX with IZ, evidence of disease, contraindication, or refusal, if any.

### **Measure Source**

HP 2020 IID-13.1

### **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	85.5%
IHS FY 2010 Performance	84.0%
IHS FY 2009 Performance	82.0%
IHS FY 2008 Performance	82.0%
IHS FY 2007 Performance	79.0%
IHS FY 2006 Performance	74.0%
IHS FY 2005 Performance	69.0%
IHS FY 2004 Performance	69.0%
IHS FY 2003 Performance	65.0%
IHS FY 2002 Performance	64.0%
HP 2020 Goal for % of patients => 65	90.0%

### **Performance Improvement Tips**

- Providers should ask about and record off-site historical immunizations (IZ type, date received and location) on PCC forms. Data entry mnemonic: HIM
- Providers should document refusals; write "Refused" in Pneumo Vax Order box on PCC form. Data entry mnemonic: REF (Immunization, Value, Date Refused).

DU November 25, 2012 Page 41  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Adult Immunizations									
	REPORT PERIOD		PREV YR PERIOD		CHG from PREV YR %				
Active Clinical Pts ages 65 & older (GPRA)	116		64			65			
Total # w/Pneumovax/ contra/NMI Refusal (GPRA) A. # w/ Contraind/ N Ref w/ % of		43.1	44	68.8	-25.6	37	56.9	-13.8	
Total IZ	4	8.0	2	4.5	+3.5	0	0.0	+8.0	
# w/Pneumovax Refusal	3	2.6	0	0.0	+2.6	0	0.0	+2.6	
Active Clinical Pts									

18-64 high risk	222		164			105			
Total # w/Pneumovax/ contra/									
NMI Refusal	55	24.8	47	28.7	-3.9	39	37.1	-12.4	
A. # w/ Contra/ NMI Ref									
w/ % of Total IZ	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
# w/ Pneumovax									
Refusal	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Active Diabetic Pts	144		97			87			
Total # w/up to date Pneumovax/contra/NMI Refusal A. # w/ Contraind/ NMI	56	38.9	49	50.5	-11.6	46	52.9	-14.0	
Ref w/ % of Total IZ # w/Pneumovax	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Refusal	0	0.0	0	0.0	+0.0	0	0.0	+0.0	

Figure 2-29: Sample Report, Adult Immunization: Pneumovax

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Adult Immunizations: List of patients =>18 yrs or DM DX with IZ, evidence of disease, contraindication, or refusal, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, DEBORAH 000001 COMMUNITY #1 F 18

UP, AC, HR TDAP/TD: 12/22/11 Imm 9 (past 10 yrs)

PATIENT2, TARA 000002 COMMUNITY #1 F 27

IDAC TDAP: 03/03/12 Imm 115 (ever): TDAP/T TDAP: 03/03/12 Imm 115 (ever); TDAP/TD: 03/03/12 UP,AC TDAP: 03/03/12 Imm 13
PATIENT3, BOBBIE 000003 COMMUNITY #1 F 41 UP,AC PATIENT4, NADINE 000004 COMMUNITY #1 F 55 UP, AC, HR, AD Pneumo: 03/27/10 Imm Pneumo: 03/27/10 Imm 33 (ever) (up-to-date); TDAP/TD: 03/27/10 Imm 9 (past 10 yrs) PATIENT5, SHERRY 000005 COMMUNITY #1 F 68 UP,AC,HR Pneumo: 02/03/12 Refused CPT 90669

Figure 2-30: Sample Patient List, Adult Immunization: Pneumovax

### 2.4.3 Childhood Immunizations

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 77.8% for the proportion of American Indian/Alaska Native children ages 19-35 months who have received the recommended immunizations.

#### **Denominators**

Active Clinical patients ages 19-35 months at end of Report Period.

User Population patients ages 19–35 months at end of Report Period.

User Population patients *active in the Immunization Package* who are 19–35 months at end of Report Period. (GPRA Denominator)

**Note:** Only values for the Current Period will be reported for this denominator since currently there is not a way to determine if a patient was active in the Immunization Package during the Previous Year or Baseline Periods.

#### **Numerators**

Patients who have received the 4:3:1:3:3 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B), including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received the 4:3:1:3:3:1 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B, 1 Varicella), including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received the 4:3:1:3:3:1:4 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B, 1 Varicella, and 4 Pneumococcal), including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals. (GPRA Numerator)

Patients who have received four doses of DTaP ever, including contraindications.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received three doses of Polio ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received one dose of MMR ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received three doses of HiB ever, including contraindications.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received three doses of Hepatitis B vaccine ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received one dose of Varicella ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received four doses of Pneumococcal conjugate vaccine ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received two doses of Hepatitis A vaccine ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received two or three doses of Rotavirus vaccine ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received two doses of Influenza vaccine ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

For each of the above numerators, the following subnumerators are included:

a. Patients with either (1) evidence of the disease, (2) a contraindication, or (3) a documented NMI refusal

Patients with documented REF refusal in PCC or Parent or Patient refusal in the IZ program

**Immunization Program Numerator**: Patients who have received the 4:3:1:3:3 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B), *not* including refusals, contraindications, and patients with evidence of disease.

**Immunization Program Numerator**: Patients who have received the 4:3:1:3:3:1 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B, and 1 Varicella), *not* including refusals, contraindications, and patients with evidence of disease.

**Immunization Program Numerator**: Patients who have received the 4:3:1:3:3:1:4 combination (i.e., 4 DTaP, 3 Polio, 1 MMR, 3 HiB, 3 Hepatitis B, 1 Varicella, and 4 Pneumococcal), *not* including refusals, contraindications, and patients with evidence of disease.

### **Logic Description**

Age definition: Age of the patient is calculated at the beginning of the Report Period. Therefore the age range will be adjusted to 7–23 months, which makes the patient between the ages of 19–35 months at the end of the Report Period. Because IZ data comes from multiple sources, any IZ codes documented on dates within 10 days of each other will be considered as the same immunization.

Active Immunization Package Patients denominator definition: Same as User Pop definition except includes only patients flagged as active in the Immunization Package.

**Note**: Only values for the Current Period will be reported for this denominator since currently there is not a way to determine if a patient was active in the Immunization Package during the Previous Year or Baseline Periods.

Dosage and types of immunization definitions:

- Four doses of DTaP: (1) four DTaP/DTP/Tdap; (2) one DTaP/DTP/Tdap and three DT/td; (3) one DTaP/DTP/Tdap and three each of Diphtheria and Tetanus; (4) four DT and four Acellular Pertussis; (5) four Td and four Acellular Pertussis; or (6) four each of Diphtheria, Tetanus, and Acellular Pertussis.
- Three doses of Polio: (1) v OPV; (2) three IPV; or (3) combination of OPV and IPV totaling 3 doses.
- One dose of MMR: (1) MMR; (2) one M/R and one Mumps; (3) one R/M and one Measles; or (4) one each of Measles, Mumps, and Rubella.
- Three doses of Hep B
- Three doses of HIB
- One dose of Varicella
- Four doses of Pneumococcal
- Two doses of Hepatitis A
- Two or three doses of Rotavirus, depending on the vaccine administered
- Two doses of Influenza

Except for the Immunization Program Numerators, NMI refusals, evidence of disease, and contraindications for individual immunizations will also count toward meeting the definition, as defined below. Refusals will count toward meeting the definition for refusal numerators only.

**Note**: NMI refusals are not counted as refusals; rather, they are counted as contraindications.

- For immunizations that allow a different number of doses (e.g. 2 or 3 Rotavirus): To count toward the numerator with the smaller number of doses, all of the patient's vaccinations must be part of the smaller dose series. For example, for a patient to count toward the Rotavirus numerator with only 2 doses, all two doses must be included in the 2-dose series codes listed in the Rotavirus definition. A patient with a mix of 2-dose and 3-dose series codes will need 3 doses to count toward the numerator.
- Each immunization must be refused and documented separately. For example, if a
  patient has an NMI refusal for Rubella only, then there must be an immunization,
  contraindication, or separate NMI refusal for the Measles and Mumps
  immunizations.
- For immunizations where required number of doses is >1, only one NMI refusal is necessary to be counted in the numerator. For example, if there is a single NMI refusal for Hepatitis B, the patient will be included in the numerator.

- For immunizations where required number of doses is >1, only one contraindication is necessary to be counted in the numerator. For example, if there is a single contraindication for HiB, the patient will be included in the numerator.
- Evidence of disease will be checked for at any time in the child's life (prior to the end of the Report Period.)
- To be counted in subnumerator A, a patient must meet the numerator definition and have a REF refusal in PCC or a Parent or Patient Refusal in the IZ program for any of the immunizations in the numerator. For example, if a patient refused Rubella only but had immunizations for Measles and Mumps, the patient would be included in subnumerator A.

For the separate numerator for REF refusal (Patient Refusal for Service) in PCC or a Parent or Patient refusal in the IZ program, all conditions shown below must be met:

- Each immunization must be refused and documented separately. For example, if a patient has an REF refusal for Rubella, then there also must be an immunization, contraindication, or separate REF refusal for Measles and Mumps.
- Where the required number of doses is >1, only one REF refusal in PCC or one Parent or Patient refusal in the IZ program is necessary to be counted in the numerator. For example, for the 4 DTaP numerator, only one refusal is necessary to be counted in the refusal numerator.

Childhood immunization refusals are defined as Parent/Patient Refusal in Immunization package or PCC Refusal type REF or NMI for IZ codes:

Immunization	Immunization Codes for Refusals	CPT Codes for Refusals
DTaP	20, 50, 106, 107, 110, 120, 130, 132, 146	90696, 90698, 90700, 90721, 90723
DTP	1, 22, 102	90701, 90711 (old code), 90720
Tdap	115	90715
DT (Diphtheria & Tetanus)	28	90702
Td (Tetanus & Diphtheria)	9, 113	90714, 90718
Tetanus	35, 112	90703
Acellular Pertussis	11	
OPV	2, 89	90712
IPV	10, 89, 110, 120, 130, 132, 146	90696, 90698, 90711 (old code), 90713, 90723
MMR	3, 94	90707, 90710; M/R: 90708
M/R (Measles/ Rubella)	4	
R/M (Rubella/ Mumps)	38	90709 (old code)
Measles	5	90705
Mumps	7	90704

Immunization	Immunization Codes for Refusals	CPT Codes for Refusals
Rubella	6	90706
HiB	17, 22, 46-49, 50, 51, 102, 120, 132, 146	90645-90648, 90698, 90720- 90721, 90737 (old code), 90748
Hepatitis B	8, 42-45, 51, 102, 104, 110, 132, 146	90636, 90723, 90731 (old code), 90740, 90743-90748, G0010, Q3021 (old code), Q3023 (old code)
Varicella	21, 94	90710, 90716
Pneumococcal	33, 100, 109, 133	90669, 90670, 90732, G0009, G8115 (old code)
Hepatitis A	1, 52, 83, 84, 85, 104	90632-90634, 90636, 90730 (old code)
Rotavirus	74, 116, 119, 122	90680
Influenza	15, 16, 88, 111, 135, 140, 141	90654-90658, 90659 (old code), 90660-90662, 90724 (old code), G0008, G8108 (old code)

Childhood immunizations are defined in the following ways:

Immunization	CPT Codes	ICD and Other Codes NOTE: IZ Program Numerators Do Not Use POV, V Procedure, Evidence of Disease, Contraindication or Refusal Codes
DTaP	90696, 90698, 90700, 90721, 90723	Immunization (CVX) Codes: 20, 50, 106, 107, 110, 120, 130, 132, 146 POV: V06.1
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
DTP	90701, 90711 (old code), 90720	Immunization (CVX) Codes: 1, 22, 102 POV: V06.1, V06.2, V06.3 V Procedure: 99.39 Contraindications: Immunization Package contraindication of "Anaphylaxis."
Tdap	90715	Immunization (CVX) Codes: 115 Contraindications: Immunization Package contraindication of "Anaphylaxis."
DT (Diphtheria & Tetanus)	90702	Immunization (CVX) Codes: 28 POV: V06.5 Contraindications: Immunization Package contraindication of "Anaphylaxis."
Td (Tetanus & Diphtheria)	90714, 90718	Immunization (CVX) Codes: 9, 113 POV: V06.5 Contraindications: Immunization Package contraindication of "Anaphylaxis."

Immunization	CPT Codes	ICD and Other Codes NOTE: IZ Program Numerators Do Not Use POV, V Procedure, Evidence of Disease, Contraindication or Refusal Codes
Diphtheria	90719	<b>POV:</b> V03.5
		V Procedure: 99.36
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Tetanus	90703	Immunization (CVX) Codes: 35, 112 POV: V03.7
		V Procedure: 99.38
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Acellular Pertussis		Immunization (CVX) Codes: 11 POV: V03.6
		V Procedure: 99.37 (old code)
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
OPV	90712	Immunization (CVX) Codes: 2, 89
		Contraindications: POV 279, V08, 042, 200-202, 203.0, 203.1, 203.8, 204-208 or Immunization Package contraindication of "Anaphylaxis."
IPV	90696, 90698, 90711 (old	Immunization (CVX) Codes: 10, 89, 110, 120, 130, 132, 146 POV: V04.0, V06.3
	code), 90713,	V Procedure: 99.41
	90723	<b>Evidence of Disease:</b> POV or PCC Problem List (active or inactive) 730.70-730.79
		Contraindications: Immunization Package contraindication of "Anaphylaxis" or "Neomycin Allergy."
MMR	90707, 90710	Immunization (CVX) Codes: 3, 94
		<b>POV:</b> V06.4
		V Procedure: 99.48
		Contraindications: POV 279, V08, 042, 200-202, 203.0, 203.1, 203.8, 204-208; or Immunization Package contraindication of "Anaphylaxis," "Immune Deficiency," "Immune Deficient," or "Neomycin Allergy."
M/R (Measles/	90708	Immunization (CVX) Codes: 4
Rubella)		Contraindications: Immunization Package contraindication of "Anaphylaxis"
R/M (Rubella/	90709 (old code)	Immunization (CVX) Codes: 38
Mumps)		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis"

Immunization	CPT Codes	ICD and Other Codes NOTE: IZ Program Numerators Do Not Use POV, V Procedure, Evidence of Disease, Contraindication or Refusal Codes
Measles	90705	Immunization (CVX) Codes: 5 POV: V04.2 V Procedure: 99.45 Evidence of Disease: POV or PCC Problem List (active or inactive) 055* Contraindications: Immunization Package contraindication of
Mumps	90704	"Anaphylaxis"  Immunization (CVX) Codes: 7 POV: V04.6 V Procedure: 99.46 Evidence of Disease: POV or PCC Problem List (active or inactive) 072* Contraindications: Immunization Package contraindication of "Anaphylaxis"
Rubella	90706	Immunization (CVX) Codes: 6 POV: V04.3 V Procedure: 99.47 Evidence of Disease: POV or PCC Problem List (active or inactive) 056*, 771.0 Contraindications: Immunization Package contraindication of "Anaphylaxis"
HiB	90645-90648, 90698, 90720- 90721, 90737 (old code), 90748	Immunization (CVX) Codes: 17, 22, 46-49, 50, 51, 102, 120, 132, 146 POV: V03.81 Contraindications: Immunization Package contraindication of "Anaphylaxis"
Hepatitis B	90636, 90723, 90731 (old code), 90740, 90743-90748, G0010, Q3021, (old code) Q3023 (old code)	Immunization (CVX) Codes: 8, 42-45, 51, 102, 104, 110, 132, 146 Evidence of Disease: POV or PCC Problem List (active or inactive) V02.61, 070.2, 070.3 Contraindications: Immunization Package contraindication of "Anaphylaxis"
Varicella	90710, 90716	Immunization (CVX) Codes: 21, 94 POV: V05.4 Evidence of Disease: 1) POV or PCC Problem List (active or inactive) 052*, 053* or 2) 2) Immunization Package contraindication of "Hx of Chicken Pox" or "Immune." Contraindications: POV 279, V08, 042, 200-202, 203.0, 203.1, 203.8, 204-208; or Immunization Package contraindication of "Anaphylaxis," "Immune Deficiency," "Immune Deficient," or "Neomycin Allergy."

Immunization	CPT Codes	ICD and Other Codes NOTE: IZ Program Numerators Do Not Use POV, V Procedure, Evidence of Disease, Contraindication or Refusal Codes
Pneumococcal	90669, 90670, 90732, G0009, G8115 (old code)	Immunization (CVX) Codes: 33, 100, 109, 133 POV: V06.6; V03.82 Contraindications: Immunization Package contraindication of "Anaphylaxis"
Hepatitis A	90632-90634, 90636, 90730 (old code)	Immunization (CVX) Codes: 31, 52, 83, 84, 85, 104  Evidence of Disease: POV or PCC Problem List (active or inactive) 070.0, 070.1  Contraindications: Immunization Package contraindication of
Rotavirus – 2- dose series	90681	"Anaphylaxis"  Immunization (CVX) Codes: 119 Contraindications: Immunization Package contraindication of "Anaphylaxis" or "Immune Deficiency"
Rotavirus – 3- dose series	90680	Immunization (CVX) Codes: 74, 116, 122 POV: V05.8 Contraindications: Immunization Package contraindication of "Anaphylaxis" or "Immune Deficiency"
Influenza	90654-90658, 90659 (old code), 90660- 90662, 90724 (old code), G0008, G8108 (old code)	Immunization (CVX) Codes: 15, 16, 88, 111, 135, 140, 141, 144 POV: V04.8 (old code), V04.81, V06.6 V Procedure: 99.52 Contraindications: Immunization Package contraindication of "Egg Allergy" or "Anaphylaxis"

### **Key Logic Changes from CRS Version 11.1**

- Added CVX code 144 to Influenza and refusal definitions.
- Added CVX code 146 to DTaP, IPV, HiB, and Hepatitis B definitions.
- Added CPT codes to refusal definitions.
- Removed "or 2 doses if documented with CPT 90743" from Hepatitis B definition.

### **Patient List Description**

List of patients 19–35 months with IZ, if any. If a patient did not have all doses in a multiple dose vaccine, the IZ will not be listed. For example, if a patient only had two DTaP, no IZ will be listed for DTaP.

**Note 1:** Because age is calculated at the beginning of the Report Period, the patient's age on the list will be between 7-23 months.

Note 2: The order of the display for the immunizations is: 4
Dtap/Dtp;3 IPV/OPV;MMR;3 HIB;3 HEP;Vari;4
PNEUMO. A blank value in the Numerator column means the patient didn't meet the requirements for any of the immunizations. Another example is "MMR; vari;4
PNEUMO," which means the patient did not have 4
Dtap/Dtp, 3 IPV/OPV, 3 HIB and 3 Hep B immunizations.

### **Measure Source**

CDC; HP 2020 IID-7, IID-8; HEDIS

# **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 GPRA Performance Active Immunization Package 4:3:1:3:3:1:4 (rate for children age 19–35 months)	75.9%
IHS FY 2010 GPRA Performance Active Immunization Package 4:3:1:3:3:1 (rate for children age 19–35 months)	79.0%
IHS FY 2009 GPRA Performance Active Immunization Package 4:3:1:3:3 (rate for children age 19–35 months)	79.0%
IHS FY 2008 GPRA Performance Active Immunization Package 4:3:1:3:3 (rate for children age 19–35 months)	78.0%
IHS FY 2008 Non-GPRA Performance Active Clinical 4:3:1:3:3 (rate for children age 19–35 months)	68.0%
IHS FY 2007 GPRA Performance Active Immunization Package 4:3:1:3:3(rate for children age 19–35 months)	78.0%
IHS FY 2006 Performance (rate for children age 19–35 months)	80.0%
IHS FY 2005 Performance (rate for children age 19–35 months)	75.0%
IHS FY 2004 Performance(baseline rate for children age 19-35 months)	72.0%
IHS FY 2004 Performance(rate for children age 3-27 months)	81.0%
IHS FY 2003 Performance(rate for children age 3-27 months)	80.0%
IHS FY 2002 Performance(rate for children age 3-27 months)	80.0%
HP 2020 goal for % of children age 19–35 months with 4:3:1:3:3:4 vaccines	80.0%
HP 2020 goal for % of children age 19–35 months with each individual vaccine	90.0%

For the IHS FY 2006 Performance (rate for children 19-35 months), the Percent (80.0) please consider: All 2002–2006 rates reported on this table were reported by the Immunization Program from the quarterly immunization reports. Effective in 2007, CRS reports the rate and not the Immunization Program. The CRS rate is reported using the CRS Active Immunization Package denominator.

### **Performance Improvement Tips**

- Providers should ask about and record off-site historical immunizations (IZ type, date received and location) on PCC forms. Data entry mnemonic: HIM
- Providers should document refusals; write "Refused" in appropriate vaccine order box on PCC form. Data entry mnemonic: REF (Immunization, Value, Date Refused).

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DU *** IHS 2012 Sele	ct ad		mber 25, 2			Fied Per		ge 51 *
1115 2012 5010	CCCO		INDIAN HO			ica kcp	,OI C	
Report	Per	iod: J	an 01, 201	.2 to	Dec 31, 20	12		
Previous Y						•		
Baselin	e Pe	riod:	Jan 01, 2	2000	to Dec 31,	20000		
Childhood Immunizations								
REP	ORT	%	PREV YR	%	CHG from	BASE	%	CHG from
PER	IOD		PERIOD		PREV YR %	PERIOD		BASE %
2 1 2 2 2								
Active Clinical Pts 19-35 months	56		39			55		
T9-33 MOHUIS	50		39			55		
# w/ 43133 combo								
or w/ Dx/ Contraind/								
NMI Refusal		19.6	3	7.7	+12.0	6	10.9	+8.7
A. # w/ Dx/Contraind/NMI Ref w/ % of								
	1	9 1	0	0 0	+9.1	0	0.0	+9.1
# w/ 43133	_	J. ±	Ü	0.0	10.1	Ü	0.0	
refusal	2	3.6	0	0.0	+3.6	0	0.0	+3.6
# w/ 431331 combo								
or w/ Dx/ Contraind/ NMI Refusal	0	16 1	2	7 7	+8.4	F	0 1	+7.0
NMI REIUSAI A. # w/ Dx/Contraind/NMI		10.1	3	1.1	+8.4	5	9.1	+ / . 0
Ref w/ % of								
Total 431331	0	0.0	0	0.0	+0.0	0	0.0	+0.0
# w/ 431331								
Refusal	1	1.8	0	0.0	+1.8	0	0.0	+1.8
# r.r / 4212214 gombo								
# w/ 4313314 combo or w/Dx/Contraind/								
NMI Refusal	3	5.4	0	0.0	+5.4	0	0.0	+5.4
A. # w/ Dx/Contraind/NMI	_	5.1	Ü			· ·	0.0	. 3 . 2
Ref w/ % of								
Total 4313314	0	0.0	0	0.0	+0.0	0	0.0	+0.0

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# w/ 4313314 Refusal	2	3.6	0	0.0	+3.6	0	0.0	+3.6	
<pre># w/ 4 doses DTaP   or w/ Contraind/   NMI Refusal A. # w/ Contraind/NMI   Ref w/ % of</pre>	16	28.6	3	7.7	+20.9	9	16.4	+12.2	
Total DTaP # w/ DTap Refusal	1 6	6.3 10.7	0	0.0	+6.3 +10.7	0	0.0	+6.3 +10.7	

Figure 2-31: Sample Report, Childhood Immunizations

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Childhood Immunizations: List of patients 19-35 months with IZ, if any. If a patient did not have all doses in a multiple dose vaccine, the IZ will not be listed. For example, if a patient only had 2 DTaP, no IZ will be listed for DTaP. NOTE: Because age is calculated at the beginning of the Report Period, the patient's age on the list will be between 7-23 months. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, ANDREA 000001 COMMUNITY #1 F 0 UP,AC,IMM 4 DTaP/DTP; 3 Polio; MMR; 3 Hib; 3 Hep B; Vari; 2 Influenza PATIENT2, HEATHER 000002 COMMUNITY #1 F 1 NMI DTaP/DTP; NMI Polio; 2 Hep A; 2 Influenza UP,AC,IMM PATIENT3, TONYA 000003 COMMUNITY #1 F 1 PATIENT4, JAMES 000004 COMMUNITY #1 M 0 UP,AC,IMM 3 Polio; MMR; 3 Hep B; Vari; NMI Rota PATIENTS, SCOTT 000005 COMMUNITY #1 M 0 UP, AC, IMM 3 Hib; 2 Hep A; 3 Rota; 2 Influenza

Figure 2-32: Sample Patient List, Childhood Immunizations

#### 2.4.4 Adolescent Immunizations

### **Denominators**

Active Clinical patients age 13.

Female *Active Clinical* patients age 13.

Active Clinical patients ages 13–17.

Female *Active Clinical* patients ages 13–17.

#### **Numerators**

Patients who have received the 2:3:1 combination (i.e., two MMR, three Hepatitis B, one Varicella), including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patient who have received the 1:3:2:1 combination (i.e., one Td/Tdap, three Hepatitis B, two MMR, one Varicella), including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received one dose of Tdap/Td ever, including contraindications and evidence of disease.

- a. Patients with (1) evidence of the disease, (2) a contraindication, or (3) a documented NMI refusal.
- b. Patients who have received one dose of Tdap ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received two doses of MMR ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received three doses of Hepatitis B ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received one dose of Varicella ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received one dose of meningococcal ever, including contraindications and evidence of disease.

**Note**: The only refusals included in this numerator are NMI refusals.

Patients who have received three doses of HPV ever, including contraindications and evidence of disease.

Note: The only refusals included in this numerator are NMI

refusals.

Note: Included for Female Active Clinical age 13 and Female

Active Clinical ages 13–17 only.

For each of the above numerators, the following subnumerators are included:

c. Patients with (1) evidence of the disease, (2) a contraindication, or (3) a documented NMI refusal.

Patients with documented REF refusal in PCC or Parent or Patient refusal in the IZ program.

### **Logic Description**

Age definition: Age of the patient is calculated at the beginning of the Report Period. Because IZ data comes from multiple sources, any IZ codes documented on dates within ten days of each other will be considered as the same immunization.

Dosage and types of immunization definitions:

- One dose of Td or Tdap
- Two doses of MMR: (1) two MMRs; (2) two M/R and two Mumps; (3) two R/M and two Measles; or (4) two each of Measles, Mumps, and Rubella
- Three doses of Hep B or two doses if documented with CPT 90743
- One dose of Varicella
- One dose of Meningococcal
- Three doses of HPV

Not Medically Indicated (NMI) refusals, evidence of disease and contraindications for individual immunizations will also count toward meeting the definition, as defined below. Refusals will count toward meeting the definition for refusal numerators only.

**Note**: NMI refusals are not counted as refusals; rather, they are counted as contraindications.

- Each immunization must be refused and documented separately. For example, if a patient has an NMI refusal for Rubella only, then there must be an immunization, contraindication, or separate refusal for the Measles and Mumps immunizations.
- For immunizations where required number of doses is >1, only one NMI refusal is necessary to be counted in the numerator. For example, if there is a single NMI refusal for Hepatitis B, the patient will be included in the numerator.
- For immunizations where required number of doses is >1, only one contraindication is necessary to be counted in the numerator. For example, if there is a single contraindication for HiB, the patient will be included in the numerator.
- Evidence of disease will be checked for at any time in the child's life (prior to the end of the Report Period).
- To be counted in subnumerator A, a patient must have evidence of disease, a contraindication, or an NMI refusal for any of the immunizations in the numerator. For example, if a patient was Rubella immune but had a Measles and Mumps immunization, the patient would be included in subnumerator A.

For the separate numerator for REF refusal (Patient Refusal for Service) in PCC or a Parent or Patient refusal in the IZ program, all conditions shown below must be met:

- Each immunization must be refused and documented separately. For example, if a patient has an REF refusal for Rubella, then there also must be an immunization, contraindication, or separate REF refusal for Measles and Mumps.
- Where the required number of doses is >1, only one REF refusal in PCC or one Parent or Patient refusal in the IZ program is necessary to be counted in the numerator. For example, for the 4 DTaP numerator, only one refusal is necessary to be counted in the refusal numerator.

Adolescent immunizations are defined in the following ways:

Immunization	CPT Codes	ICD and Other Codes
MMR	90707, 90710	Immunization codes: 3, 94
		<b>POV:</b> V06.4
		V Procedure: 99.48
		Contraindications: POV 279, V08, 042, 200-202, 203.0, 203.1, 203.8, 204-208; or Immunization Package contraindication of "Anaphylaxis," "Immune Deficiency," "Immune Deficient," or "Neomycin Allergy."  Refusals: Immunization codes 3, 94; CPT codes 90707, 90710
M/R (Measles/	90708	Immunization code: 4
Rubella)	90700	
raboliaj		Refusals: Immunization code 4; CPT code 90708
		Contraindications: Immunization Package contraindication of "Anaphylaxis."

Immunization	CPT Codes	ICD and Other Codes
R/M (Rubella/	90709 (old code)	Immunization code: 38
Mumps)		Refusals: Immunization code 38; CPT code 90709 (old code)
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Measles	90705	Immunization code: 5
		<b>POV:</b> V04.2
		V Procedure: 99.45
		<b>Evidence of Disease:</b> POV or PCC Problem List (active or inactive) 055*
		Refusals: Immunization code 5; CPT code 90705
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Mumps	90704	Immunization code: 7
		<b>POV:</b> V04.6
		V Procedure: 99.46
		<b>Evidence of Disease:</b> POV or PCC Problem List (active or inactive) 072*
		Refusals: Immunization code 7; CPT code 90704
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Rubella	90706	Immunization code: 6
		<b>POV:</b> V04.3
		V Procedure: 99.47
		<b>Evidence of Disease:</b> POV or PCC Problem List (active or inactive) 056*, 771.0
		Refusals: Immunization code 6; CPT code 90706
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Hepatitis B	90736, 90723,	Immunization codes: 8, 42-45, 51, 102, 104, 110, 132, 146
	90731 (old code), 90740, 90743-90748, G0010, Q3021, Q3023	<b>Evidence of Disease:</b> POV or PCC Problem List (active or inactive) V02.61, 070.2, 070.3
		<b>Refusals:</b> Immunization codes 8, 42-45, 51, 102, 104, 110, 132, 146; CPT codes 90736, 90723, 90731 (old code), 90740, 90743-90748, G0010, Q3021, Q3023
		Contraindications: Immunization Package contraindication of "Anaphylaxis."

Immunization	CPT Codes	ICD and Other Codes
Varicella	90710, 90716	Immunization codes: 21, 94
		POV: V05.4
		<b>Evidence of Disease:</b> 1) POV or PCC Problem List (active or inactive) 052*, 053*; or 2) Immunization Package contraindication of "Hx of Chicken Pox" or "Immune."
		Refusals: Immunization codes 21, 94; CPT codes 90710, 90716
		Contraindications: POV 279, V08, 042, 200-202, 203.0, 203.1, 203.8, 204-208; or Immunization Package contraindication of "Anaphylaxis," "Immune Deficiency," "Immune Deficient," or "Neomycin Allergy.
Tdap	90715	Immunization code: 115
		Refusals: Immunization code 115; CPT code 90715
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Td	90714, 90718	Immunization codes: 9, 113
		<b>POV:</b> V06.5
		Refusals: Immunization codes 9, 113; CPT codes 90714, 90718
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
Meningococcal	90733, 90734	Immunization codes: 32, 108, 114, 136
		<b>Refusals:</b> Immunization codes 32, 108, 114, 136; CPT codes 90733, 90734
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."
HPV	90649, 90650	Immunization codes: 62, 118, 137
		<b>Refusals:</b> Immunization codes 62, 118, 137; CPT codes 90649, 90650
		<b>Contraindications</b> : Immunization Package contraindication of "Anaphylaxis."

# **Key Logic Changes from CRS Version 11.1**

- Added CVX code 146 to Hepatitis B definition.
- Added CPT codes to refusal definitions.

### **Patient List Description**

List of patients 13–17 with IZ, if any. If a patient did not have all doses in a multiple dose vaccine, the IZ will not be listed. For example, if a patient only had two Hep B, no IZ will be listed for Hep B.

Note: An absent value in the Numerator column means the patient did not meet the requirements for any of the immunizations. An example for a female patient age 13 with a value of ";2 MMR" which means the patient did not have one Td/Tdap, three Hepatitis B, one Varicella, one Meningococcal, and three HPV immunizations.

#### **Measure Source**

HEDIS, HP 2020 IID-11

### Measure Past Performance and Long-Term Targets:

Target	Percent
HP 2020 goal for each individual IZ: Tdap, Meningococcal, HPV	80.0%
HP 2020 goal for each individual IZ: varicella	90.0%

### **Performance Improvement Tips**

- Providers should ask about and record off-site historical immunizations (IZ type, date received and location) on PCC forms. Data entry mnemonic: HIM
- Providers should document refusals; write "Refused" in appropriate vaccine Order box on PCC form. Data entry mnemonic: REF (Immunization, Value, Date Refused).

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DH
                                                                    Page 66
       *** IHS 2012 Selected Measures with Community Specified Report ***
                            DEMO INDIAN HOSPITAL
                 Report Period: Jan 01, 2012 to Dec 31, 2012
             Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                Baseline Period: Jan 01, 2000 to Dec 31, 2000
Adolescent Immunizations
                    REPORT
                              % PREV YR
                                            % CHG from BASE
                                                                   % CHG from
                              PERIOD
                    PERIOD
                                               PREV YR % PERIOD
                                                                      BASE %
Active Clinical patients
                        20
                                      17
age 13
                                                             28
# w/2:3:1 Combo or w/
 Dx/Contraind/
```

NMI Refusal A. # w/ Dx/ Contraind/ NMI Ref w/ % of Total	2	10.0	0	0.0	+10.0	0	0.0	+10.0	
2:3:1 # w/ 2:3:1	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Refusal	4	20.0	0	0.0	+20.0	0	0.0	+20.0	
# w/1:3:2:1 Combo or w/ Dx/Contraind/									
NMI Refusal A. # w/ Dx/ Contraind/	1	5.0	0	0.0	+5.0	0	0.0	+5.0	
NMI Ref w/ % of Total 1:3:2:1 # w/ 1:3:2:1	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Refusal	4	20.0	0	0.0	+20.0	0	0.0	+20.0	
# w/ 1 dose Tdap/Td or w/ Dx/ Contraind/									
NMI Refusal A. # w/ Dx/ Contraind/ N	3 MI	15.0	4	23.5	-8.5	6	21.4	-6.4	
Ref w/ % of Total Tdap/Td B. # w/ Tdap or w/	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Dx/ Contraind/ NMI Refusal	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
# w/ Tdap/Td Refusal	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
# w/ 2 doses MMR or w/									
DX/ Contraind/ NMI Refusal A. # w/ Dx/ Contraind/ NMI Ref w/ % of	6	30.0	0	0.0	+30.0	0	0.0	+30.0	
Total MMR # w/ MMR Refusal	0 2	0.0	0	0.0	+0.0 +10.0	0	0.0	+0.0 +10.0	

Figure 2-33: Sample Report, Adolescent Immunizations

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Adolescent Immunizations: List of patients 13-17 with IZ, if any. If a patient did not have all doses in a multiple dose vaccine, the IZ will not be listed. For example, if a patient only had 2 Hep B, no IZ will be listed for Hep B. HRN PATIENT NAME COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1,LINDA 000001 COMMUNITY #3 F 13 2 MMR; 2 Hep B + 90743AC 2 MMR; 2 Hep B + 9074 PATIENT2, SHERRY 000002 COMMUNITY #3 F 13 NMI Tdap; NMI MMR; NMI Hep B; NMI Vari PATIENT22, JESSICA 000022 COMMUNITY #4 F 13 Ref MMR; 3 Hep B; Evid Vari PATIENT23, SAMANTHA 000023 COMMUNITY #4 F 13 Td; 3 Hep B; NMI Meningococcal 000024 COMMUNITY #4 PATIENT24,NINA F 13

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AC	Contra MMR; Contra Vari
PATIENT25,RHONDA	000025 COMMUNITY #4 F 13
AC	Td; 3 HPV

Figure 2-34: Sample Patient List, Adolescent Immunizations

# 2.5 Childhood Diseases Group

# 2.5.1 Appropriate Treatment for Children with Upper Respiratory Infection

#### **Denominators**

Active Clinical patients who were ages three months through 18 years of age who were diagnosed with an *upper respiratory infection* during the period 6 months (180 days) prior to the report period through the first six months of the report period.

*User Population patients* who were ages three months through 18 years of age who were diagnosed with an *upper respiratory infection* during the period 6 months (180 days) prior to the report period through the first six months of the report period.

#### Numerator

Patients who were *not* prescribed an antibiotic on or within three days after diagnosis. In this measure, appropriate treatment is *not* to receive an antibiotic.

### **Logic Description**

Age is calculated as follows: Children three months as of six months (180 days) of the year prior to the Report Period to 18 years as of the first six months of the Report Period.

In order to be included in the denominator, *all* of the following conditions must be met:

- 1. Patient's diagnosis of an upper respiratory infection (URI) must have occurred at an outpatient visit. Upper Respiratory Infection defined as POV 460 or 465.\*. Outpatient visit defined as Service Category A, S, or O.
- 2. If outpatient visit was to Clinic Code 30 (Emergency Medicine), it must not have resulted in a hospitalization, defined as Service Category H, either on the same day or the next day with URI diagnosis.
- 3. Patient's visit must only have a diagnosis of URI. If any other diagnosis exists, the visit will be excluded.

- 4. The patient did not have a new or refill prescription for antibiotics within 30 days prior to the URI visit date.
- 5. The patient did not have an active prescription for antibiotics as of the URI visit date. "Active" prescription defined as:

```
Rx Days Supply >= (URI Visit Date–Prescription Date)
```

If multiple visits exist that meet the above criteria, the first visit will be used.

Antibiotic medications defined with medication taxonomy BGP HEDIS ANTIBIOTIC MEDS or V Procedure 99.21. (Medications are: Amoxicillin, Amox/Clavulanate, Ampicillin, Azithromycin, Cefaclor, Cefadroxil hydrate, Cefazolin, Cefdinir, Cefixime, Cefditoren, Ceftibuten, Cefpodoxime proxetil, Cefprozil, Ceftriaxone, Cefuroxime, Cephalexin, Cephradine, Ciprofloxacin, Clindamycin, Dicloxacillin, Doxycycline, Erythromycin, Ery E-Succ/Sulfisoxazole, Gatifloxacin, Levofloxacin, Lomefloxacin, Loracarbef, Minocycline, Ofloxacin, Penicillin VK, Penicillin G, Sparfloxacin, Sulfisoxazole, Tetracycline, Trimethoprim, Trimethoprim-Sulfamethoxazol.) Medications must not have a comment of RETURNED TO STOCK.

### **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients three months to 18 years of age with upper respiratory infection, with antibiotic prescription, if any.

#### **Measure Source**

**HEDIS** 

### Measure Past Performance and Long-Term Targets

None

```
DIJ
                             November 25, 2012
                                                                      Page 72
       *** IHS 2012 Selected Measures with Community Specified Report ***
                             DEMO INDIAN HOSPITAL
                  Report Period: Jan 01, 2012 to Dec 31, 2012
              Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                 Baseline Period: Jan 01, 2000 to Dec 31, 2000
Appropriate Treatment for Children with Upper Respiratory Infection (con't)
                     REPORT
                                   PREV YR
                                              % CHG from BASE
                                                                     % CHG from
                                                 PREV YR % PERIOD
                     PERIOD
                                   PERIOD
                                                                        BASE %
```

Active Clinical 3 months-18 yrs								
w/Upper Respiratory Infection	38		36			30		
# w/o Antibiotic Rx	37	97.4	35	97.2	+0.1	27	90.0	+7.4
User Pop 3 months-18 yrs w/Upper Respiratory Infection	43		38			35		
# w/o Antibiotic	13		30			33		
Rx	42	97.7	37	97.4	+0.3	32	91.4	+6.2

Figure 2-35: Sample Report, Appropriate Treatment for Children with Upper Respiratory Infection

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Appropriate Treatment for Children with Upper Respiratory Infection: List of patients 3 months to 18 years with upper respiratory infection, with antibiotic prescription, if any.

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

PATIENT1, PAMELA 00001 COMMUNITY #3 F 3

UP, AC MEETS MEASURE
PATIENT2, ALICIA 000002 COMMUNITY #3 F 7

UP, AC MEETS MEASURE
PATIENT3, JAMES 000003 COMMUNITY #3 M 0

UP, AC MEETS MEASURE
PATIENT4, HENRY 000004 COMMUNITY #3 M 12

UP, AC MEETS MEASURE
PATIENT25, HEATHER 000025 COMMUNITY #4 F 7

UP, AC MEETS MEASURE
PATIENT26, DYLAN 000026 COMMUNITY #4 M 3

UP, AC MEETS MEASURE
PATIENT27, CODY 000027 COMMUNITY #4 M 4

UP, AC MEETS MEASURE
PATIENT27, CODY 000027 COMMUNITY #4 M 4

UP, AC MEETS MEASURE
PATIENT28, KAREN 000028 COMMUNITY #5 F 0

UP, AC MEETS MEASURE
PATIENT27, KAREN 000028 COMMUNITY #5 F 0

up, AC MEETS MEASURE
PATIENT28, KAREN 000028 COMMUNITY #5 F 0

up, AC MEETS MEASURE
PATIENT28, KAREN 000028 COMMUNITY #5 F 0

up, AC MEETS MEASURE

Figure 2-36: Sample Patient List, Appropriate Treatment for Children with Upper Respiratory Infection

# 2.5.2 Appropriate Testing for Children with Pharyngitis

#### **Denominators**

Active Clinical patients who were ages 2–18 years who were diagnosed with pharyngitis and prescribed an antibiotic during the period six months (180 days) prior to the Report Period through the first six months of the Report Period.

*User Population patients* who were ages 2–18 years who were diagnosed with *pharyngitis* and prescribed an antibiotic during the period six months (180 days) prior to the Report Period through the first six months of the Report Period.

#### **Numerator**

Patients who received a Group A strep test.

### **Logic Description**

Age is calculated as follows: Children two years as of six months (180 days) of the year prior to the Report Period to 18 years as of the first six months of the Report Period.

In order to be included in the denominator, *all* of the following conditions must be met:

- 1. Patient's diagnosis of pharyngitis must have occurred at an outpatient visit. Pharyngitis defined as POV 462, 463, or 034.0. Outpatient visit defined as Service Category A, S, or O.
- 2. If outpatient visit was to Clinic Code 30 (Emergency Medicine), it must not have resulted in a hospitalization, defined as Service Category H, either on the same day or the next day with pharyngitis diagnosis.
- 3. Patient's visit must only have a diagnosis of pharyngitis. If any other diagnosis exists, the visit will be excluded.
- 4. The patient did not have a new or refill prescription for antibiotics within 30 days prior to the pharyngitis visit date.
- 5. The patient did not have an active prescription for antibiotics as of the pharyngitis visit date. "Active" prescription defined as:
- 6. Rx Days Supply >= (URI Visit Date Prescription Date)
- 7. The patient filled a prescription for antibiotics on or within three days after the pharyngitis visit.

If multiple visits exist that meet the above criteria, the first visit will be used.

Antibiotic medications defined with medication taxonomy BGP HEDIS ANTIBIOTIC MEDS or V Procedure 99.21. (Medications are: Amoxicillin, Amox/Clavulanate, Ampicillin, Azithromycin, Cefaclor, Cefadroxil hydrate, Cefazolin, Cefdinir, Cefixime, Cefditoren, Ceftibuten, Cefpodoxime proxetil, Cefprozil, Ceftriaxone, Cefuroxime, Cephalexin, Cephradine, Ciprofloxacin, Clindamycin, Dicloxacillin, Doxycycline, Erythromycin, Ery E-Succ/Sulfisoxazole, Gatifloxacin, Levofloxacin, Lomefloxacin, Loracarbef, Minocycline, Ofloxacin, Penicillin VK, Penicillin G, Sparfloxacin, Sulfisoxazole, Tetracycline, Trimethoprim, Trimethoprim-Sulfamethoxazol.) Medications must not have a comment of RETURNED TO STOCK.

To be included in the numerator, a patient must have received a Group A Streptococcus test within the seven-day period beginning three days prior through three days after the Pharyngitis visit date.

*Group A Streptococcus* test defined as: CPT 87430 (by enzyme immunoassay), 87650-87652 (by nucleic acid), 87880 (by direct optical observation), 87081 (by throat culture), 3210F (Group A Strep Test); site-populated taxonomy BGP GROUP A STREP TESTS; and LOINC taxonomy.

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients 2–18 years of age with pharyngitis and a Group A Strep test, if any.

#### **Measure Source**

**HEDIS** 

#### Measure Past Performance and Long-Term Targets

None

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                             DEMO INDIAN HOSPITAL
                 Report Period: Jan 01, 2012 to Dec 31, 2012
              Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                Baseline Period: Jan 01, 2000 to Dec 31, 2000
Appropriate Testing for Children with Pharyngitis (con't)
                     REPORT
                                  PREV YR
                                              % CHG from BASE
                                                                     % CHG from
                                  PERIOD
                                                PREV YR % PERIOD
                                                                        BASE %
Active Clinical 2-18 yrs w/
```

Pharyngitis and Antibiotic Rx	10		5			8		
# w/Group A Strep Test	9	90.0	3	60.0	+30.0	2	25.0	+65.0
User Pop 2-18 yrs w/ Pharyngitis and Antibiotic Rx	11		7			10		
# w/Group A Strep Test	9	81.8	4	57.1	+24.7	2	20.0	+61.8

Figure 2-37: Sample Report, Appropriate Testing for Children with Pharyngitis

UP-User Pop; AC-Active Clinical; AD-Active Diabetic; AAD-Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Appropriate Testing for Children with Pharyngitis: List of patients 2-18 years with pharyngitis and a Group A Strep test, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, MICHAEL 000001 COMMUNITY #1 M 9 03/19/12 RAPID ANTIGEN (STREP A) UP,AC 03/19/12 RAPID ANTIGER PATIENT2,JOSEPH 000002 COMMUNITY #1 M 12 05/01/12 RAPID ANTIGEN (STREP A) UP,AC PATIENT3, LESTER 000003 COMMUNITY #1 M 13 PATIENT24, MONICA 000024 COMMUNITY #2 F 5 01/23/12 RAPID ANTIGEN (STREP A) PATIENT25, MICHAEL JAMES 000025 COMMUNITY #2 M 7 UP,AC 03/12/12 RAPID ANTIGEN (STREP A)

Figure 2-38: Sample Patient List, Appropriate Testing for Children with Pharyngitis

# 2.6 Cancer Related Measure Topics

# 2.6.1 Cancer Screening: Pap Smear Rates

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 59.5% for the proportion of female patients ages 21 through 64 without a documented history of hysterectomy who have had a Pap screen within the previous three years.

#### **Denominators**

Female Active Clinical patients ages 21 through 64 without documented history of Hysterectomy. (GPRA Denominator)

*Female User Population patients* ages 21 through 64 without a documented history of Hysterectomy.

### **Numerators**

Patients with a Pap smear documented in the past three years.

**Note**: This numerator does *not* include refusals. (GPRA Denominator)

Patients with documented Pap smear refusal in past year.

### **Logic Description**

Age of the patient is calculated at the beginning of the report period. Patients must be at least 21 years of age at the beginning of the report period and less than 65 years of age as of the end of the report period.

	CPT Codes	II( I) and ()ther ( odes	LOINC Codes	Taxonomy
Hysterectomy	51925, 56308 (old code), 57540, 57545, 57550, 57555, 57556, 58150, 58152, 58200- 58294, 58548, 58550-58554, 58570-58573, 58951, 58953- 58954, 58956, 59135	V Procedure: 68.4-68.8 V POV: 618.5, V88.01, V88.03 Women's Health: Procedure called Hysterectomy.		

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Pap Smear	88141-88167, 88174-88175, G0123, G0124, G0141, G0143- G0145, G0147, G0148, P3000, P3001, Q0091	V Lab: PAP SMEAR POV: V67.01 Follow-up Vaginal Pap Smear V76.2 Screen Mal Neop- Cervix V72.32 Encounter for Pap Cervical Smear to Confirm Findings of Recent Normal Smear Following Initial Abnormal Smear V72.3 Gynecological Examination, Pap Cervical Smear as Part of General Gynecological Exam, Pelvic Exam (annual) (periodic) (old code, to be counted for visits prior to 10/1/04 only) V76.47 Vaginal Pap Smear for Post-Hysterectomy Patients 795.0*, 795.10-16, 795.19 V Procedure: 91.46 Women's Health: Procedure called Pap Smear and where the result does NOT have "ERROR/DISREGARD"	Yes	BGP PAP SMEAR TAX
Refusal (in past year)	Refusal of codes: 88141- 88167, 88174- 88175, G0123, G0124, G0141, G0143-G0145, G0147, G0148, P3000, P3001, Q0091	Refusals: Lab Test Value Pap Smear		

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of women 21-64 with documented Pap Smear or refusal, if any.

### **Measure Source**

HP 2020 C-15

# **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	58.1%
IHS FY 2010 Performance	59.0%
IHS FY 2009 Performance	59.0%
IHS FY 2008 Performance	59.0%
IHS FY 2007 Performance	59.0%
IHS FY 2006 Performance	59.0%
IHS FY 2005 Performance	60.0%
IHS FY 2004 Performance	58.0%
IHS FY 2003 Performance	61.0%
IHS FY 2002 Performance	62.0%
HP 2020 Goal	93.0%

# **Performance Improvement Tips**

- Providers should ask about and record off-site tests (date received and location) on PCC forms. Data entry mnemonic: HPAP
- Providers should document refusals; write "Refused" in Pap Order box on PCC form. Data entry mnemonic: REF (Lab Test Value, Date Refused).

DU November 25, 2012 Page 77  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
	REPORT	%	PREV YR		CHG from PREV YR %				
Female Active Clinica 21-64 yrs (GPRA)	='		377			320			
# w/Pap Smear recorded w/in 3 years-No Refu	sals								
(GPRA)	202	40.4	179	47.5	-7.1	147	45.9	-5.5	
# w/ Pap Smear Refusal	8	1.6	0	0.0	+1.6	0	0.0	+1.6	
# Female User Pop 21-64 years	833		699			614			
<pre># w/Pap Smear recorder w/in 3 years -No Refusals # w/ Pap Smear</pre>		26.4	196	28.0	-1.6	159	25.9	+0.5	

Refusal 8 1.0 0 0.0 +1.0 0 0.0 +1.0
-------------------------------------

Figure 2-39: Sample Report, Cancer Screening: Pap Smear Rates

Figure 2-40: Sample Patient List, Cancer Screening: Pap Smear rates

# 2.6.2 Cancer Screening: Mammogram Rates

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 51.7% for the proportion of female patients ages 52 through 64 who have had mammography screening within the last two years.

#### **Denominators**

Female Active Clinical patients ages 52 through 64 without a documented history of bilateral mastectomy or two separate unilateral mastectomies. (GPRA Denominator)

Female Active Clinical patients ages 42 and older without a documented history of bilateral mastectomy or two separate unilateral mastectomies.

Female User Population patients ages 52 through 64 without a documented history of bilateral mastectomy or two separate unilateral mastectomies.

Female User Population patients ages 42 and older without a documented history of bilateral mastectomy or two separate unilateral mastectomies. (GPRA Developmental Denominator)

#### **Numerators**

All patients who had a Mammogram documented in the past two years.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

Patients with documented mammogram refusal in the past year

### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period. For all denominators, patients must be at least the minimum age as of the beginning of the Report Period. For the 52–64 denominators, the patients must be less than 65 years of age as of the end of the Report Period.

	CPT Codes	ICD and Other Codes
Bilateral Mastectomy	19300.50-19307.50 or old code 19180, 19200, 19220 <b>OR</b> 19300-19307 w/modifier 09950 (.50 and 09950 modifiers indicate bilateral) <b>OR</b> 19240, with modifier of .50 or 09950	V Procedure: 85.42, 85.44, 85.46, 85.48
Unilateral Mastectomy	Must have 2 separate occurrences on 2 different dates of service. 19300-19307, or old codes 19180, 19200, 19220, 19240	Must have 2 separate occurrences on 2 different dates of service. 85.41, 85.43, 85.45, 85.47
Mammogram	V Rad or VCPT: 77052-77059, 76090 (old code), 76091 (old code), 76092 (old code), G0206, G0204, G0202	POV: V76.11, V76.12, 793.80 Abnormal mammogram, unspecified; 793.81 Mammographic microcalcification; 793.89 Other abnormal findings on radiological exam of breast V Procedure: 87.36-87.37 Women's Health: Mammogram Screening, Mammogram Dx Bilat, Mammogram Dx Unilat and where the mammogram result does NOT have
		"ERROR/DISREGARD"
Refusal (in past year)	<b>V Rad Mammogram for CPT:</b> 77052-77059, 76090 (old code), 76091 (old code), 76092 (old code), G0206, G0204, G0202	

# **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of women 42+ with mammogram/refusal, if any.

### **Measure Source**

HP 2020 C-17

### **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	49.8%
IHS FY 2010 Performance	48.0%
IHS FY 2009 Performance	45.0%
IHS FY 2008 Performance	45.0%
IHS FY 2007 Performance	43.0%
IHS FY 2006 Performance	41.0%
IHS FY 2005 Performance	41.0%
IHS FY 2004 Performance	40.0%
IHS FY 2003 Performance	40.0%
IHS FY 2002 Performance	42.0%
HP 2020 Goal	81.1%

# **Performance Improvement Tips**

- Providers should ask about and record off-site mammogram procedures (date received and location) on PCC forms. Data entry mnemonic: HRAD.
- Providers should document refusals; write "Refused" in Mammogram Order box on PCC form. Data entry mnemonic: REF (Mammogram, Procedure (CPT) Code, Date Refused).

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Cancer Screening: Mammog	ram Rate	s (con't)						
		PREV YR PERIOD		CHG from E			CHG from BASE %	
Female Active Clinical 52-64 (GPRA)	97	60			47			
# w/Mammogram recorded w 2 years-No Refusals	/in							
V = /	31 32.	0 22	36.7	-4.7	22	46.8	-14.8	
# w/ Mammogram Refusal	7 7.	2 0	0.0	+7.2	0	0.0	+7.2	
# Female Active Clinical	# Female Active Clinical							

42+ (GPRA Dev.)	293		188			163			
# w/Mammogram recorded									
w/in 2 years-No Refusa	ıls								
(GPRA Dev.)	63	21.5	62	33.0	-11.5	54	33.1	-11.6	
# w/ Mammogram									
Refusal	9	3.1	0	0.0	+3.1	0	0.0	+3.1	
# Female User Pop	1.0.0		100			100			
52-64	175		122			102			
# w/Mammogram recorded									
w/in 2 years-No									
Refusals	34	19.4	26	21.3	-1.9	23	22.5	-3.1	
# w/ Mammogram	31	10.1	20	21.5	1.5	23	22.5	3.1	
Refusal	7	4.0	0	0.0	+4.0	0	0.0	+4.0	
# Female User Pop									
42+	513		380			333			
# w/Mammogram recorded									
w/in 2 years-No									
Refusals	67	13.1	69	18.2	-5.1	58	17.4	-4.4	
# w/ Mammogram									
Refusal	9	1.8	0	0.0	+1.8	0	0.0	+1.8	

Figure 2-41: Sample Report, Cancer Screening: Mammogram rates

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Cancer Screening: Mammogram Rates: List of women 42+ with mammogram/refusal, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, CARLA 000001 COMMUNITY #1 F 43 UP, AC 10/01/11 CPT 77052 PATIENT2, CRYSTAL 000002 COMMUNITY #1 F 42 PATIENT3, ALEXA 000003 COMMUNITY #1 F 45
UP, AC 04/24/11 CPT 76090
PATIENT4, HANNAH 000004 COMMUNITY #1 F 42 PATIENT5, MARTHA 000005 COMMUNITY #1 F 43 UP
PATIENT6, TARA 000006 COMMUNITY #1 F 44
UP, AC 01/15/12 Refused CPT (
PATIENT7, CAROL LYNN 000007 COMMUNITY #1 F 44 01/15/12 Refused CPT G0206 PATIENT8, MARY ANN 03/05/12 RAD 76092 000008 COMMUNITY #1 F 52 UP,AC PATIENT9, BARBARA 000009 COMMUNITY #1 F 52 04/22/12 CPT 77057 UP,AC

Figure 2-42: Sample Patient List, Cancer Screening: Mammogram rates

# 2.6.3 Colorectal Cancer Screening

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 43.2% for the proportion of clinically appropriate patients ages 51–80 who have received colorectal screening.

#### **Denominators**

All Active Clinical patients ages 51–80 without a documented diagnosis of colorectal cancer or total colectomy. Broken down by gender. (GPRA Denominator)

All *User Population patients* ages 51–80 without any documented diagnosis of colorectal cancer or total colectomy.

#### **Numerators**

Patients who have had any CRC screening, defined as any of the following: (1) Fecal Occult Blood Test (FOBT) or Fecal Immunochemical Test (FIT) during the Report period; (2) flexible sigmoidoscopy or double contrast barium enema in the past five years; or (3) colonoscopy in the past 10 years.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

Patients with documented CRC screening refusal in the past year.

Patients with FOBT or FIT during the Report Period.

Patients with a flexible sigmoidoscopy or double contrast barium enema in the past five years or a colonoscopy in the past 10 years.

Patients with a flexible sigmoidoscopy in the past five years or a colonoscopy in the past 10 years.

Patients with a flexible sigmoidoscopy and double contrast barium enema in the past five years or a colonoscopy in the past 10 years.

### **Logic Description**

Age is calculated at the beginning of the Report Period.

**Denominator Exclusions** 

Any diagnosis ever of one of the following:

- 1. **Colorectal Cancer:** POV: 153.\*, 154.0, 154.1, 197.5, V10.05; CPT G0213–G0215 (old codes), G0231 (old code).
- 2. **Total Colectomy:** CPT 44150–44151, 44152 (old code), 44153 (old code), 44155–44158, 44210–44212; V Procedure 45.8 (old code).

Colorectal cancer screening definition: The most recent of any of the following tests and procedures during applicable timeframes. CRS identifies the tests and procedures described in the numerators above with the following codes:

# Colorectal Cancer Screening (CRS looks for the most recent of any of the following during timeframes specified in numerator section above)

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Fecal Occult Blood lab test (FOBT) or Fecal Immuno- chemical Test (FIT)	82270, 82274, 89205 (old code), G0107 (old code), G0328, G0394 (old code)		Yes	BGP GPRA FOB TESTS
Flexible Sigmoidoscopy	45330-45345, G0104	V Procedure: 45.24		
Double contrast barium enema	VRad 74280, G0106, G0120			
Colonoscopy	44388-44394, 44397, 45355, 45378-45387, 45391, 45392, G0105, G0121	V POV: V76.51 Colon screening V Procedure: 45.22, 45.23, 45.25, 45.42, 45.43		

# Refusal definition: Any of the following in the past year:

	CPT Codes	ICD and Other Codes	Taxonomy
Refusals	FOBT or FIT: 82270, 82274, 89205 (old code), G0107 (old code), G0328, or G0394 (old code)	Flexible Sigmoidoscopy V Procedure:	V Lab Fecal Occult Blood Test
	Flexible Sigmoidoscopy: 45330-45345, G0104 DCBE: 74280, G0106, G0120	45.24, 45.42 Colonoscopy	
	<b>Colonoscopy:</b> 44388-44394, 44397, 45355, 45378-45387, 45391, 45392, G0105, G0121	V Procedure: 45.22, 45.23, 45.25, 45.42, 45.43	

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients 51–80 with CRC screening or refusal, if any.

#### **Measure Source**

HEDIS, HP 2020 C-16

### **Measure Past Performance and Long-Term Targets:**

Performance	Percent
IHS FY 2011 Performance	41.7%
IHS FY 2010 Performance	37.0%
IHS FY 2009 Performance	33.0%
IHS FY 2008 Performance	29.0%
IHS FY 2007 Performance	26.0%
IHS FY 2006 Performance	22.0%
HP 2020 Goal	70.5%

### **Performance Improvement Tip**

Providers should ask about and record off-site historical tests (test type, date received and location) on PCC forms. Data entry mnemonics: HBE (barium enema); HCOL (colonoscopy); HFOB (Fecal Occult Blood); HSIG (sigmoidoscopy). Providers should also enter as a refusal if the patient refuses the colorectal cancer screening. Refusals may be entered with the data entry mnemonic of REF (refusal).

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                Report Period: Jan 01, 2012 to Dec 31, 2012
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               Baseline Period: Jan 01, 2000 to Dec 31, 2000
Colorectal Cancer Screening (con't)
                   REPORT % PREV YR % CHG from BASE % CHG from PREV YR % PERIOD BASE %
                   PERIOD PERIOD
AC Pts 51-80 w/o
colorectal cancer or total
colectomy (GPRA) 316
                                    195
                                                          152
# w/ CRC screening
-No Refusals (GPRA) 64 20.3
                                     49 25.1
                                                  -4.9
                                                          28 18.4
                                                                        +1.8
# w/ CRC Screening
```

Refusal # w/FOBT/FIT during	10	3.2	0	0.0	+3.2	0	0.0	+3.2
Report period # w/Flex Sig, DCBE,	12	3.8	11	5.6	-1.8	0	0.0	+3.8
or Colonoscopy # w/Flex Sig or	55	17.4	40	20.5	-3.1	28	18.4	-1.0
Colonoscopy	49	15.5	31	15.9	-0.4	20	13.2	+2.3
# w/Flex Sig & DCBE or Colonoscopy	46	14.6	29	14.9	-0.3	18	11.8	+2.7
Male Active Clinical								
51-80	151		93			65		
# w/ CRC screening								
-No Refusals	27	17.9	18	19.4	-1.5	9	13.8	+4.0
# w/ CRC Screening Refusal	5	3.3	0	0.0	+3.3	0	0.0	+3.3
# w/FOBT/FIT during Report period	7	4.6	3	3.2	+1.4	0	0.0	+4.6
<pre># w/Flex Sig, DCBE,   or Colonoscopy</pre>	22	14.6	15	16.1	-1.6	9	13.8	+0.7
# w/Flex Sig or Colonoscopy # w/Flex Sig & DCBE	20	13.2	14	15.1	-1.8	8	12.3	+0.9
or Colonoscopy	20	13.2	14	15.1	-1.8	8	12.3	+0.9

Figure 2-43: Sample Report, Colorectal Cancer Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Colorectal Cancer Screening: List of patients 51-80 with CRC screening or refusal, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, DANIELLE 000001 COMMUNITY #1 F 51
UP FOB: 08/19/12 CPT G0107
PATIENT2, MARIE 000002 COMMUNITY #1 F 51
UP, AC COLO: 02/12/12 Refused CPT
PATIENT3, MARY ANN 000003 COMMUNITY #1 F 52
UP, AC PATIENT4, BOBBIE 000004 COMMUNITY #1 F 52
UP, AC BE: 01/25/12 RAD 74280
PATIENT5, WINONA 000005 COMMUNITY #1 F 53 BE: 01/25/12 RAD 74280 UP,AC PATIENT6, DARLENE 000006 COMMUNITY #1 F 54 SIG: 04/07/09 45.24; BE: 03/31/10 RAD 74280 PATIENT7, JOYCE 000007 COMMUNITY #1 F 57 UP,AC COLO: 07/07/12 POV V76.51

Figure 2-44: Sample Patient List, Colorectal Cancer Screening

# 2.6.4 Comprehensive Cancer Screening

### **GPRA Measure Description**

Increase the proportion of patients ages 21–80 who received a comprehensive cancer screening.

### **Denominators**

Active Clinical patients ages 21–80 who are eligible for cervical cancer, breast cancer, and/or colorectal cancer screening. (GPRA Developmental Denominator)

- a. Active Clinical female patients ages 21–80.
- b. Active Clinical male patients ages 51–80.

#### **Numerators**

Patients who have had all screenings for which they are eligible.

**Note**: This numerator does *not* include refusals. (GPRA Developmental Numerator)

- a. Female patients with cervical cancer, breast cancer, and/or colorectal cancer screening.
- b. Male patients with colorectal cancer screening.

### **Logic Description**

Age is calculated at the beginning of the Report Period.

Cervical Cancer Screening definition: To be eligible for this screening, patients must be female Active Clinical ages 21 through 64 and not have a documented history of hysterectomy. Patients must be at least 21 years of age at the beginning of the Report Period and less than 65 years of age as of the end of the Report Period. To be counted as having the screening, the patient must have had a Pap Smear documented in the past three years.

CRS identifies the tests and procedures described in the numerators above with the following codes:

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Hysterectomy	51925, 56308 (old code), 57540, 57545, 57550, 57555, 57556, 58150, 58152, 58200-58294, 58548, 58550-58554, 58570-58573, 58951, 58953-58954, 58956, 59135	V Procedure: 68.4-68.8 V POV: 618.5, V88.01, V88.03 Women's Health: Procedure called Hysterectomy.		
Pap Smear	88141-88167, 88174-88175, G0123, G0124, G0141, G0143- G0145, G0147, G0148, P3000, P3001, Q0091	V Lab: PAP SMEAR POV: V67.01 Follow-up Vaginal Pap Smear V76.2 Screen Mal Neop-Cervix V72.32 Encounter for Pap Cervical Smear to Confirm Findings of Recent Normal Smear Following Initial Abnormal Smear V72.3 Gynecological Examination, Pap Cervical Smear as Part of General Gynecological Exam, Pelvic Exam (annual) (periodic) (old code, to be counted for visits prior to 10/1/04 only) V76.47 Vaginal Pap Smear for Post-Hysterectomy Patients 795.0*, 795.10-16, 795.19 V Procedure: 91.46 Women's Health: Procedure called Pap Smear	Yes	BGP PAP SMEAR TAX

Breast Cancer Screening definition: To be eligible for this screening, patients must be female Active Clinical ages 52 through 64 and not have a documented history ever of bilateral mastectomy or two separate unilateral mastectomies. Patients must be at least age 52 as of the beginning of the Report Period and must be less than 65 years of age as of the end of the Report Period. To be counted as having the screening, the patient must have had a Mammogram documented in the past two years.

CRS identifies the tests and procedures described in the numerators above with the following codes:

	CPT Codes	ICD and Other Codes
Bilateral Mastectomy	19300.50-19307.50 or old code 19180, 19200, 19220 <b>OR</b> 19300-19307 w/modifier 09950 (.50 and 09950 modifiers indicate bilateral) <b>OR</b> 19240, with modifier of .50 or 09950	V Procedure: 85.42, 85.44, 85.46, 85.48
Unilateral Mastectomy	Must have 2 separate occurrences on 2 different dates of service. 19300-19307, or old codes 19180, 19200, 19220, 19240	Must have 2 separate occurrences on 2 different dates of service. 85.41, 85.43, 85.45, 85.47
Mammogram	<b>V Rad or VCPT:</b> 77052-77059, 76090 (old code), 76091 (old code), 76092 (old code), G0206, G0204, G0202	POV: V76.11, V76.12, 793.80 Abnormal mammogram, unspecified; 793.81 Mammographic microcalcification; 793.89 Other abnormal findings on radiological exam of breast
		V Procedure: 87.36-87.37 Women's Health: Mammogram
		Screening, Mammogram Dx Bilat, Mammogram Dx Unilat and where the mammogram result does NOT have "ERROR/DISREGARD".

Colorectal cancer screening definition: To be eligible for this screening, patients must be Active Clinical ages 51–80 and not have a documented history ever of colorectal cancer or total colectomy. To be counted as having the screening, patients must have had any of the following: (1) FOBT or FIT during the Report Period; (2) flexible sigmoidoscopy or double contrast barium enema in the past 5 years; or (3) colonoscopy in the past 10 years.

The most recent of any of the following tests and procedures during applicable timeframes. CRS identifies the tests and procedures described in the numerators above with the following codes:

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Colorectal Cancer	G0213-G0215, G0231	<b>V POV:</b> 153.*, 154.0, 154.1, 197.5, V10.05		
Total Colectomy	44150-44151, 44152 (old code), 44153 (old code), 44155-44158, 44210-44212	V Procedure: 45.8 (old code)		
Fecal Occult Blood lab test (FOBT) or Fecal Immuno- chemical Test (FIT)	82270, 82274, 89205 (old code), G0107 (old code), G0328, G0394 (old code)		Yes	BGP GPRA FOB TESTS

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Flexible Sigmoidoscopy	45330-45345, G0104	V Procedure: 45.24		
Double contrast barium enema	VRad 74280, G0106, G0120			
Colonoscopy	44388-44394, 44397, 45355, 45378-45387, 45391, 45392, G0105, G0121	V POV: V76.51 Colon screening V Procedure: 45.22, 45.23, 45.25, 45.42, 45.43		

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients 21–80 with comprehensive cancer screening, if any.

#### **Measure Source**

Not Available

## Measure Past Performance and Long-Term Targets

None

### **Performance Improvement Tip**

- Providers should ask about and record off-site Pap tests (date received and location) on PCC forms. Data entry mnemonic: HPAP
- Providers should ask about and record off-site mammogram procedures (date received and location) on PCC forms. Data entry mnemonic: HRAD.
- Providers should ask about and record off-site historical colorectal cancer tests (test type, date received and location) on PCC forms. Data entry mnemonics: HBE (barium enema); HCOL (colonoscopy); HFOB (Fecal Occult Blood); HSIG (sigmoidoscopy).

```
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DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000

Comprehensive Cancer Screening

REPORT % PREV YR % CHG from BASE % CHG from
```

PE	RIOD		PERIOD		PREV YR %	PERIOD	В	ASE %	
Active Clinical 21-80 (GPRA Dev.)	721		509			424			
# w/ Comprehensive Cand Screening-No Refusals (GPRA Dev.)		31.3	195	38.3	-7.0	153	36.1	-4.7	
A. Female 21-80	570		416			359			
A. # Female w/all Screens	199	34.9	177	42.5	-7.6	144	40.1	-5.2	
B. Male 51-80	151		93			65			
B. # Male w/CRC Screen	27	17.9	18	19.4	-1.5	9	13.8	+4.0	

Figure 2-45: Sample Report, Comprehensive Cancer Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Comprehensive Cancer Screening: List of patients 21-80 with comprehensive cancer screening, if any. DENOMINATOR HRN COMMUNITY SEX AGE PATIENT NAME NUMERATOR PATIENT1, DANIELLE 000001 COMMUNITY #1 F 21 PAP: 05/05/11 POV 795
PATIENT2, MARIE 000002 COMMUNITY #1 F 51
AC.PAP PAP: 05/05/11 POV 795.0 AC,PAP
PATIENT3,MARY ANN
AC,MAM
PATIENT4,BOBBIE
AC,PAP,MAM,CRCS
PATIENT5,WINONA
AC,PAP,MAM,CRCS
PATIENT6,HARRY
AC,CRCS
PATIENT6,HARRY
AC,CRCS
PATIENT7,LARRY

O00007 COMMUNITY #1 F 57
CRCS: 07/20/12 POV V76
MAM: 10/01/11 CPT 7705
CRCS: 04/07/08 Proc 45
PATIENT7,LARRY
O00007 COMMUNITY #1 M 57 AC, PAP MAM: 07/06/11 CPT 77055 CRCS: 07/20/12 POV V76.51 MAM: 10/01/11 CPT 77052 CRCS: 04/07/08 Proc 45.24 AC, CRCS PATIENT8, BARRY 000008 COMMUNITY #1 M 63 CRCS: 02/18/12 CPT 45330 AC, CRCS

Figure 2-46: Sample Patient List, Comprehensive Cancer Screening

## 2.6.5 Tobacco Use and Exposure Assessment

#### **Denominators**

Active Clinical patients ages five and older. Broken down by gender and age groups (5–13, 14–17, 18–24, 25–44, 45–64, and 65 and older), based on HP 2010 age groups.

All Pregnant female User Population patients with no documented miscarriage or abortion.

All *User Population patients* ages five and older. Broken down by gender.

#### **Numerators**

Patients who have been screened for tobacco use during the Report Period.

Patients identified as current tobacco users during the Report Period, both smokers and smokeless users.

- a. Patients identified as current smokers during the Report Period.
- b. Patients identified as current smokeless tobacco users during the Report Period.

Patients identified as exposed to environmental tobacco smoke (ETS) (second-hand smoke) during the Report Period.

## **Logic Description**

Ages are calculated at beginning of Report Period.

For screening, an additional eight months is included for patients who were pregnant during the Report Period but who had their tobacco assessment prior to that.

CRS uses the following codes to define the denominators and numerators:

	CPT Codes	ICD and Other Codes
Pregnancy (At least two visits during the past 20 months. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the past 20 months, CRS will use the first two visits in the 20-month period. The patient must not have a documented miscarriage or abortion occurring after the second pregnancy-related visit. In addition, the patient must have at least one pregnancy-related visit occurring during the reporting period.) An additional 8 months is included for patients who were pregnant during the Report period but who had their tobacco assessment prior to that.		<b>V POV:</b> V22.0-V23.9, V72.42, 640.*-649.*, 651.*-676.*
Miscarriage (after second pregnancy POV in past 20 months)	59812, 59820, 59821, 59830	<b>V POV:</b> 630, 631, 632, 633*, 634*
Abortion (after second pregnancy POV in past 20 months)	59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260-S2267	V POV: 635*, 636* 637* Procedure: 69.01, 69.51, 74.91, 96.49
Screened for Tobacco Use (time frame for pregnant patients is past 20 months)	D1320, 99406, 99407, G0375 (old code), G0376 (old code), G8453 (old code), G8455-G8457 (old codes), G8402 (old code), 1034F (Current Tobacco Smoker), 1035F (Current Smokeless Tobacco User), 1036F (Current Tobacco Non-User), 1000F (Tobacco Use Assessed)	V POV or current Active Problem List: 305.1, 305.1* (old codes), 649.00-649.04, V15.82  Patient Education codes: "TO-", "-TO", "-SHS", 305.1, 305.1* (old codes), 649.00-649.04, V15.82, D1320, 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, 1036F, 1000F, G8455-G8457 (old codes), G8402 (old code), or G8453 (old code)  Dental code: 1320
Tobacco Users (time frame for pregnant patients is past 20 months)	99406, 99407, G0375 (old code), G0376 (old code), 1034F (Current Tobacco Smoker), 1035F (Current Smokeless Tobacco User), G8455 (old code), G8456 (old code), G8453 (old code)	V POV or current Active Problem List: 305.1, 305.10-305.12 (old codes), or 649.00-649.04  Dental code: 1320

	CPT Codes	ICD and Other Codes
Current Smokers (time frame for pregnant patients is past 20 months)	99406, 99407, G0375 (old code), G0376 (old code), 1034F (Current Tobacco Smoker), G8455 (old code), G8402 (old code), G8453 (old code)	V POV or current Active Problem List: 305.1, 305.10-305.12 (old codes), or 649.00-649.04  Dental code: 1320
Current Smokeless (time frame for pregnant patients is past 20 months)	1035F (Current Smokeless Tobacco User), G8456 (old code)	

For numerator definitions, all existing national Tobacco, TOBACCO (SMOKING), TOBACCO (SMOKELESS-CHEWING/DIP), and TOBACCO (EXPOSURE) Health Factors are listed below with the numerator to which they apply.

Health Factor	Numerator
Ceremonial	Screened (does NOT count as Smoker)
Cessation-Smokeless	Screened; Tobacco Users; Smokeless User
Cessation-Smoker	Screened; Tobacco Users; Smoker
Current Smokeless	Screened; Tobacco Users; Smokeless User
Current Smoker	Screened; Tobacco Users; Smoker
Current Smoker, status unknown	Screened; Tobacco Users; Smoker
Current smoker, every day	Screened; Tobacco Users; Smoker
Current smoker, some day	Screened; Tobacco Users; Smoker
Non-Tobacco User	Screened
Previous Smokeless	Screened
Previous (Former) Smokeless	Screened
Previous Smoker	Screened
Previous (Former) Smoker	Screened
Smoke Free Home	Screened
Smoker In Home	Screened; ETS
Current Smoker & Smokeless	Screened; Tobacco Users; Smoker; Smokeless User
Exposure To Environmental Tobacco Smoke	Screened; ETS

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients five and older with documented tobacco screening, if any.

## **Measure Source**

HP 2020 TU-1.1 Cigarette smoking 18 and older; TU-1.2 Smokeless tobacco use 18 and older; TU-11 Exposure to ETS-nonsmokers three and older

# **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance (Screening)	62.0%
IHS FY 2010 Performance (Screening)	60.0%
IHS FY 2009 Performance (Screening)	57.0%
IHS FY 2008 Performance (Screening)	54.0%
IHS FY 2005 Performance (Screening)	34.0%
IHS FY 2004 Performance (Screening)	27.0%

Performance	Percent
IHS FY 2008 Performance (Tobacco Users)	29.0%
HP 2020 Goals: TU-1.1 (Cigarette smoking 18 and older): 12%; TU-1.2 (Smokeless tobacco use 18 and older): 0.3%; TU-11 (Exposure to ETS-non smokers 18 and older): 68%	

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Tobacco Use and Exposure Ass	sessment (con't	)			
		% CHG from PREV YR			
% PREV YR % CHG from PERIOD		HG from PREV YR	% PERIOD	:	BASE %
# Active Clinical Pts					
=> 5 1,357	1,031		911		
# w/Tobacco					
Screening 624	46.0 426	41.3 +4.7	328	36.0	+10.0
# Tobacco Users w/ % of					
Total Screened 294	47.1 165	38.7 +8.4	130	39.6	+7.5
A. # Smokers w/ % of Total Tobacco Users 277 B. # Smokeless Tobacco Users w/ % of Total	94.2 164	99.4 -5.2	130	100.0	-5.8
Tobacco Users 29	9.9 5	3.0 +6.8	3	2.3	+7.6

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<pre># exposed to ETS/ smoker in home w/ % of</pre>									
	4	0.6	2	0.5	+0.2	1	0.3	+0.3	
# Male Active Clinical									
ages => 5	570		430			374			
# w/Tobacco									
Screening		40.4	152	35.3	+5.0	128	34.2	+6.1	
<pre># Tobacco Users w/ % of Total Screened</pre>		59.1	69	45.4	+13.7	59	46.1	+13.0	
A. # Smokers w/ % of		37.1	0,5	10.1	. 20.7	3,2		123.0	
Total Tobacco Users	124	91.2	69	100.0	-8.8	59	100.0	-8.8	
B. # Smokeless Tobacco Users w/ % of Total									
	22	16.2	2	2.9	+13.3	3	5.1	+11.1	
<pre># exposed to ETS/ smoker in home w/ % of</pre>									
Total Screened	1	0.4	0	0.0	+0.4	1	0.8	-0.3	
# Female Active Clinical	l								
ages => 5	787		601			537			
# w/Tobacco									
Screening		50.1	274	45.6	+4.5	200	37.2	+12.8	
<pre># Tobacco Users w/ % of Total Screened</pre>		40.1	96	35.0	+5.1	71	35.5	+4.6	
A. # Smokers w/ % of		10.1	, ,	33.0			33.3		
Total Tobacco Users	153	96.8	95	99.0	-2.1	71	100.0	-3.2	
B. # Smokeless Tobacco Users w/ % of Total									
Tobacco Users	7	4.4	3	3.1	+1.3	0	0.0	+4.4	
<pre># exposed to ETS/ smoker in home w/ % of</pre>									
Total Screened	3	0.8	2	0.7	+0.0	0	0.0	+0.8	

Figure 2-47: Sample Report, Tobacco Use Assessment Tobacco Use and Exposure Assessment

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Tobacco Use and Exposure	e Asses	ssment	(con't)				
			_	Distri	bution	65 and	al dan
	5-13	14-1/	18-24	25-44	45-64	os and	older
CURRENT REPORT PERIOD							
# Active Clinical	171	74	177	448	371	116	
# w/Tobacco Screening	10	19	102	243	199	51	
% w/Tobacco Screening	5.8	25.7	57.6	54.2	53.6	44.0	
# Tobacco Users	3	8	46	118	104	15	

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% Tobacco Users w/ % of							
Total Screened		42.1	45.1	48.6	52.3	29.4	
# Smokers	2	8	43	108	102	14	
% Smokers w/ % of							
Total Tobacco Users	66.7	100.0	93.5	91.5	98.1	93.3	
# Smokeless	1	0	3	15	8	2	
% Smokeless w/ % of	_	· ·	J		ŭ	_	
Total Tobacco Users	33 3	0 0	6 5	12 7	7 7	13.3	
TOTAL TODACCO OBCID	33.3	0.0	0.5	,	, • ,	13.3	
# ETS/Smk Home	0	0	0	3	1	0	
% ETS/Smk Home w/ % of		· ·	· ·	3	_	Ŭ	
Total Screened		0.0	0 0	1 2	0.5	0.0	
100al bol celled	0.0	0.0	0.0	1.2	0.5	0.0	
PREVIOUS YEAR PERIOD							
	176	62	168	323	238	64	
# Active Clinical # w/Tobacco Screening	11	14	27	147	12Ω		
% w/Tobacco Screening					53.8		
. w/iobacco screening	0.3	44.0	51.0	10.0	55.0	00.9	
# Tobacco Users	Λ	5	41	60	5.0	9	
% Tobacco Users w/ % of		3	-11	00	30		
Total Screened		35.7	17 1	40 0	20 1	22 1	
Total Screened	0.0	33.7	4/.1	40.0	39.1	23.1	
# Smokers	0	5	40	60	50	9	
% Smokers w/ % of	U	5	40	00	50	9	
Total Tobacco Users	0 0	100 0	07 6	100 0	100 0	100 0	
TOTAL TODACCO USELS	0.0	100.0	97.0	100.0	100.0	100.0	
# Smokeless	0	0	1	2	1	0	
% Smokeless w/ % of	U	U	1	3	1	U	
Total Tobacco Users	0 0	0 0	2 4	E 0	2 0	0.0	
TOTAL TODACCO USERS	0.0	0.0	2.4	5.0	2.0	0.0	
# ETC/Cmlr Homo	0	0	0	2	0	0	
# ETS/Smk Home		U	U	2	0	U	
% ETS/Smk Home w/ % of		0.0	0 0	1 4	0 0	0 0	
Total Screened	0.0	0.0	0.0	1.4	0.0	0.0	
CHANCE EDOM DDEM 37D 0							
CHANGE FROM PREV YR %	0 4	. 2 1		. 0 7	0 1	17 0	
# w/Tobacco Screening					-0.1		
Tobacco Users					+13.2		
	+66.7	+0.0			-1.9		
	+33.3	+0.0			+5.7		
ETS	+0.0	+0.0	+0.0	-0.1	+0.5	+0.0	

Figure 2-48: Sample Age Breakdown Report, Tobacco Use Assessment

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Tobacco Use and Exposure Assessment: List of patients 5 and older with documented tobacco screening, if any.

PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR

PATIENT1, CHESTER 000001 COMMUNITY #1 M 7

UP, AC 01/10/12 SCREEN
PATIENT2, JUAN 000002 COMMUNITY #1 M 19
```

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```
PATIENT3, BEN
                      000003 COMMUNITY #1 M 22
ΠP
PATIENT4, MARY
                      000004 COMMUNITY #1 F 35
UP, AC, PREG
                            04/10/12 SCREEN, 04/10/12 USER, 04/10/12 SMOKELESS
                      000005 COMMUNITY #1 M 13
PATIENT5, HARRY B
                            03/15/12 SCREEN
PATIENT6, EMERSON
                      000006 COMMUNITY #1 M 15
                            05/21/12 SCREEN, 05/21/11 USER, 05/21/12 SMOKER,
UP,AC
05/21/12 ETS
PATIENT7, EUGENE JAY
                      000007 COMMUNITY #1 M 29
                      000008 COMMUNITY #1 M 31
PATIENT8, ROGER
UP,AC
                           01/21/12 SCREEN, 01/21/12 USER, 01/21/12 SMOKER
PATIENT9, ANDREW
                       000009 COMMUNITY #1 M 42
```

Figure 2-49: Sample Patient List, Tobacco Use Assessment

### 2.6.6 Tobacco Cessation

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 30.0% for the proportion of tobacco-using patients who receive tobacco cessation intervention.

#### **Denominators**

Active Clinical patients identified as current tobacco users prior to the Report Period. Broken down by gender and age groups (<12, 12–17, 18 and older). (GPRA Denominator)

*User Population patients* identified as *current tobacco users* prior to the report period. Broken down by gender.

#### **Numerators**

Patients who have received tobacco cessation counseling or received a prescription for a smoking cessation aid during the Report Period.

```
Note: This numerator does not include refusals. (GPRA Numerator)
```

Patients who refused tobacco cessation counseling during the Report Period.

Patients identified during the Report Period as having quit their tobacco use.

Patients who have received tobacco cessation counseling, received a prescription for a smoking cessation aid, or who quit their tobacco use during the Report Period.

**Note**: This numerator does *not* include refusals.

# **Logic Description**

Age is calculated at the beginning of the Report Period.

	ICD and Other Codes
Tobacco Users (documented prior to the Report Period)	Tobacco Health Factors (looks at the last documented health factor in the Tobacco, TOBACCO (SMOKING) and TOBACCO (SMOKELESS—CHEWING/DIP) categories): Current Smoker, Current Smokeless, Current Smoker and Smokeless, Cessation-Smoker, Cessation-Smoker, Cessation-Smoker, Cessation-Smoker, every day, or Current smoker, some day
	CPT code (looks at the last documented): 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, G8455 (old code), G8456 (old code), G8402 (old code) or G8453 (old code)
	V POV or current Active Problem List (looks at the last documented): 305.1, 305.10-305.12 (old codes), or 649.00-649.04
Tobacco Cessation Counseling (documented during the Report Period)	Patient education codes containing: "TO-", "-TO", "-SHS", 305.1, 305.1* (old codes), 649.00-649.04, D1320, 99406, 99407, G0375 (old code), G0376 (old code), 4000F, G8402 (old code) or G8453 (old code)
	Dental code: 1320
	Clinic code: 94 (tobacco cessation clinic)
	<b>CPT code:</b> D1320, 99406, 99407, G0375 (old code), G0376 (old code), 4000F, G8402 (old code) or G8453 (old code)
Prescription for Tobacco Cessation Aid (documented	<b>Taxonomy:</b> Medication in the site-populated BGP CMS SMOKING CESSATION MEDS taxonomy
during the Report Period)	Medication Name: Any medication with name containing "NICOTINE PATCH", "NICOTINE POLACRILEX", "NICOTINE INHALER", or "NICOTINE NASAL SPRAY"
	CPT Code: 4001F
Quit Tobacco User (documented during the Report Period)	<b>V POV or current Active Problem List:</b> 305.13 Tobacco use in remission (old code) or V15.82
	Health Factors documented during the Report Period (looks at the last documented health factor): Previous Smoker, Previous Smokeless, Previous (Former) Smoker, Previous (Former) Smokeless.

### **Tobacco Cessation Counseling Refusal Definition**

Documented refusal of patient education code containing "TO-", "-TO", or "-SHS" or CPT code D1320, 99406, 99407, G0375 (old code), G0376 (old code), 4000F, G8402 (old code) or G8453 (old code). Refusals will only be counted if a patient did not receive counseling or a prescription for tobacco cessation aid.

## Key Logic Changes from CRS Version 11.1

None

### **Patient List Description**

List of tobacco users with tobacco cessation intervention, if any, or who have quit tobacco use.

### **Measure Source**

Smoking Cessation Attempts: HP 2020 TU-4; Smoking Cessation Counseling: HP 2020 TU-10

### **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	29.4%
IHS FY 2010 Performance	25.0%
IHS FY 2009 Performance	24.0%
IHS FY 2008 Performance	21.0%
IHS FY 2007 Performance	16.0%
IHS FY 2006 Performance	12.0%
HP 2020 goal for increasing smoking cessation attempts for adult smokers	80.0%

```
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                               DEMO INDIAN HOSPITAL
                   Report Period: Jan 01, 2012 to Dec 31, 2012
              Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                 Baseline Period: Jan 01, 2000 to Dec 31, 2000
Tobacco Cessation (con't)
                      REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %
Active Clinical Tobacco
                                         236
                         301
                                                                 182
Users (GPRA)
# w/tobacco cessation counseling
 or Rx for cessation-No
```

Refusals (GPRA)	54	17.9	46	19.5	-1.6	48	26.4	-8.4	
# w/refusal of									
counseling	8	2.7	0	0.0	+2.7	0	0.0	+2.7	
# who quit	13	4.3	3	1.3	+3.0	1	0.5	+3.8	
# w/ cessation counseli	ng,								
cessation aid, or quit									
-No Refusals	66	21.9	48	20.3	+1.6	49	26.9	-5.0	
Male Active Clinical									
Tobacco Users	140		117			94			
# w/tobacco cessation c	ounse	ling							
or RX for cessation-No									
Refusals	29	20.7	20	17.1	+3.6	25	26.6	-5.9	
# w/refusal of									
counseling	2	1.4	0	0.0	+1.4	0	0.0	+1.4	
# who quit	5	3.6	1	0.9	+2.7	1	1.1	+2.5	
# w/ cessation counseli	ng,								
cessation aid, or quit									
-No Refusals	34	24.3	21	17.9	+6.3	26	27.7	-3.4	
Female Active Clinical									
Tobacco Users	161		119			88			
# w/tobacco cessation c	ounse	ling							
or RX for cessation-No									
Refusals	25	15.5	26	21.8	-6.3	23	26.1	-10.6	
# w/refusal of									
counseling	6	3.7	0	0.0	+3.7	0	0.0	+3.7	
# who quit	8	5.0	2	1.7	+3.3	0	0.0	+5.0	
# w/ cessation counseli	ng,								
cessation aid, or quit									
-No Refusals	32	19.9	27	22.7	-2.8	23	26.1	-6.3	

Figure 2-50: Sample Report, Tobacco Cessation

Tobacco Cessation (con't)							
ACTIVE CLINICAL TOBACCO USERS  Age Distribution							
	<12	12-17	=>18				
CURRENT REPORT PERIOD Active Clin Tobacco Users	0	6	295				
<pre># w/tobacco cessation counseling   or Rx for cessation aid   -No Refusals % w/ tobacco cessation counseling</pre>	0	0	54				
or Rx for cessation aid -No Refusals	0.0	0.0	18.3				
# w/refusal of counseling	0	0	8				
% w/refusal of counseling	0.0	0.0	2.7				

# who quit	0	1	12	
% who quit	0.0	16.7	4.1	
% WIIO QUIC	0.0	10.7	7.1	
<pre># w/tobacco cessation counseling,</pre>				
or Rx for cessation aid or quit				
-No Refusals	0	1	65	
% w/ tobacco cessation counseling,				
or Rx for cessation aid or quit				
-No Refusals	0.0	16.7	22.	
-NO Relusais	0.0	10.7	22.	
PREVIOUS YEAR PERIOD				
Active Clin Tobacco Users	1	4	231	
# w/tobacco cessation counseling				
or Rx for cessation aid				
-No Refusals	0	0	46	
	U	U	40	
% w/tobacco cessation counseling				
or Rx for cessation aid				
-No Refusals	0.0	0.0	19.9	
# w/refusal of				
counseling	0	0	0	
% w/refusal of	U	0	0	
	0 0	0 0	0 0	
counseling	0.0	0.0	0.0	
# who quit	0	0	3	
% who quit	0.0	0.0	1.3	
<pre># w/tobacco cessation counseling,</pre>				
Rx for cessation aid or quit				
-No Refusals	0	0	48	
		U	40	
% w/ tobacco cessation counseling,				
Rx for cessation aid or quit				
-No Refusals	0.0	0.0	20.8	
CHANGE FROM PREV YR %				
w/tobacco cessation counseling				
or Rx for cessation aid				
	+0.0	+0.0	-1.6	
	.0.0	+0.0	-1.0	
w/refusal of	. 0 . 0	2	o =	
3	+0.0	+0.0	+2.7	
	+0.0	+16.7	+2.8	
w/tobacco cessation counseling,				
Rx for cessation aid or quit				
=	+0.0	+16.7	+1.3	
		. 10.7		

Figure 2-51: Sample Age Breakdown Report, Tobacco Cessation

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient

Tobacco Cessation: List of tobacco users with tobacco cessation intervention,
if any, or who have quit tobacco use.

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

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```
PATIENT1, BRITNEY 000001 COMMUNITY #1 F 22
                           COUNSEL/RX: 06/10/12 CPT G0375
PATIENT2, LORETTA
                      000002 COMMUNITY #1 F 22
UP,AC
                           COUNSEL/RX: 01/13/12 305.1-DP
PATIENT3, HALEY 000003 COMMUNITY #1 F 25
UP,AC
                           COUNSEL/RX: 02/19/12 TO-LA
PATIENT4, ANGEL 000004 COMMUNITY #1 F 30
UP,AC
                           COUNSEL/RX: 03/05/12 CPT 4000F
PATIENT5, JOYCE 000005 COMMUNITY #1 F 31
                           QUIT: PREVIOUS (FORMER) SMOKER 05/31/12
UP,AC
PATIENT6, ESTHER
                      000006 COMMUNITY #1 F 32
UP,AC
                           COUNSEL/RX: 03/05/12 CESSATION MED - NICOTINE 14MG
TRANSDERMAL PATCH PATIENT7, SARAH
                      000007 COMMUNITY #1 F 33
UP,AC
PATIENT8, PAULA
                      000008 COMMUNITY #1 F 34
                            COUNSEL/RX: 03/17/12 TO-QT
UP,AC
```

Figure 2-52: Sample Patient List Tobacco Cessation

# 2.7 Behavioral Health Related Performance Measure Topics

# 2.7.1 Alcohol Screening (FAS Prevention)

## **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 58.7% for the proportion of female patients ages 15 to 44 who receive screening for alcohol use.

### **Denominators**

Female Active Clinical patients ages 15 to 44. (GPRA Denominator)

Female User Population patients ages 15 to 44.

### **Numerators**

Patients screened for alcohol use, had an alcohol-related diagnosis or procedure, or received alcohol-related patient education during the report period.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

- a. Patients with alcohol screening during the report period.
- b. Patients with alcohol-related diagnosis or procedure during the report period
- c. Patients with alcohol-related patient education during the report period.
- d. Patients with documented alcohol screening refusal in past year.

## **Logic Description**

Ages are calculated at beginning of Report Period.

Alcohol screening definition: Any of the following during the Report Period: (a) Alcohol Screening Exam, any CAGE Health Factor, or Screening Diagnosis; (b) Alcohol-related diagnosis in POV, Current PCC or BHS Problem List; (c) Alcohol-related procedure; or (d) Patient education.

	ICD and Other Codes
Alcohol Screening	PCC Exam Code: 35
	<b>CPT code:</b> 99408, 99409, G0396, G0397, H0049, H0050, 3016F
	Any CAGE Health Factor
	V POV: V11.3 (history of alcoholism), V79.1 (screening for
	alcoholism)
	BHS Problem Code: 29.1 (Screening for Alcoholism)
	V Measurement in PCC or BHS: AUDT, AUDC, or CRFT
Alcohol-related	V POV, Current PCC or BHS Problem List: 303.*, 305.0*,
Diagnosis	291.*, 357.5*
	BHS POV: 10, 27, 29
Alcohol-related	<b>V Procedure:</b> 94.46, 94.53, 94.61-94.63, 94.67-94.69
Procedure	
Alcohol-related	Patient Education codes: "AOD-" or "-AOD", "CD-" or "-CD" (old
Education	codes), or containing V11.3, V79.1, 303.*, 305.0*, 291.* 357.5*,
	99408, 99409, G0396, G0397, H0049, or H0050, or 3016F

Alcohol screening may be documented with either an exam code or the CAGE health factor in PCC or BHS. BHS problem codes can also currently be used.

Refusal definition: Refusal of PCC Exam Code 35 in the past year

## **Recommended Brief Screening Tool**

Single Alcohol Screening Question (SASQ) (below).

For Women:

When was the last time you had more than four drinks in one day?

For Men:

When was the last time you had more than five drinks in one day?

Any time in the past three months is a positive screen; further evaluation indicated. Provider should note the screening tool used was the SASQ in the COMMENT section of the Exam Code.

#### **Alcohol Health Factors**

The existing Health Factors for alcohol screening are based on the CAGE questionnaire, which asks the following four questions:

- 1. Have you ever felt the need to Cut down on your drinking?
- 2. Have people Annoyed you by criticizing your drinking?
- 3. Have you ever felt bad or Guilty about your drinking?
- 4. Have you ever needed an **E**ye opener the first thing in the morning to steady your nerves or get rid of a hangover?
- 5. Based on how many YES answers are received, document Health Factor:
  - HF-CAGE 0/4 (all "No" answers)
  - HF-CAGE 1/4
  - HF-CAGE 2/4
  - HF–CAGE 3/4
  - HF-CAGE 4/4

#### Optional values:

- Level/Severity: Mild, Moderate, or Severe
- Quantity: # of drinks daily

### **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of female patients with documented alcohol screening or refusal, if any.

#### **Measure Source**

HP 2010 16-17a

### Measure Past Performance and Long-term Targets

Performance	Percent
IHS FY 2011 Performance	57.8%
IHS FY 2010 Performance	55.0%
IHS FY 2009 Performance	52.0%
IHS FY 2008 Performance	47.0%
IHS FY 2007 Performance	41.0%

Performance	Percent
IHS FY 2006 Performance	28.0%
IHS FY 2005 Performance	11.0%
IHS FY 2004 Performance	7.0%

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Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Alcohol Screening (FAS	 Preve	ntion)							
	PORT RIOD				CHG from PREV YR %			CHG from BASE %	
Female Active Clinical ages 15-44 (GPRA)	434		348			304			
# w/ alcohol screening/ Dx/Proc/Pt Ed		11 г	2	0.6	.10.0	1	0 2	.11 0	
-No Refusals (GPRA) A. w/alcohol	50	11.5	2	0.6	+10.9	1	0.3	+11.2	
screening B. # w/alcohol related	43	9.9	1	0.3	+9.6	0	0.0	+9.9	
Dx or procedure	3	0.7	1	0.3	+0.4	1	0.3	+0.4	
C. # w/alcohol related patient education	10	2.3	0	0.0	+2.3	0	0.0	+2.3	
<pre># w/alcohol screening   refusal</pre>	3	0.7	0	0.0	+0.7	0	0.0	+0.7	
Female User Population ages 15-44	725		636			588			
# w/ alcohol screening/	Dx/								
Proc/Pt Ed -No Refusals A. # w/alcohol	54	7.4	2	0.3	+7.1	2	0.3	+7.1	
screening	46	6.3	1	0.2	+6.2	0	0.0	+6.3	
B. # w/alcohol related Dx or procedure	4	0.6	1	0.2	+0.4	2	0.3	+0.2	
C. # w/alcohol related patient education	11	1.5	0	0.0	+1.5	0	0.0	+1.5	
# w/alcohol screening						-			
refusal	3	0.4	0	0.0	+0.4	0	0.0	+0.4	

Figure 2-53: Sample Report, Alcohol Screening (FAS Prevention)

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

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```
Alcohol Screening (FAS Prevention): List of female patients with
documented alcohol screening or refusal, if any.
PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR
PATIENT1, CHRISTINE S 000001 COMMUNITY #1 F 15
PATIENT2, RITA A 000002 COMMUNITY #1 F 15
                           SCREEN: 03/06/12
UP,AC
POV V11.3
POV V11.3

PATIENT3, DIANE L 000003 COMMUNITY #1 F 15

SCREEN: 02/02/12 Refr
                                SCREEN: 02/02/12 Refused Ex 35
PATIENT4, ALICIA 000004 COMMUNITY #1 F 15
IJP . AC
DP,AC
PATIENT5,MELISSA

UP,AC
PATIENT6,LISA MARIE

UP,AC

DO0005 COMMUNITY #1 F 16
PT ED: 02/13/12 99408-P
O00006 COMMUNITY #1 F 16
UP,AC

SCREEN: 10/13/12 HF CAGE 1/4
PATIENT7, RUTH NELLIE 000007 COMMUNITY #1 F 16
PATIENT8, ALISHA DAWN 000008 COMMUNITY #1 F 16
              SCREEN: 03/03/12 CPT 3016F
UP,AC
```

Figure 2-54: Sample Patient List, Alcohol Screening (FAS Prevention)

# 2.7.2 Alcohol Screening and Brief Intervention (ASBI) in the ER

#### **Denominators**

Number of visits for *Active Clinical patients* age 15–34 seen in the ER for injury during the report period. Broken down by gender and age groups (15–24 and 25–34).

Number of visits for *Active Clinical patients* age 15–34 seen in the ER for injury and screened positive for hazardous alcohol use during the report period. Broken down by gender and age groups (15–24 and 25–34).

Number of visits for *User Population patients* age 15–34 seen in the ER for injury during the report period. Broken down by gender and age groups (15–24 and 25–34).

Number of visits for *User Population patients* age 15–34 seen in the ER for injury and screened positive for hazardous alcohol use during the report period. Broken down by gender and age groups (15–24 and 25–34).

#### **Numerators**

Number of visits where patients were screened in the ER for hazardous alcohol use.

a. Number of visits where patients were screened positive.

Number of visits where patients were provided a brief negotiated interview (BNI) at or within seven days of the ER visit.

- a. Number of visits where patients were provided a BNI at the ER visit.
- b. Number of visits where patients were provided a BNI not at the ER visit but within 7 days of the ER visit.

## **Logic Description**

Age of the patient is calculated as of the beginning of the Report Period.

Emergency room visit definition: Clinic Code 30.

Multiple visits definition: If a patient has multiple ER visits for injury during the Report Period, each visit will be counted in the denominator. For the screening numerator, each ER visit with injury at which the patient was screened for hazardous alcohol use will be counted. For the positive alcohol use screen numerator, each ER visit with injury at which the patient screened positive for hazardous alcohol use will be counted. For the BNI numerators, each visit where the patient was either provided a BNI at the ER or within seven days of the ER visit will be counted. An example of this logic is shown below.

	Denom	Scrn	Pos Scrn	BNI Num
ER Visit w/Injury	Count	Num	Num Count	Count
*****	******	*****	*****	*****
John Doe, 07/17/09,	Screened Posi	tive at	ER, BNI at E	R
John Doe, 09/01/09,	Screened Posi	tive at	ER, No BNI	
John Doe, 11/15/09,	No Screen			
******	******	*****	*****	* * * * * * * *
COUNTS:	3	2	2	1

CRS uses the following codes:

	ICD and Other Codes						
Injury	<b>V POV (primary or secondary):</b> 800.0–999.9 or E800.0–E989.						
ER Screening for	Any conducted during an ER visit:						
Hazardous Alcohol	PCC Exam Code: 35						
Use	Any Alcohol Health Factor (i.e., CAGE)						
	/ POV: V79.1 Screening for Alcoholism						
	<b>CPT:</b> G0396, G0397, H0049, 99408, 99409, 3016F						
	V Measurement in PCC: AUDT, AUDC, or CRFT						
Positive Screen for	Any of the following for the screening conducted during an ER visit:						
Hazardous Alcohol	PCC Exam Code: 35 Alcohol Screening result of "Positive"						
Use	Health Factor: CAGE result of 1/4, 2/4, 3/4 or 4/4						
	<b>CPT:</b> G0396, G0397, 99408, 99409						
	<b>V Measurement Result in PCC:</b> AUDT result of => 8, AUDC result of => 4 for men and =>3 for women, CRFT result of 2-6						

	ICD and Other Codes
BNI	Any of the following documented at the ER visit or within seven days of the ER visit at a face-to-face visit, which excludes chart reviews and telecommunication visits:
	<b>CPT:</b> G0396, G0397, H0050, 99408, 99409
	Patient Education Code: AOD-BNI or containing G0396, G0397, H0050, 99408, or 99409

## **Recommended Brief Screening Tool**

SASQ (below).

For Women:

When was the last time you had more than four drinks in one day?

For Men:

When was the last time you had more than five drinks in one day?

Any time in the past three months is a positive screen; further evaluation indicated. Provider should note the screening tool used was the SASQ in the COMMENT section of the Exam Code.

### **Alcohol Health Factors**

The existing Health Factors for alcohol screening are based on the CAGE questionnaire, which asks the following four questions:

- 1. Have you ever felt the need to Cut down on your drinking?
- 2. Have people Annoyed you by criticizing your drinking?
- 3. Have you ever felt bad or **G**uilty about your drinking?
- 4. Have you ever needed an **E**ye opener the first thing in the morning to steady your nerves or get rid of a hangover?

Based on how many YES answers are received, document Health Factor:

- HF–CAGE 0/4 (all No answers)
- HF–CAGE 1/4
- HF-CAGE 2/4
- HF–CAGE 3/4
- HF–CAGE 4/4

# Optional values:

- Level/Severity: Mild, Moderate, or Severe
- Quantity: # of drinks daily

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of visits for patients seen in the ER for an injury, with screening for hazardous alcohol use, results of screen and BNI, if any.

## **Measure Source**

None

# **Measure Past Performance and Long-Term Targets**

None

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Alcohol Screening and Br	ief	Interv	rention (AS	BBI)	in the ER				
REPORT % PREV YR % CHG from BASE % CHG from PERIOD PREV YR % PERIOD BASE %									
# ER Injury Visits for AC Pts 15-34	33		33			32			
<pre># Visits w/ ER Hazardous Alcohol Screening A. # Visits w/Positive</pre>		57.6	0	0.0	+57.6	0	0.0	+57.6	
Screen	15	45.5	0	0.0	+45.5	0	0.0	+45.5	
# ER Injury Visits for Male AC Pts 15-34	12		18			20			
<pre># Visits w/ ER Hazardous Alcohol Screening A. # Visits w/Positive</pre>		58.3	0	0.0	+58.3	0	0.0	+58.3	
Screen	5	41.7	0	0.0	+41.7	0	0.0	+41.7	
# ER Injury Visits for Female AC Pts 15-34	21		15			12			

# Visits w/ ER Hazardous								
Alcohol Screening A. # Visits w/Positive	12	57.1	0	0.0	+57.1	0	0.0	+57.1
Screen	10	47.6	0	0.0	+47.6	0	0.0	+47.6
# of ER Injury Visits for AC Pts 15-24	17		16			21		
<pre># Visits w/ ER Hazardous Alcohol Screening A. # Visits w/Positive</pre>	10	58.8	0	0.0	+58.8	0	0.0	+58.8
Screen	9	52.9	0	0.0	+52.9	0	0.0	+52.9
# ER Injury Visit for AC Pts 25-34	16		17			11		
<pre># Visits w/ ER Hazardous Alcohol Screening A. # Visits w/Positive</pre>		56.3	0	0.0	+56.3	0	0.0	+56.3
Screen	6	37.5	0	0.0	+37.5	0	0.0	+37.5

Figure 2-55: Sample Report, Alcohol Screening and Brief Intervention (ASBI) in the ER

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Alcohol Screening and Brief Intervention (ASBI) in the ER: List of visits for patients seen in the ER for an injury, with screening for hazardous alcohol use, results of screen and BNI, if any. PATIENT NAME COMMUNITY DENOMINATOR NUMERATOR -----PATIENT1, DARLENE S 000001 COMMUNITY #1 F 33 ER 1) 02/02/12 POV 816.02, SCREEN: Neg/No Res CPT 3016F UP,AC UP, AC
PATIENT2, RITA A 000002 COMMUNITY #1 F 33
FR 1) 07/12/12 POV 875.0 ER 1) 07/12/12 POV 875.0, SCREEN: Pos Ex 35, BNI: No PATIENT3, DIANE L 000003 COMMUNITY #1 F 15
UP.AC ER 1) 09/08/12 POV 815. ER 1) 09/08/12 POV 815.00, SCREEN: None PATIENT4, ALICIA 000004 COMMUNITY #1 F 18

IID ER 1) 04/20/12 POV 959.7 ER 1) 04/20/12 POV 959.7, SCREEN: Neg/No Res CPT H0049 PATIENT5, MELISSA 000005 COMMUNITY #1 F 16 UP,AC PATIENT6, LISA MARIE 000006 COMMUNITY #1 F 20 ER 1) 12/16/12 POV 873.42, SCREEN: None; ER 2) 12/18/12 POV 800.10, SCREEN: Pos Ex 35, BNI: 12/18/12 Yes ER AOD-BNI

Figure 2-56: Sample Patient List, Alcohol Screening and Brief Intervention (ASBI) in the ER

# 2.7.3 Intimate Partner (Domestic) Violence Screening

## **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 55.3% for the proportion of female patients ages 15 to 40 who receive screening for domestic violence.

### **Denominators**

Female Active Clinical patients ages 13 and older.

Female Active Clinical patients ages 15–40. (GPRA Denominator)

Female User Population patients ages 13 and older.

#### **Numerators**

Patients screened for intimate partner (domestic) violence at any time during the Report Period.

**Note**: This numerator does *not* include refusals. (GPRA Numerator)

- a. Patients with documented IPV/DV exam
- b. Patients with IPV/DV related diagnosis
- c. Patients provided with education or counseling about IPV/DV
- d. Patients with documented refusal in past year of an IPV/DV exam or IPV/DV-related education.

## **Logic Description**

Age of the patient is calculated at the beginning of the report period. CRS uses the following codes to define numerators.

	CPT Codes	ICD and Other Codes
IPV/DV Screening		V Exam: Code 34
		BHS Exam: IPV/DV
IPV/DV Diagnosis		V POV or current PCC or BHS Problem List: 995.80-995.83, 995.85, V15.41, V15.42, V15.49 BHS POV: 43.*, 44.*
IPV/DV Education		<b>Patient education codes:</b> "DV-" or "-DV", 995.80-83, 995.85, V15.41, V15.42, or V15.49
IPV/DV Counseling		<b>V POV:</b> V61.11
Refusals		V Exam: Code 34 BHS IPV/DV exam Patient education codes containing "DV-" or "-DV"

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of female patients 13 and older with documented IPV/DV screening or refusal, if any.

### **Measure Source**

HP 2010 15-34

# **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	55.3%
IHS FY 2010 Performance	53.0%
IHS FY 2009 Performance	48.0%
IHS FY 2008 Performance	42.0%
IHS FY 2007 Performance	36.0%
IHS FY 2006 Performance	28.0%
IHS FY 2005 Performance	13.0%
IHS FY 2004 Performance	4.0%

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Intimate Partner (Domestic) Violence Screening (con't)								
					CHG from E			HG from ASE %
# Female Active Clinical ages 13 and older	712		519			460		
# w/IPV/DV Screening								
-No Refusals	13	1.8	1	0.2	+1.6	0	0.0	+1.8
A. # w/documented IPV/DV exam	8	1.1	0	0.0	+1.1	0	0.0	+1.1
B. # w/ IPV/DV related diagnosis	4	0.6	0	0.0	+0.6	0	0.0	+0.6
C. # provided DV education	4	0.6	1	0.2	+0.4	0	0.0	+0.6

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# w/ IPV/DV refusal	3	0.4	0	0.0	+0.4	0	0.0	+0.4
# Female Active Clinical ages 15-40								
(GPRA)	380		311			267		
# w/IPV/DV screening -No Refusals								
(GPRA) A. # w/ documented	11	2.9	1	0.3	+2.6	0	0.0	+2.9
IPV/DV exam  B. # w/ IPV/DV related	6	1.6	0	0.0	+1.6	0	0.0	+1.6
diagnosis C. # provided DV	3	0.8	0	0.0	+0.8	0	0.0	+0.8
education # w/ IPV/DV	4	1.1	1	0.3	+0.7	0	0.0	+1.1
refusal	2	0.5	0	0.0	+0.5	0	0.0	+0.5
# Female User Pop 13 and older	010		1 005			917		
13 and order	1,218		1,005			917		
# w/IPV/DV Screening								
-No Refusals A. # w/ documented	13	1.1	1	0.1	+1.0	1	0.1	+1.0
IPV/DV exam	8	0.7	0	0.0	+0.7	0	0.0	+0.7
B. # w/ IPV/DV related diagnosis	4	0.3	0	0.0	+0.3	1	0.1	+0.2
C. # provided DV education	4	0.3	1	0.1	+0.2	0	0.0	+0.3
# w/ IPV/DV refusal	3	0.2	0	0.0	+0.2	0	0.0	+0.2

Figure 2-57: Sample Report, Intimate Partner (Domestic) Violence Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Intimate Partner (Domestic) Violence Screening: List of female patients 13 and older with documented IPV/DV screening or refusal, if any. HRN COMMUNITY SEX AGE PATIENT NAME DENOMINATOR NUMERATOR PATIENT1, ELVIRA 000001 COMMUNITY #1 F 13
UP 03/18/12 Refused Ex 3
PATIENT2, SHARON KAY 000002 COMMUNITY #1 F 14 03/18/12 Refused Ex 34 PATIENT3, KRISTINA 000003 COMMUNITY #1 F 15 PATIENT4, RITA 000004 COMMUNITY #1 F 15 EXAM: 05/06/12 Ex 34 UP,AC PATIENT5, DIANE LOUISE 000005 COMMUNITY #1 F 15 EXAM: 02/24/12 Ex 34 PATIENT6, ALICE LILA 000006 COMMUNITY #1 F 15 UP,AC

Figure 2-58: Sample Patient List, Intimate Partner (Domestic) Violence Screening

# 2.7.4 Depression Screening

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 56.5% for the proportion of adults ages 18 and older who receive annual screening for depression.

#### **Denominators**

Active Clinical patients ages 18 and older. Broken down by gender. (GPRA Denominator)

a. Active Clinical patients ages 65 and older. Broken down by gender.

User Population patients ages 18 and older. Broken down by gender.

b. User Population patients ages 65 and older. Broken down by gender.

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes prior to the Report Period, and at least two visits during the Report Period, and two DM-related visits ever. Broken down by gender.

Active IHD patients, defined as all Active Clinical patients diagnosed with ischemic heart disease (IHD) prior to the Report Period, and at least two visits during the Report Period, and two IHD-related visits ever. Broken down by gender.

#### **Numerators**

Patients screened for depression or diagnosed with a mood disorder at any time during the Report Period.

**Note**: This numerator does not include refusals. (GPRA Numerator)

- a. Patients screened for depression during the Report Period.
- b. Patients with a diagnosis of a mood disorder during the Report Period.
- c. Patients with documented depression screening refusal in past year.
- d. Patients with depression-related education or refusal of education in past year.

**Note:** Depression-related patient education does not count toward the GPRA numerator and is included as a separate numerator only.

#### **Logic Description**

Age is calculated at beginning of the Report Period.

CRS uses the following codes and taxonomies to define the denominator and numerators.

	ICD and Other Codes					
Diabetes	V POV: 250.00–250.93					
Ischemic Heart Disease	V POV: 410.0–412.*, 414.0–414.9, 429.2					
Depression Screening	V Exam: Exam Code 36					
	<b>V POV:</b> V79.0					
	<b>CPT:</b> 1220F					
	BHS Problem Code: 14.1 (Screening for Depression)					
	V Measurement in PCC or BHS: PHQ2 or PHQ9					
Mood Disorders	At least 2 visits in PCC or BHS for: Major Depressive Disorder, Dysthymic Disorder, Depressive Disorder NOS, Bipolar I or II Disorder, Cyclothymic Disorder, Bipolar Disorder NOS, Mood Disorder Due to a General Medical Condition, Substance-induced Mood Disorder, or Mood Disorder NOS.  V POV: 296.*, 291.89, 292.84, 293.83, 300.4, 301.13, or 311  BHS POV: 14, 15					
Depression-related Patient Education (does not count	Documented education of any of the following during the Report Period:					
toward GPRA numerator)	Patient education codes: containing "DEP-" (depression), 296.2* or 296.3*, "BH-" (behavioral and social health), 290319, 995.5*, or 995.80–995.85, "SB-" (suicidal behavior) or 300.9, or "PDEP-" (postpartum depression) or 648.44.					
Screening Refusals	V Exam: Exam Code 36, in past year					
Refusal of Depression- related Patient Education (does not count toward GPRA numerator)	Documented refusal of any of the following during the Report Period:  Patient education codes: containing "DEP-" (depression), "BH-" (behavioral and social health), "SB-" (suicidal behavior), or "PDEP-" (postpartum depression).					

## **Recommended Brief Screening Tool**

A sample of a Patient Health Questionnaire (PHQ-2 Scaled Version) appears below.

Over the past two weeks, how often have you been bothered by any of the following problems?

- 1. Little interest or pleasure in doing things
  - Not at all Value: 0
  - Several days Value: 1
  - More than half the days Value: 2
  - Nearly every day Value: 3
- 2. Feeling down, depressed, or hopeless

• Not at all Value: 0

• Several days Value: 1

More than half the days Value: 2

• Nearly every day Value: 3

Total Possible PHQ-2 Score: Range: 0-6

0–2: Negative

3-6: Positive; further evaluation indicated

Provider should note the screening tool used was the PHQ-2 Scaled in the COMMENT section of the Exam Code.

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients with documented depression screening or refusal/diagnosed with mood disorder, if any.

### **Measure Source**

USPSTF (US Preventive Services Task Force), HP 2010 developmental indicator 18–6

## **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	56.5%
IHS FY 2010 Performance	52.0%
IHS FY 2009 Performance	44.0%
IHS FY 2008 Performance	35.0%
IHS FY 2007 Performance	24.0%
IHS FY 2006 Performance	15.0%

```
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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000

Depression Screening (con't)

REPORT % PREV YR % CHG from BASE % CHG from
```

PER	IOD		PERIOD		PREV YR % I	PERIOD	В	BASE %
2								
Active Clinical Pts =>18 (GPRA) 1,	112		793			666		
# w/Depression screening or Mood Disorder DX-No								
Refusals (GPRA) A. # screened for	81	7.3	42	5.3	+2.0	17	2.6	+4.7
depression B. # w/mood disorder	39	3.5	0	0.0	+3.5	0	0.0	+3.5
DX # w/depression	43	3.9	42	5.3	-1.4	17	2.6	+1.3
<pre>screening refusal # w/depression education</pre>	5	0.4	0	0.0	+0.4	0	0.0	+0.4
or refusal	11	1.0	3	0.4	+0.6	0	0.0	+1.0
Male Active Clinical								
Pts >=18	448		308			250		
# w/ Depression screenin or Mood Disorder DX-No	g							
Refusals A. # screened for	25	5.6	7	2.3	+3.3	1	0.4	+5.2
depression B. # w/Mood Disorder	14	3.1	0	0.0	+3.1	0	0.0	+3.1
DX # with depression	11	2.5	7	2.3	+0.2	1	0.4	+2.1
screening refusal # w/depression education	3	0.7	0	0.0	+0.7	0	0.0	+0.7
or refusal	3	0.7	1	0.3	+0.3	0	0.0	+0.7
Female Active Clinical Pts >=18	664		485			416		
# w/ Depression screenin	g							
or Mood Disorder DX-No Refusals	56	8.4	35	7.2	+1.2	16	3.8	+4.6
A. # screened for depression	25	3.8	0	0.0	+3.8	0	0.0	+3.8
B. # w/Mood Disorder DX	32	4.8	35	7.2	-2.4	16	3.8	+1.0
# with depression screening refusal	2	0.3	0	0.0	+0.3	0	0.0	+0.3
<pre># w/depression education   or refusal</pre>	8	1.2	2	0.4	+0.8	0	0.0	+1.2
A. Active Clinical Pts	116		64			65		
# w/ Depression screenin								
or Mood Disorder DX-No Refusals	9	7.8	6	9.4	-1.6	2	3.1	+4.7
A. # screened for depression	3	2.6	0	0.0	+2.6	0	0.0	+2.6
B. # w/mood disorder DX	6	5.2	6	9.4	-4.2	2	3.1	+2.1
# with depression	0	J. Z	0	2.4	-1.2	۷	J•1	12.1

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screening refusal # w/depression education	1	0.9	0	0.0	+0.9	0	0.0	+0.9
or refusal	2	1.7	2	3.1	-1.4	0	0.0	+1.7

Figure 2-59: Sample Report, Depression Screening

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Depression Screening: List of patients with documented depression
screening or refusal/diagnosed with mood disorder, if any.
                        HRN COMMUNITY SEX AGE
PATIENT NAME
 DENOMINATOR
                            NUMERATOR
PATIENT55, LORETTA LYNN 000055 COMMUNITY #1 F 78
PATIENT56, TINA MARIE 000056 COMMUNITY #1 F 78
UP,AC,AD,IHD SCREEN: 05/22/12 Meas
PATIENT57,DANIELLE 000057 COMMUNITY #1 F 79
UP,AC PATIENT58,LESLIE ANN 000058 COMMUNITY #1 F 80
UP,AC SCREEN: 04/15/12 POV V
PATIENT59,DONNA SUE 000059 COMMUNITY #1 F 86
UP,AC SCREEN: 01/15/12 POV V
                                   SCREEN: 05/22/12 Meas PHQ9
                                   PT ED: 02/06/12 296.20-DP
                                   SCREEN: 04/15/12 POV V79.0
                                   SCREEN: 01/15/12 POV V79.0
PATIENT60, TAYLOR OLIVIA 000060 COMMUNITY #1 F 87
PATIENT61, DENNIS GERALD 000061 COMMUNITY #1 M 18
                      PT ED: 02/01/12 296.20-DP
PATIENT62, JOSHUA DALE 000062 COMMUNITY #1 M 18
UP,AC
```

Figure 2-60: Sample Patient List, Depression Screening

# 2.7.5 Antidepressant Medication Management

#### **Denominators**

As of the 120th day of the Report Period, *Active Clinical* patients 18 years of age and older who were diagnosed with a new episode of depression and treated with antidepressant medication in the past year.

As of the 120th day of the Report Period, *User Population* patients 18 years of age and older who were diagnosed with a new episode of depression and treated with antidepressant medication in the past year.

#### **Numerators**

Optimal Practitioner Contacts: Patients with at least three mental health visits with a non-mental health or mental health provider within 12 weeks (84 days) after diagnosis, two of which must be face-to-face visits and one of which must be with a prescribing provider.

Effective Acute Phase Treatment: Patients who filled a sufficient number of separate prescriptions/refills of antidepressant medication for continuous treatment of at least 84 days (12 weeks).

Effective Continuation Phase Treatment: Patients who filled a sufficient number of separate prescriptions/refills of antidepressant medication treatment to provide continuous treatment for at least 180 days (6 months).

## **Logic Description**

Age is calculated at the beginning of the report period. To be included in the denominator, patient must meet *both* of the following conditions:

- 1. One of the following from the 121st day of the year prior to the Report Period to the 120th day of the Report Period:
  - a. One visit in any setting with major depression DX (see list of codes below) as primary POV.
  - b. Two outpatient visits occurring on different dates of service with secondary POV of major depression.
  - c. An inpatient visit with secondary POV of major depression.

For example, if Report Period is July 1, 2010–June 30, 2011, the patient must have one of the three scenarios above during 11/1/2009–10/29/2010.

Major depression is defined as POV 296.2\*, 296.3\*, 298.0, 300.4, 309.1, 311. The Index Episode Start Date is date of the patient's earliest visit during this period. For inpatient visits, the discharge date will be used.

2. Filled a prescription for an antidepressant medication (see list of medications below) within 30 days before the Index Episode Start Date or 14 days on or after that date. In V Medication, Date Discontinued must not be equal to the prescription (i.e., visit) date. The Index Prescription Date is the date of the earliest prescription for antidepressant medication filled during that time period.

#### **Denominator Exclusions**

- 1. Patients who have had any diagnosis of depression within the previous 120 days (four months) of the Index Episode Start Date. The POVs to be checked for prior depressive episodes is more comprehensive and include the following: POV 296.2\*-296.9\*, 298.0, 300.4, 309.0, 309.1, 309.28, 311, or
- 2. Patients who had a new or refill prescription for antidepressant medication (see the list of medications below) within 90 days (3 months) prior to the Index Prescription Date are excluded as they do not represent new treatment episodes, or

3. Patients who had an acute mental health or substance abuse inpatient stay during the 245 days after the Index Episode Start Date treatment period. Acute mental health stays are defined as Service Category of H and primary POV 290\*, 293\*–302\*, 306\*–316\*. Substance abuse inpatient stays are defined as Service Category of H and primary POV 291\*–292\*, 303\*–305\* or primary POV 960\*–979\* and secondary POV of 291\*–292\*, 303\*–305\*.

Optimal Practitioner Contacts Numerator

Patient must have one of the following:

- 1. Three face-to-face follow-up outpatient, non-ER visits (clinic code not equal to 30) or intermediate treatment with either a non-mental-health or mental health provider within 84 days after the Index Episode Start Date, *or*...
- 2. Two face-to-face outpatient, non-ER visits (clinic code not equal to 30) and one telephone visit (Service Category T), with either a non-mental-health or mental health provider within 84 days after the Index Episode Start Date.

For either option, one of the visits must be to a prescribing provider, defined as Provider Codes 00, 08, 11, 16-18, 21, 24-25, 30, 33, 41, 44-45, 47, 49, 64, 67-68, 70-83, 85-86, A1, A9, or B1-B6.

**Note**: If patient was diagnosed with two secondary diagnoses of depression, the second visit may be counted toward the numerator.

**Outpatient mental health provider visits** are defined as BHS or PCC visit with primary provider code of 06, 12, 19, 48, 49, 50, 62, 63, 81, or 92–96, *and*:

- a. (1) Service category A, S, or O, and (2A) CPT 90801, 90802, 90804–90819, 90821–90824, 90826–90829, 90845, 90847, 90849, 90853, 90857, 90862, 90870, 90871 (old code), 90875, 90876, 99384–99387, 99394–99397, 99401–99404, G0155, G0176, G0177, H0002, H0004, H0031, H0034, H0036, H0037, H0039, H0040, H2000, H2010, H2011, H2013–H2020, M0064, S9484, S9485 or (2B) POV 290\*, 293\*–302\*, 306\*–316, or
- b. (1) Service category of A, S, or O and (2A) Location of Encounter = Home (as designated in Site Parameters) or (2B) Clinic Code = 11, or
- c. Service category of T.

**Outpatient non-mental-health provider visits** are defined as BHS or PCC visits with:

- a. (1) Service category A, S, or O, a (2) CPT 90801, 90802, 90804–90819, 90821–90824, 90826–90829, 90845, 90847, 90849, 90853, 90857, 90862, 90870, 90871 (old code), 90875, 90876, G0155, G0176, G0177, H0002, H0004, H0031, H0034, H0036, H0037, H0039, H0040, H2000, H2010, H2011, H2013–H2020, M0064, S9484, S9485, or...
- b. (1) Service category A, S, O, or T or (2) Location of Encounter = Home (as designated in Site Parameters) or (3A) clinic code 11 and (3B) POV 290\*, 293\*–302\*, 306\*–316\*, or...
- c. (1) Service category A, S, or O, and (2) CPT 99384–99387, 99394–99397, 99401–99404 and (3) POV 290\*, 293\*–302\*, 306\*–316\*.

### Effective Acute Phase Treatment Numerator

For all antidepressant medication prescriptions filled (see list of medications below) within 114 days of the Index Prescription Date, from V Medication CRS counts the days prescribed (i.e., treatment days) from the Index Prescription Date until a total of 84 treatment days has been established. If the patient had a total gap exceeding 30 days or if the patient does not have 84 treatment days within the 114-day time frame, the patient is not included in the numerator.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2010, Discontinued Date=11/19/2010, Recalculated # Days Prescribed=4.

### **Example of Patient Included in Numerator:**

- First RX is Index Rx Date: 11/1/2010, # Days Prescribed=30
- Rx covers patient through 12/1/2010
- Second RX: 12/15/2010, # Days Prescribed=30
- Gap #1 = (12/15/2010-12/1/2010) = 14 days
- Rx covers patient through 1/14/2011
- Third RX: 1/10/2011, # Days Prescribed=30
- No gap days
- Rx covers patient through 2/13/2011
- Index Rx Date  $\frac{11}{12010} + \frac{114}{400} = \frac{2}{23}$
- Patient's 84th treatment day occurs on 2/7/2011, which is <= 2/23/2011 and # gap days of 14 is less than 30

### **Example of Patient Not Included in Numerator:**

- First Rx is Index Rx Date: 11/1/2010, # Days Prescribed=30
- Rx covers patient through 12/1/2010
- Second Rx: 12/15/2010, # Days Prescribed=30
- Gap #1 = (12/15/2010-12/1/2010) = 14 days
- Rx covers patient through 1/14/2011
- Third Rx: 2/01/2011, # Days Prescribed=30
- Gap #2 = (2/01/2011-1/14/2011) = 18, total # gap days = 32, so patient is not included in the numerator

Effective Continuation Phase Treatment Numerator

For all antidepressant medication prescriptions (see list of medications below) filled within 231 days of the Index Prescription Date, CRS counts the days prescribed (i.e., treatment days) (from V Medication) from the Index Prescription Date until a total of 180 treatment days has been established. If the patient had a total gap exceeding 51 days or if the patient does not have 180 treatment days within the 231 day time frame, the patient is not included in the numerator.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2010, Discontinued Date=11/19/2010, Recalculated # Days Prescribed=4.

Antidepressant medications defined with medication taxonomy BGP HEDIS ANTIDEPRESSANT MEDS. (Medications are: Tricyclic antidepressants (TCA) and other cyclic antidepressants, Selective serotonin reuptake inhibitors (SSRI), Monoamine oxidase inhibitors (MAOI), Serotonin-norepinepherine reuptake inhibitors (SNRI), and other antidepressants.) Medications must not have a comment of RETURNED TO STOCK.

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients with new depression DX and optimal practitioner contact (OPC), acute phase treatment (APT) and continuation phase treatment (CONPT), if any.

#### **Measure Source**

HEDIS, HP 2010 18-9b

## Measure Past Performance and Long-Term Targets

None

DU November 25, 2012 Page 141  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000										
Antidepressant Medication	Antidepressant Medication Management (con't)									
REPORT % PREV YR % CHG from BASE % CHG from PERIOD PREV YR % PERIOD BASE %										
Active Clinical Pts =>1 depression DX and antidepressant meds	8 w/n 17	ew	6			2				
<pre># w/3 outpt mental health visits within 12 weeks</pre>	5	29.4	1	16.7	+12.7	0	0.0	+29.4		
<pre># w/12 week treatment   meds # w/180 day treatment</pre>	9	52.9	4	66.7	-13.7	0	0.0	+52.9		
meds	4	23.5	3	50.0	-26.5	0	0.0	+23.5		
User Pop Pts =>18 w/new depression DX and antidepressant meds	18		7			3				
<pre># w/3 outpt mental health visits within 12 weeks # w/12 week treatment</pre>	5	27.8	1	14.3	+13.5	0	0.0	+27.8		
meds	9	50.0	4	57.1	-7.1	0	0.0	+50.0		
# w/180 day treatment meds	4	22.2	3	42.9	-20.6	0	0.0	+22.2		

Figure 2-61: Sample Report, Antidepressant Medication Management

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Antidepressant Medication Management: List of patients with new depression DX and optimal practitioner contact (OPC), acute phase treatment (APT) and continuation phase treatment (CONPT), if any.

PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR

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```
PATIENT1, MICHELLE D 000001 COMMUNITY #1 F 22

UP, AC IESD: 06/06/11; NOT OPC; NOT APT: DAYS=60, GAP=1; NOT

CONPT: DAYS=60, GAP=1

PATIENT2, PAULA KAY 000002 COMMUNITY #1 F 34

UP, AC IESD: 10/29/11; NOT OPC; NOT APT: DAYS=68, GAP=28;

CONPT

PATIENT3, RHONDA SUE 000003 COMMUNITY #1 F 35

UP IESD: 04/21/12; NOT OPC; NOT APT: DAYS=74, GAP=0; NOT

CONPT: DAYS=74, GAP=0

PATIENT4, KATHLEEN 000004 COMMUNITY #1 F 38

UP, AC IESD: 11/15/11; NOT OPC; APT; CONPT
```

Figure 2-62: Sample Patient List, Antidepressant Medication Management

# 2.8 Cardiovascular Disease Related Measure Topics

# 2.8.1 Obesity Assessment

#### **Denominators**

*Active Clinical patients* ages 2 through 74. Broken down by gender and age groups (2-5, 6-11, 12-19, 20-24, 25-34, 35-44, 45-54, 55-74).

All User Population patients ages 2 through 74. Broken down by gender.

#### **Numerators**

Patients for whom a Body Mass Index (BMI) could be calculated.

**Note**: This numerator does *not* include refusals.

- 1. For those with a BMI calculated, those considered overweight but not obese using BMI and standard tables
- 2. For those with a BMI calculated, those considered obese using BMI and standard tables
- 3. Total of overweight and obese

Patients with documented refusal in past year.

### **Logic Description**

Age is calculated at beginning of the Report Period.

BMI calculation definition: CRS calculates BMI at the time the report is run, using NHANES II. For age 18 and under, a height and weight must be taken on the same day any time during the Report Period. For 19 through 50, height and weight must be recorded within last five years, not required to be on the same day. For over 50, height and weight within last two years, not required to be recorded on same day. Overweight but not obese is defined as BMI of 25 through 29 for adults 19 and older. Obese is defined as BMI of 30 or more for adults 19 and older. For ages 2-18, definitions are based on standard tables. Refusals include REF, NMI, and UAS (unable to screen) and must be documented during the past year. For ages 18 and under, both the height and weight must be refused on the same visit at any time during the past year. For ages 19 and older, the height and weight must be refused during the past year and are not required to be on the same visit.

Patients whose BMI either is greater or less than the Data Check Limit range shown in the BMI Standard Reference Data Table in PCC will not be included in the report counts for Overweight or Obese.

# **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients with current BMI, if any.

#### **Measure Source**

HP 2020: NWS-9 Obesity in Adults 20+, NWS-10.1 (Obesity in Children 2-5), NWS-10.2 Overweight or Obesity in Children 6-11, NWS-10.3 Overweight or Obesity in Adolescents 12-19, NWS-10.4 Overweight or Obesity in Children 2-19

### **Measure Past Performance and Long-Term Targets**

Performance	Percent
Assessed as Obese–IHS FY 2011 Performance	46.9%
Assessed as Obese–IHS FY 2010 Performance	47.0%
Assessed as Obese–IHS FY 2009 Performance	47.0%
Assessed as Obese–IHS FY 2008 Performance	46.0%
BMI Measured–IHS FY 2011 Performance	78.0%
BMI Measured–IHS FY 2010 Performance	76.0%
BMI Measured–IHS FY 2009 Performance	75.0%
BMI Measured–IHS FY 2008 Performance	74.0%
BMI Measured– FY 2005 Performance	64.0%
BMI Measured–IHS FY 2004 Performance	60.0%
HP 2020 Goal: Obesity in Adults 20+ (NWS-9)	30.6%

Performance	Percent
HP 2020 Goal: Overweight or Obesity in Children 2–5 (NWS-10.1)	9.6%
HP 2020 Goal: Overweight or Obesity in Children 6–11 (NWS-10.2)	15.7%
HP 2020 Goal: Overweight or Obesity in Adolescents 12–19 (NWS-10.3)	16.1%
HP 2020 Goal: Overweight or Obesity in Children 2–19 (NWS-10.4)	14.6%

### **Performance Improvement Tips**

- 1. A Body Mass Index report can be run from your PCC Management Reports menu. This report can be run for all patients or for a specific template of patients that has been pre-defined with a QMan search. The BMI report will provide you with patient height, weight, date weight taken, BMI and NHANES percentile.
- 2. Recent guidelines indicate that height for adults must be taken at least once every five years, rather than once after age 18. Your BMI rates may be lower than anticipated because of height data that is over five years old.
- 3. If height and weight measurements are being recorded as cm/kg vs. in/lbs ensure providers are *noting* they are cm/kg *and* that data entry is entering the measurements correctly in PCC, as shown below.
  - Use mnemonics of CHT and KWT (vs. HT and WT), or
  - Add "c" after height value and "k" after weight value (e.g. 100c, 50k)

Previo	port Per us Year	Measu DEMC iod: J Period	) INDIAN H Jan 01, 20 l: Jan 01	Commun OSPITA 12 to , 2011	nity Specif	12	_	ge 143
Obesity Assessment (	con't)							
					CHG from PREV YR %			
Active Clinical Pts ages 2-74	1,400		1,096			982		
<pre># w/BMI calculated   -No Refusals A. # Overweight w/</pre>	880	62.9	824	75.2	-12.3	712	72.5	-9.6
% of Total BMI B. # Obese w/	242	27.5	237	28.8	-1.3	191	26.8	+0.7
% of Total BMI		42.3	339	41.1	+1.1	267	37.5	+4.8
C. # Overweight/Obes % of Total BMI		69.8	576	69.9	-0.1	458	64.3	+5.4
# w/BMI refusal (No BMI)	4	0.3	0	0.0	+0.3	0	0.0	+0.3

Male Active Clinical									
Pts 2-74	595		465			410			
# w/BMI calculated									
-No Refusals	341	57.3	331	71.2	-13.9	283	69.0	-11.7	
A. # Overweight w/ % of Total BMI	103	30.2	9.8	29.6	+0.6	73	25.8	+4.4	
B. # Obese w/	103	30.2	70	20.0	10.0	75	23.0	14.4	
% of Total BMI C. #Overweight/Obese w/	155	45.5	141	42.6	+2.9	117	41.3	+4.1	
% of Total BMI	258	75.7	239	72.2	+3.5	190	67.1	+8.5	
<pre># w/BMI refusal   (no BMI)</pre>	2	0.3	0	0.0	+0.3	0	0.0	+0.3	
· · · · · · · · · · · · · · · · · · ·									
Female Active Clinical Pts 2-74	805		631			572			
# w/BMI calculated -No Refusals	539	67.0	493	78.1	-11.2	429	75.0	-8.0	
A. # Overweight w/									
% of Total BMI B. # Obese w/	139	25.8	139	28.2	-2.4	118	27.5	-1.7	
% of Total BMI	217	40.3	198	40.2	+0.1	150	35.0	+5.3	
<pre>C. #Overweight/Obese w/ % of Total BMI</pre>	356	66.0	337	68.4	-2.3	268	62.5	+3.6	
# w/BMI refusal	330	00.0	331	00.4	-2.3	200	04.3	+3.0	
(No BMI)	2	0.2	0	0.0	+0.2	0	0.0	+0.2	

Figure 2-63: Sample Report, Obesity Assessment

Obesity Assessment (con	ı't)								
	TOTAL	ACTIVE							
	2-5	6-11	_	Distri 20-24		35-44	45-54	55-74	
CURRENT REPORT PERIOD									
Total # Active Clin # w/BMI calculated	109	112	157	133	233	215	216	225	
-No Refusals	52	44	89	117	183	146	128	121	
% w/BMI calculated -No Refusals	47.7	39.3	56.7	88.0	78.5	67.9	59.3	53.8	
# A. Overweight	9	10	21	33	44	39	38	48	
<pre>% A. Overweight w/ % Total BMI</pre>	17.3	22.7	23.6	28.2	24.0	26.7	29.7	39.7	
# B. Obese	7	13	28	39	86	86	61	52	
% B. Obese w/ % of Total BMI	13.5	29.5	31.5	33.3	47.0	58.9	47.7	43.0	
# C. Overweight									
or Obese % C. Overweight or Obes	16 se w/	23	49	72	130	125	99	100	
% Total BMI	30.8	52.3	55.1	61.5	71.0	85.6	77.3	82.6	
# w/BMI refusal									

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(No BMI) % w/BMI refusal	1	0	0	0	0	1	1	1	
(No BMI)	1.9	0.0	0.0	0.0	0.0	0.7	0.8	0.8	
PREVIOUS YEAR PERIOD Total # Active Clin	111	120	137	128	172	151	140	137	
# w/BMI calculated	111	120	137	120	1/2	131	140	137	
-No Refusals % w/BMI calculated	49	56	88	114	155	129	113	120	
-No Refusals	44.1	46.7	64.2	89.1	90.1	85.4	80.7	87.6	
# A. Overweight % A. Overweight w/	7	11	20	38	47	33	36	45	
% Total BMI	14.3	19.6	22.7	33.3	30.3	25.6	31.9	37.5	
# B. Obese % B. Obese w/	14	14	26	35	65	77	56	52	
% of Total BMI	28.6	25.0	29.5	30.7	41.9	59.7	49.6	43.3	
# C. Overweight									
or Obese	21	25	46	73	112	110	92	97	
% C. Overweight or Obe									
% Total BMI	42.9	44.6	52.3	64.0	72.3	85.3	81.4	80.8	
# w/BMI refusal									
(No BMI) % w/BMI refusal	0	0	0	0	0	0	0	0	
(No BMI)	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
CHANGE FROM PREV YR %									
w/BMI calculated									
-No Refusals	+3.6	-7.4	-7.5	-1.1	-11.6	-17.5	-21.5	-33.8	
A. Overweight	+3.0	+3.1	+0.9	-5.1	-6.3	+1.1	-2.2	+2.2	
B. Obese	-15.1	+4.5	+1.9	+2.6	+5.1	-0.8	-1.9	-0.4	
C. Overweight									
or Obese w/BMI refusal	-12.1	+7.6	+2.8	-2.5	-1.2	+0.3	-4.1	+1.8	
(No BMI)	+1.9	+0.0	+0.0	+0.0	+0.0	+0.7	+0.8	+0.8	

Figure 2-64: Sample Report, Age Breakout, Obesity Assessment

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Obesity Assessment: List of patients with current BMI, if any. PATIENT NAME HRN DENOMINATOR COMMUNITY SEX AGE PATIENT NAME NUMERATOR -----PATIENT1, PAMELA 000001 COMMUNITY #1 F 3 UP,AC 16.03 PATIENT2, GLENDA 000002 COMMUNITY #1 F 3
UP, AC 17.49 PATIENT3, SHIRLEY 000003 COMMUNITY #1 F 5 PATIENT4, MARY ANNE 000004 COMMUNITY #1 F 5
UP, AC Refused
PATIENT5, JACKIE 000005 COMMUNITY #1 F 9
UP PATIENT6,ZINNIA 000006 COMMUNITY #1 F 15
UP 29.41 [OVERWEIGHT]
PATIENT7,MARY RYAN 000007 COMMUNITY #1 F 15
UP,AC 33.69 [OBESE]

Figure 2-65: Sample Patient List, Obesity Assessment

# 2.8.2 Childhood Weight Control

### **GPRA Description**

During FY 2012, achieve the tentative long-term target rate of 24% for the proportion of children with a BMI of 95% or higher.

#### **Denominators**

Active Clinical patients aged 2–5 for whom a BMI could be calculated. Broken down by gender and age groups (2, 3, 4, 5).

#### **Numerators**

Patients with BMI in the 85th to 94th percentile.

Patients with a BMI at or above the 95th percentile.

Patients with a BMI at or above the 85th percentile.

### **Logic Description**

BMI calculation definition: All patients for whom a BMI could be calculated and who are between the ages of two and five at the beginning of the Report Period and who do not turn age six during the Report Period are included in this measure. Age in the age groups is calculated based on the date of the most current BMI found. For example, a patient may be two years of age at the beginning of the time period, but is three years old at the time of the most current BMI found. That patient will fall into the age three group. CRS looks for the most recent BMI in the Report Period. CRS calculates BMI at the time the report is run using NHANES II. A height and weight must be taken on the same day any time during the Report Period. The BMI values for this measure are reported differently than in Obesity Assessment since this age group is children ages 2-5, whose BMI values are age-dependent. The BMI values are categorized as Overweight for patients with a BMI in the 85th to 94th percentile and Obese for patients with a BMI at or above the 95th percentile.

Patients whose BMI either is greater or less than the Data Check Limit range shown below will not be included in the report counts for Overweight or Obese.

#### **BMI Standard Reference Data**

Low-High Ages	Sex	BMI >= (OVERWT)	BMI >= (OBESE)		Data Check Limits BMI <
2-2	MALE	17.7	18.7	36.8	7.2
2-2	FEMALE	17.5	18.6	37.0	7.1
3-3	MALE	17.1	18.0	35.6	7.1
3-3	FEMALE	17.0	18.1	35.4	6.8
4-4	MALE	16.8	17.8	36.2	7.0
4-4	FEMALE	16.7	18.1	36.0	6.9
5-5	MALE	16.9	18.1	36.0	6.9
<b>5-</b> 5	FEMALE	16.9	18.5	39.2	6.8

### **Key Logic Changes from CRS Version 11.1**

None

#### Patient List Description

List of patients ages 2–5, with current BMI.

#### **Measure Source**

CDC, National Center for Health Statistics, HP 2020 NWS-10.1

# **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	24.1%
IHS FY 2010 Performance	25.0%
IHS FY 2009 Performance	25.0%
IHS FY 2008 Performance	24.0%
IHS FY 2007 Performance	24.0%
IHS FY 2006 Performance	24.0%
HP 2020 Goal	9.6%

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DEMO INDIAN HOSPITAL										
Report Period: Jan 01, 2012 to Dec 31, 2012 Previous Year Period: Jan 01, 2011 to Dec 31, 2011										
					to Dec 31,					
Childhood Weight Cor	ntrol (co	n't)								
	REPORT PERIOD		PREV YR PERIOD		CHG from PREV YR %					
Active Clinical Pts										
2-5 w/BMI	44		39			40				
# w/BMI 85-94%	7	15.9	5	12.8	+3.1 -11.7	10	25.0			
# w/BMI =>95%								-1.1		
# w/BMI =>85%	12	27.3	14	35.9	-8.6	15	37.5	-10.2		
Active Clinical Pts										
Age 2	2		8			5				
# w/BMI 85-94%	1	50.0	0	0.0	+50.0	1	20.0	+30.0		
# w/BMI =>95%	0	0.0			-25.0					
# w/BMI =>85%	1	50.0	2	25.0	+25.0	1	20.0	+30.0		
Active Clinical Pts										
Age 3	23		15			8				
# w/BMI 85-94%	2	8.7	2	13.3	-4.6	3	37.5	-28.8		
# w/BMI =>95%	3	13.0	3	20.0	-7.0	2	25.0	-12.0		
# w/BMI =>85%	5	21.7	5	33.3	-11.6	5	62.5	-40.8		
Active Clinical Pts										
Age 4	12		10			17				
# w/BMI 85-94%	1	8.3	2	20.0	-11.7	3	17.6	-9.3		
# w/BMI =>95%		8.3	2	20.0	-11.7	2	11.8	-3.4		
# w/BMI =>85%	2	16.7	4	40.0	-23.3	5	29.4	-12.7		
Active Clinical Pts										
Age 5	7		6			10				

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# w/BMI 85-94%	3 42	.9 1	16.7	+26.2	3	30.0	+12.9
# w/BMI =>95%	1 14	.3 2	33.3	-19.0	1	10.0	+4.3
# w/BMI =>85%	4 57	.1 3	50.0	+7.1	4	40.0	+17.1

Figure 2-66: Sample Report, Childhood Weight Control

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Childhood Weight Control: List of patients ages 2-5, with current BMI.
                         HRN COMMUNITY
PATIENT NAME
DENOMINATOR
                                                     SEX AGE
                              NUMERATOR
PATIENT1, MELISSA ANN 000001 COMMUNITY #1 F 4
AC Age at BMI: 4; 08/20,
PATIENT2,RANDY 000002 COMMUNITY #1 M 2
AC Age at BMI: 2; 05/06,
PATIENT3,PAUL BARRY 000003 COMMUNITY #1 M 2
AC Age at BMI: 2; 08/05,
PATIENT4,TYLER 000004 COMMUNITY #1 M 4
                                   Age at BMI: 4; 08/20/12 16.03
                                   Age at BMI: 2; 05/06/12 17.96 [OVERWEIGHT]
                                   Age at BMI: 2; 08/05/12 19.87 [OBESE]
                                    Age at BMI: 4; 02/19/12 15.67
AC Age at BMI: 4; 02/19 PATIENT5, SAMUEL III 000005 COMMUNITY #1 M 5
                                    Age at BMI: 5; 11/29/12 19.07 [OBESE]
AC Age at BMI: 5; 11/29 PATIENT21, JOSEPHINE 000021 COMMUNITY #2 F 4
                                  Age at BMI: 4; 05/30/12 15.71
```

Figure 2-67: Sample Patient List, Childhood Weight Control

### 2.8.3 Nutrition and Exercise Education for At Risk Patients

#### **Denominators**

Active Clinical patients ages six and older considered overweight (including obese). Broken down by gender.

a. Active Clinical patients ages 6 and older *considered obese*. Broken down by gender and age groups (6-11, 12-19, 20-39, 40-59, 60+).

Active Diabetic patients, defined as all Active Clinical patients diagnosed with diabetes at least one year prior to the end of the Report Period, and at least two visits in the past year, and two diabetes-related visits ever.

#### **Numerators**

Patients provided with medical nutrition therapy during the Report Period.

Patients provided specific nutrition education during the Report Period.

Patients provided specific exercise education during the Report Period.

Patients provided with other related exercise and nutrition (lifestyle) education.

# **Logic Description**

Age of the patient is calculated at beginning of Report Period.

*Diabetes:* First DM Purpose of Visit 250.00–250.93 recorded in the V POV file prior to the Report period.

*Overweight*: Ages 19 and older, BMI equal to or greater than (=>) 25. Overweight is defined as including both obese and overweight categories calculated by BMI.

Obese: Ages 19 and older, BMI equal to or greater than (=>) 30. For ages 18 and under, the definition is based on standard tables. CRS calculates BMI at the time the report is run, using NHANES II. For 18 and under, a height and weight must be taken on the same day any time in the year prior to the end of the Report Period. For 19 through 50, height and weight must be recorded within last five years, not required to be on the same day. For over 50, height and weight within last two years; not required to be recorded on same day.

CRS uses any of the following codes to define the numerators.

	CPT Codes	ICD and Other Codes
Medical nutrition therapy	97802-97804, G0270, G0271	Primary or secondary provider codes: 07, 29 Clinic codes: 67 (dietary) or 36 (WIC)
Nutrition education		V POV: V65.3 dietary surveillance and counseling Patient education codes: ending "-N" (nutrition), "-MNT" (medical nutrition therapy), (or old code "-DT" (diet)) or containing V65.3, 97802-97804, G0270, or G0271.
Exercise education		V POV: V65.41 exercise counseling Patient education codes: ending "-EX" (exercise) or containing V65.41.
Related exercise and nutrition education		Patient education codes: ending "-LA" (lifestyle adaptation) or containing "OBS-" (obesity) or 278.00 or 278.01.

### **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

A list of at risk patients with education, if any.

### **Measure Source**

HP 2010 19-17

# **Measure Past Performance and Long-Term Targets for Diabetic Education**

Performance	Percent
HP 1997 data	42.0%

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_					Dec 31, 2				
					1 to Dec 3. to Dec 31,				
Nutrition and Exercise	 Educa	tion f	or At Ris	k Pat	 i ent				
	REPORT	%	PREV YR PERIOD		CHG from PREV YR %			CHG from BASE %	
# Overweight Active Cl	inical								
patients =>6	598		555			442			
<pre># w/medical nutrition</pre>									
therapy	44	7.4	23	4.1	+3.2	27	6.1	+1.2	
<pre># specific nutrition education provided</pre>	83	13.9	79	14.2	-0.4	78	17.6	-3.8	
# w/exercise educ	34	5.7	28	5.0	+0.6	35	7.9	-2.2	
# w/ other exec or nutrition educ	75	12.5	59	10.6	+1.9	24	5.4	+7.1	
# Male Overweight Acti	.ve								
Clinical pts =>6	251		230			182			
# w/medical nutrition									
therapy # specific nutrition	18	7.2	8	3.5	+3.7	10	5.5	+1.7	
education provided	36	14.3	32	13.9	+0.4	28	15.4	-1.0	
<pre># w/exercise educ # w/ other exec</pre>	16	6.4	12	5.2	+1.2	16	8.8	-2.4	
or nutrition educ	41	16.3	22	9.6	+6.8	11	6.0	+10.3	
# Female Overweight Ac	ctive								
Clinical pts =>6	347		325			260			
# w/medical nutrition									
therapy	26	7.5	15	4.6	+2.9	17	6.5	+1.0	
# specific nutrition education provided	47	13.5	47	14.5	-0.9	50	19.2	-5.7	

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# w/exercise educ	18	5.2	16	4.9	+0.3	19	7.3	-2.1
# w/ other exec or nutrition educ	34	9.8	37	11.4	-1.6	13	5.0	+4.8

Figure 2-68: Sample Report, Nutrition and Exercise Education for At Risk Patients

Nutrition and Exercise Ed	ducation for A	At Risk P	atient (c	on't)		
TOTA	AL OBESE ACTI		AL POPULA stributio			
# Obese Active Clinical	6-11	_	20-39	40-59	=>60	
CURRENT REPORT PERIOD						
# Obese Active Clinical	13	28	167	125	32	
# w/medical nutrition	0	0	1.5	1.0	2	
therapy % w/medical nutrition	0	2	15	10	3	
therapy	0.0	7.1	9.0	8.0	9.4	
cherapy	0.0	7.1	9.0	0.0	9.4	
<pre># # w/specific nutrition</pre>	education					
provided	0	3	22	28	7	
% # w/specific nutrition	education					
provided	0.0	10.7	13.2	22.4	21.9	
					_	
# w/exercise educ	0	1	8	14	5	
% w/exercise educ	0.0	3.6	4.8	11.2	15.6	
# w/other exec or						
nutrition educ	0	3	18	16	7	
% w/other exec or	0	3	10	10	,	
nutrition educ	0.0	10.7	10.8	12.8	21.9	
PREVIOUS YEAR PERIOD						
# Obese Active Clinical	14	26	137	116	32	
# w/medical nutrition	0	2	0	2	0	
therapy % w/medical nutrition	0	3	8	3	2	
therapy	0.0	11.5	5.8	2.6	6.3	
01101 ap 1	3.0	11.5	3.0	2.0	0.5	
# # w/specific nutrition	education					
provided	0	2	19	22	7	
$% \ \# \ w/specific nutrition$						
provided	0.0	7.7	13.9	19.0	21.9	
W /	^	0	4	1.4	4	
<pre># w/exercise educ % w/exercise educ</pre>	0.0	0.0	4 2.9	14 12.1	12 5	
% w/exercise educ	0.0	0.0	2.9	12.1	12.5	
# w/other exec or						
nutrition educ	0	2	13	23	3	
% w/other exec or						
nutrition educ	0.0	7.7	9.5	19.8	9.4	
CHANGE FROM PREV YR %						
medical nutrition	. 0 0	1 1	. 2 1	4	. 2 1	
therapy Spec nutr ed	+0.0	-4.4	+3.1 -0.7	+5.4	+3.1	
w/exercise educ	+0.0 +0.0	+3.0 +3.6	-0.7 +1.9	+3.4 -0.9	+0.0 +3.1	
w/exercise educ	TU.0	13.0	11.9	0.9	13.1	

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Figure 2-69: Sample Age Breakout Report, Nutrition and Exercise Education for At Risk Patients

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Nutrition and Exercise Education for At Risk Patients: List of at risk patients, with education if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, SANDRA KAY

AC-OW, AC-OB

PATIENT2, CAITLYN

AC-OW

PATIENT3, BRITNEY

AC-OW, AC-OB

PATIENT4, LORETTA

AC-OW, AC-OB

PATIENT5, HALEY

AC-OW AC-OR

PATIENT5, HALEY

AC-OW AC-OB

PATIENT5, HALEY

AC-OW AC-OB

PATIENT5, HALEY

AC-OW AC-OR

O00001 COMMUNITY #1 F 21

D00002 COMMUNITY #1 F 22

D00003 COMMUNITY #1 F 22

D00004 COMMUNITY #1 F 22

D00005 COMMUNITY #1 F 25

D00005 COMMUNITY #1 F 25 NUTR: 03/15/12 UTI-N SN NUTR: 05/07/12 HTN-N SN; EXER ED: 05/07/12 HTN-EX AC-OW, AC-OB PATIENT6, BRITTANY 000006 COMMUNITY #1 F 25 AC-OW, AC-OB EXER ED: 01/15/12 278.00-EX; LIFE: 01/15/12 278.00-EX

Figure 2-70: Sample Patient List, Nutrition, and Exercise Education for At Risk Patients

# 2.8.4 Physical Activity Assessment

#### **Denominators**

*Active Clinical patients* ages 5 and older. Broken down by gender and age groups (5-11, 12-19, 20-24, 25-34, 35-44, 45-54, 55-74, >75).

Numerator 1 (Active Clinical Patients assessed for physical activity during the Report Period). Broken down by gender and age groups (5-11, 12-19, 20-24, 25-34, 35-44, 45-54, 55-74, >75).

*User Population patients* ages 5 and older. Broken down by gender.

Numerator 1 (User Population Patients assessed for physical activity during the Report Period). Broken down by gender.

#### **Numerators**

Patients assessed for physical activity during the Report Period.

Patients from Numerator 1 who have received exercise education following their physical activity assessment.

# **Logic Description**

Age of the patient is calculated at beginning of Report Period.

CRS uses any of the following codes to define the numerators.

	ICD and Other Codes
Physical Activity Assessment	<b>Health Factors</b> : Any health factor for category Activity Level documented during the Report Period.
Exercise education	V POV: V65.41 exercise counseling
	Patient education codes: ending "-EX" (exercise) or containing V65.41.

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients with physical activity assessment and any exercise education.

DU November 25, 2012 Page 171  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000								
Physical Activity Ass	essment							
					CHG from PREV YR %			
Active Clinical Pts 5 and older	1,357		1,031			911		
# w/ exercise educ	19	1.4	0	0.0	+1.4	0	0.0	+1.4
<pre>w/ % of physical act assessment</pre>	-	21.1	0	0.0	+21.1	0	0.0	+21.1
Male Active Clinical =>5	570		430			374		
<pre># w/ physical activit assessment # w/ exercise educ</pre>	7	1.2	0	0.0	+1.2	0	0.0	+1.2
<pre>w/ % of physical act assessment</pre>	-	14.3	0	0.0	+14.3	0	0.0	+14.3
Female Active Clinica =>5	1 787		601			537		

# w/ physical activity								
# w/ exercise educ	1.5	0	0.0	+1.5	0	0.0	+1.5	
<pre>w/ % of physical activity assessment 3</pre>	25.0	0	0.0	+25.0	0	0.0	+25.0	
User Pop ages 5 and older 2,592		2,142			2,025			
2,052		_,			2,025			
# w/ physical activity assessment 19 # w/ exercise educ	0.7	0	0.0	+0.7	0	0.0	+0.7	
w/ % of physical activity assessment 4	21.1	0	0.0	+21.1	0	0.0	+21.1	
Male User Pop Pts								
=>5 1,211		988			945			
# w/ physical activity assessment 7	0.6	0	0.0	+0 6	0	0 0	+0.6	
# w/ exercise educ	0.0	U	0.0	+0.0	U	0.0	+0.0	
w/ % of physical activity								
assessment 1	14.3	0	0.0	+14.3	0	0.0	+14.3	
Female User Pop Pts								
=>5 1,381		1,154			1,080			
# w/ physical activity assessment 12	0.9	0	0 0	+0.9	0	0 0	+0.9	
# w/ exercise educ	0.9	V	0.0		3	0.0		
w/ % of physical activity								
assessment 3	25.0	0	0.0	+25.0	0	0.0	+25.0	

Figure 2-71: Sample Report, Physical Activity Assessment

Physical Activity Asses	sment	(con't)							
	TOTAL	ACTIVE		AL5 AND Distri	_				
	5-11	12-19	_			45-54	55-74	>74 yrs	
CURRENT REPORT PERIOD									
Total # AC Pts =>5 # w/ physical activity	132	157	133	233	215	216	225	46	
assessment	2	6	3	1	2	0	3	2	
% w/ physical activity assessment	1.5	3.8	2.3	0.4	0.9	0.0	1.3	4.3	
<pre># w/ exercise educ w/ % of physical activit</pre>	<b>.</b> .								
assessment % w/ exercise educ w/	y 1	2	1	0	0	0	0	0	
<pre>% of physical activit assessment</pre>	У 50.0	33.3	33.3	0.0	0.0	0.0	0.0	0.0	
PREVIOUS YEAR PERIOD Total # AC Pts =>5	141	137	128	172	151	140	137	25	

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<pre># w/ physical activity   assessment % w/ physical activity</pre>	0	0	0	0	0	0	0	0	
assessment		0.0	0.0	0.0	0.0	0.0	0.0	0.0	
<pre># w/ exercise educ w/ % of physical activi</pre>	ty								
<pre>assessment % w/ exercise educ w/ % of physical activi</pre>	0 tv	0	0	0	0	0	0	0	
assessment	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
CHANGE FROM PREV YR % # w/ physical activity									
assessment w/ exercise educ w/ % of physical activi		+3.8	+2.3	+0.4	+0.9	+0.0	+1.3	+4.3	
assessment	-	+33.3	+33.3	+0.0	+0.0	+0.0	+0.0	+0.0	

Figure 2-72: Sample Age Breakout Report, Physical Activity Assessment

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Physical Activity Assessment: List of patients with physical activity assessment and any exercise education. HRN COMMUNITY SEX AGE PATIENT NAME DENOMINATOR NUMERATOR PATIENT1, MISTY DAWN 000001 COMMUNITY #1 F 5 PHYS ACT: 08/08/12 VERY ACTIVE; EXER ED: 08/08/12 OBS-UP,AC ΕX PATIENT2, RITA ANN 000002 COMMUNITY #1 F 15 UP,AC PHYS ACT: 03/06/12 SOME ACTIVITY; EXER ED: 03/06/12 TO-EX PATIENT3, RHONDA SUE 000003 COMMUNITY #1 F 22
UP, AC PATIENT4, MARY 000004 COMMUNITY #1 F 28
UP, AC PHYS ACT: 12/12/12 SON PHYS ACT: 04/02/12 ACTIVE; EXER ED: 04/02/12 V65.41 PHYS ACT: 12/12/12 SOME ACTIVITY; PATIENT5, JOSEPH HENRY 000005 COMMUNITY #1 M 12 UP,AC PHYS ACT: 08/02/12 SOM PATIENT6,BOB 000006 COMMUNITY #1 M 17 PHYS ACT: 08/02/12 SOME ACTIVITY; UP,AC PHYS ACT: 05/05/12 INACTIVE; EXER ED: 05/05/12 OBS-EX

Figure 2-73: Sample Patient List, Physical Activity Assessment

# 2.8.5 Comprehensive Health Screening

#### **Denominators**

Active Clinical patients ages 2 and older.

Active Clinical patients ages 12 to 75.

Active Clinical patients ages 18 and older.

Female Active Clinical patients ages 15–40.

Active Clinical patients ages 5 and older.

Active Clinical patients ages 2 through 74.

Active Clinical patients ages 20 and over.

Active Clinical patients ages 5 and older.

#### **Numerators**

ALL Comprehensive Health Screening: Patients with Comprehensive Health Screening for which they are eligible, defined as having alcohol, depression, and Intimate Partner Violence/Domestic Violence (IPV/DV) screening, BMI calculated, and tobacco use, BP, and physical activity assessed.

**Note**: This does *not* include refusals.

Comprehensive Health Screening: Patients with Comprehensive Health Screening minus physical activity assessment for which they are eligible, defined as having alcohol, depression, and IPV/DV screening, BMI calculated, and tobacco use and BP assessed.

**Note**: This does *not* include physical activity assessment and does *not* include refusals.

Alcohol Screening: Patients screened for alcohol use or had an alcohol-related diagnosis or procedure during the Report Period.

**Note**: This numerator does *not* include refusals or alcohol-related patient education.

Depression Screening: Patients screened for depression or diagnosed with a mood disorder at any time during the Report Period.

**Note**: This numerator does *not* include refusals.

IPV/DV Screening: Patients screened for IPV/DV at any time during the Report Period.

**Note**: This numerator does *not* include refusals.

Tobacco Use Assessed: Patients who have been screened for tobacco use during the Report period.

BMI Available: Patients for whom a BMI could be calculated.

**Note**: This numerator does *not* include refusals.

BP Assessed: Patients with BP value documented at least twice in prior two years.

Physical Activity Assessed: Patients assessed for physical activity during the Report Period.

# **Logic Description**

Age of the patient is calculated at beginning of Report Period.

Alcohol screening definition: Any of the following during the Report Period: (a) Alcohol Screening Exam, any CAGE Health Factor, or Screening Diagnosis; (b) Alcohol-related diagnosis in POV, Current PCC or BHS Problem List; (c) Alcohol-related procedure; or (d) Patient education.

	ICD and Other Codes
Alcohol Screening	PCC Exam Code: 35
	<b>CPT code:</b> 99408, 99409, G0396, G0397, H0049, H0050, 3016F
	Any CAGE Health Factor
	V POV: V11.3 (history of alcoholism), V79.1 (screening for
	alcoholism)
	BHS Problem Code: 29.1 (Screening for Alcoholism)
	V Measurement in PCC or BHS: AUDT, AUDC, or CRFT
Alcohol-related	V POV, Current PCC or BHS Problem List: 303.*, 305.0*,
Diagnosis	291.*, 357.5*
	BHS POV: 10, 27, 29
Alcohol-related Procedure	<b>V Procedure:</b> 94.46, 94.53, 94.61-94.63, 94.67-94.69

Alcohol screening may be documented with either an exam code or the CAGE health factor in PCC or BHS. BHS problem codes can also currently be used.

Depression screening definition: CRS uses the following codes to define the numerator.

	ICD and Other Codes
Depression Screening	V Exam: Exam Code 36
	<b>V POV:</b> V79.0
	<b>CPT:</b> 1220F
	BHS Problem Code: 14.1 (Screening for Depression)
	V Measurement in PCC or BHS: PHQ2 or PHQ9
Mood Disorders	At least two visits in PCC or BHS for: Major Depressive Disorder, Dysthymic Disorder, Depressive Disorder NOS, Bipolar I or II Disorder, Cyclothymic Disorder, Bipolar Disorder NOS, Mood Disorder Due to a General Medical Condition, Substance-induced Mood Disorder, or Mood Disorder NOS.  V POV: 296.*, 291.89, 292.84, 293.83, 300.4, 301.13, or 311 BHS POV: 14, 15

IPV/DV screening definition: CRS uses the following codes to define the numerator.

	ICD and Other Codes
IPV/DV Screening	V Exam: Code 34
	BHS Exam: IPV/DV
IPV/DV Diagnosis	<b>V POV or current PCC or BHS Problem List:</b> 995.80-995.83, 995.85, V15.41, V15.42, V15.49 <b>BHS POV:</b> 43.*, 44.*
IPV/DV Education	<b>Patient education codes:</b> "DV-" or "-DV", 995.80-83, 995.85, V15.41, V15.42, or V15.49
IPV/DV Counseling	<b>V POV:</b> V61.11

Tobacco screening definition: CRS uses the following codes to define the numerator.

	CPT Codes	ICD and Other Codes
Screened for Tobacco Use	D1320, 99406, 99407, G0375 (old code), G0376 (old code), G8455-G8457 (old codes), G8402 (old code), G8453 (old code), 1034F (Current Tobacco Smoker), 1035F (Current Smokeless Tobacco User), 1036F (Current Tobacco Non-User), 1000F (Tobacco Use Assessed)	V POV or current Active Problem List: 305.1, 305.1* (old codes), 649.00-649.04, V15.82 Patient Education codes: "TO-", "-TO", "-SHS", 305.1, 305.1* (old codes), 649.00-649.04, V15.82, D1320, 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, 1036F, 1000F, G8455-G8457 (old codes), G8402 (old code), or G8453 (old code) Dental code: 1320 Health Factor categories: Tobacco, TOBACCO (SMOKING), TOBACCO (SMOKELESS – CHEWING/DIP), or TOBACCO (EXPOSURE)

BMI calculation definition: CRS calculates BMI at the time the report is run, using NHANES II. For 18 and under, a height and weight must be taken on the same day any time during the report period. For 19 through 50, height and weight must be recorded within last five years, not required to be on the same day. For over 50, height and weight within last two years, not required to be recorded on same day.

Blood pressure definition: CRS uses mean of last three Blood Pressures documented on non-ER visits in the past two years. If three BPs are not available, uses mean of last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) blood pressures and dividing by three (or two). If the systolic and diastolic values do not both meet the current category, then the value that is least controlled determines the category.

If CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on a non-ER visit during the Report Period.

*Physical Activity Assessment definition:* CRS uses the following codes to define the numerator.

	ICD and Other Codes
Physical Activity Assessment	<b>Health Factors</b> : Any health factor for category Activity Level documented during the Report Period.

# **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

# **Patient List Description**

List of patients with assessments received, if any.

DU November 25, 2012 Page 181  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Comprehensive Health	Screenin	ng							
					CHG from PREV YR %				
Active Clinical ages 2 and older			1,121			1,007			
# w/ Comprehensive H Screening-No Refusals # w/ Comprehensive H	56 ealth	3.9	39	3.5	+0.4	40	4.0	-0.1	
Screening-No Refusa or Phys Activity		4.6	46	4.1	+0.5	53	5.3	-0.6	
Active Clinical ages 12-75	1,180		867			753			
<pre># w/ alcohol screeni:   Dx/Proc/-No Refusal   or Pt Ed</pre>	s	8.1	12	1.4	+6.7	3	0.4	+7.7	
Active Clinical => 18	1,112		793			666			
# w/ Depression scre or Mood Disorder Dx No Refusals	-	7.3	42	5.3	+2.0	17	2.6	+4.7	
# Female Active Clinical ages 15-40	380		311			267			
# w/IPV/DV screening	-								

No Refusals	11	2 0	1	0.2	+2.6	0	0.0	+2.9
NO RELUSAIS	11	2.9	1	0.3	+2.0	U	0.0	+2.9
# Active Clinical Pts =>5	1,357		1,031			911		
# w/Tobacco Screening	624	46.0	426	41.3	+4.7	328	36.0	+10.0
Active Clinical Pts 2-74	1,400		1,096			982		
# w/BMI calculated- No Refusals	880	62.9	824	75.2	-12.3	712	72.5	-9.6
Active Clinical Pati	ents							
ages 20 and older	1,068		753			640		
<pre># w/ BPs documented w/in 2 yrs</pre>	643	60.2	557	74.0	-13.8	478	74.7	-14.5
Active Clinical Pts 5 and older	1,357		1,031			911		
# w/ physical activi assessment	-	1.4	0	0.0	+1.4	0	0.0	+1.4

Figure 2-74: Sample Report, Comprehensive Health Screening

UP-User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Comprehensive Health Screening: List of patients with assessments received, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, SANDRA KAY 000001 COMMUNITY #1 F 15 ALL COMP HEALTH: ALC: 03/06/12 POV V11.3; IPV: 03/06/12 Ex 34; TOB: 09/05/12 NEVER SMOKED; BMI: 17.49; PHYS ACT: 03/06/12 SOME ACTIVITY 000002 COMMUNITY #1 F 16 PATIENT2, CAITLYN PATIENT3, BRITNEY 000003 COMMUNITY #1 F 16 TOB: 10/26/12 CESSATION-SMOKER 000004 COMMUNITY #1 PATIENT4, LORETTA F 17 ALC: 10/14/12 HF CAGE 1/4 000005 COMMUNITY #1 F 18 PATIENT5, HALEY BMI: 19.79; BP: 125/67 PATIENT6, BRITTANY 000006 COMMUNITY #1 F 19 ALC: 10/30/12 CPT G0397; TOB: 08/11/12 CURRENT SMOKER, STATUS UNKNOWN; BMI: 21.01

Figure 2-75: Sample Patient List, Comprehensive Health Screening

# 2.8.6 Cardiovascular Disease and Cholesterol Screening

#### **Denominators**

Active Clinical patients ages 23 and older. Broken down by gender.

User Population patients ages 23 and older. Broken down by gender.

Active IHD patients, defined as all Active Clinical patients diagnosed with ischemic heart disease (IHD) prior to the Report Period, and at least two visits during the Report Period, and two IHD-related visits ever. Broken down by gender.

#### **Numerators**

Patients with documented blood total cholesterol screening any time in the past five years.

a. Patients with high total cholesterol levels, defined as equal to or greater than (=>) 240

Patients with LDL completed in the past five years, regardless of result.

- a. Patients with LDL <=100
- b. Patients with LDL 101-130
- c. Patients with LDL 131-160
- d. Patients with LDL >160

# **Logic Description**

Age is calculated at the beginning of the Report Period.

CRS uses the following codes to define the IHD denominator.

	ICD and Other Codes
Ischemic Heart Disease	<b>V POV</b> : 410.0-412.*, 414.0-414.9, 429.2

Total Cholesterol definition: Searches for most recent cholesterol test with a result during the Report Period. If more than one cholesterol test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If a cholesterol test with a result is not found, CRS searches for the most recent cholesterol test without a result.

LDL Cholesterol definition: Searches for most recent LDL test with a result during the Report Period. If more than one LDL test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If an LDL test with a result is not found, CRS searches for the most recent LDL test without a result.

CRS uses the following codes to define LDL and total cholesterol.

Test	CPT Codes	LOINC Codes	Taxonomy
LDL	80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F For numerator LDL =<100, CPT 3048F will count as meeting the measure.	Yes	DM AUDIT LDL CHOLESTEROL TAX
Total Cholesterol	82465	Yes	DM AUDIT CHOLESTEROL TAX

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients with cholesterol or LDL value, if any.

### **Measure Source**

HP 2020 HDS-6, HDS-7

# **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2006 Performance (blood total cholesterol screening)	48.0%
IHS FY 2005 Performance (blood total cholesterol screening)	43.0%
HP 1998 baseline	67.0%
HP 2020 goal for adults who have had blood cholesterol checked (HDS-6)	82.1%
HP 2020 goal for adults with high cholesterol (HDS-7)	13.5%

```
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                             November 25, 2012
                                                                     Page 185
       *** IHS 2012 Selected Measures with Community Specified Report ***
                             DEMO INDIAN HOSPITAL
                 Report Period: Jan 01, 2012 to Dec 31, 2012
             Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                Baseline Period: Jan 01, 2000 to Dec 31, 2000
Cardiovascular Disease and Cholesterol Screening
                                                                    % CHG from
                    REPORT
                                  PREV YR
                                             % CHG from BASE
                    PERIOD
                                  PERIOD
                                                PREV YR % PERIOD
                                                                       BASE %
```

Active Clinical Pts									
=> 23	986		676			569			
, 23	200		0,0			307			
# w/ Total Cholesterol									
	0.40	25 2	017	20 1	6.0	001	25 2	10.0	
screen w/in 5 yrs	248	25.2	21/	32.1	-6.9	201	35.3	-10.2	
A. # w/ High Chol =>240									
w/ % of Total Chol									
Screen	21	8.5	23	10.6	-2.1	28	13.9	-5.5	
# w/LDL done									
in past 5 yrs	251	25.5	185	27.4	-1.9	114	20.0	+5.4	
A. # w/LDL =<100									
w/ % of Total LDL									
Screen	108	43.0	95	51.4	-8.3	46	40.4	+2.7	
B. # w/LDL 101-130	100	13.0	) )	31.4	0.5	40	10.1	12.7	
w/ % of Total LDL	п.	00 1	4.4	00 0	. 5 2	2.5	20 17	1 6	
Screen	/3	29.1	44	23.8	+5.3	35	30.7	-1.6	
C. # w/LDL 131-160									
w/ % of Total LDL									
Screen	25	10.0	25	13.5	-3.6	13	11.4	-1.4	
D. # w/LDL >160									
w/ % of Total LDL									
Screen	15	6.0	9	4.9	+1.1	10	8.8	-2.8	
Male Active Clinical									
Pts =>23	408		272			220			
103 -725	100		2/2			220			
#/ Matal Chalastanal									
# w/ Total Cholesterol	100	06.0	0.7	25 5	۰	0.5	20.6	10.5	
screen w/in 5 yrs	106	26.0	9.7	35.7	-9.7	85	38.6	-12.7	
A. # w/ High Chol =>240									
w/ % of Total Chol									
Screen	11	10.4	14	14.4	-4.1	8	9.4	+1.0	
# w/LDL done									
in past 5 yrs	115	28.2	91	33.5	-5.3	59	26.8	+1.4	
A. # w/LDL =<100									
w/ % of Total LDL									
Screen	53	46.1	45	49.5	-3.4	23	39.0	+7.1	
B. # w/LDL 101-130	33	10.1	13	10.5	5.1	23	55.0	. , • ±	
w/ % of Total LDL									
	2.5	01 7	1.0	10 0	. 2 0	1.0	20 F	-8.8	
Screen	25	21.7	18	19.8	+2.0	TR	30.5	-8.8	
C. # w/LDL 131-160									
w/ % of Total LDL									
Screen	8	7.0	12	13.2	-6.2	5	8.5	-1.5	
D. # w/LDL >160									
w/ % of Total LDL									
Screen	10	8.7	8	8.8	-0.1	4	6.8	+1.9	

Figure 2-76: Sample Report, CVD and Cholesterol Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Cardiovascular Disease and Cholesterol Screening: List of patients with cholesterol or LDL value if any.

PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR

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```
PATIENT100, JASON AARON 000100 COMMUNITY #1
                                             M 46
PATIENT101, JOHN THOMAS 000101 COMMUNITY #1 M 47
PATIENT102, DAKOTA CHEY 000102 COMMUNITY #1 M 47
PATIENT103, TRAVIS CLINT 000103 COMMUNITY #1 M 47
                             CHOL 04/13/12 210
PATIENT104, TRACY MITCHE 000104 COMMUNITY #1 M 47
                             CHOL 03/15/12 167; LDL 08/15/12 105
UP,AC,IHD
PATIENT105, RUSSELL DALE 000105 COMMUNITY #1 M 48
                             LDL 04/01/12 CPT 3048F
PATIENT106, CURTIS DWAYN 000106 COMMUNITY #1 M 49
                            CHOL 03/04/09 139; LDL 06/04/10 68
UP,AC CHOL 03/04/09 139; LDI PATIENT107,RONALD 000107 COMMUNITY #1 M 49
UP,AC
                      CHOL 01/07/08 213; LDL 08/01/11 122
```

Figure 2-77: Sample Patient List, CVD and Cholesterol Screening

### 2.8.7 Cardiovascular Disease and Blood Pressure Control

#### **Denominators**

All Active Clinical patients ages 20 and over. Broken down by gender.

All *User Population patients* ages 20 and older. Broken down by gender.

Active IHD patients, defined as all Active Clinical patients diagnosed with IHD prior to the Report Period, and at least two visits during the Report Period, and two IHD-related visits ever. Broken down by gender.

#### **Numerators**

Patients with Blood Pressure value documented at least twice in prior two years.

- a. Patients with normal BP, defined as < 120/80, i.e., the mean systolic value is less than (<) 120 *and* the mean diastolic value is less than (<) 80.
- b. Patients with Pre Hypertension I BP, defined as => 120/80 and < 130/80, i.e., the mean systolic value is equal to or greater than (=>) 120 and less than (<) 130 and the mean diastolic value is equal to 80.
- c. Patients with Pre Hypertension II BP, defined as => 130/80 and <140/90, i.e., the mean systolic value is equal to or greater than (=>) 130 and less than (<) 140 *and* the mean diastolic value is equal to or greater than (=>) 80 and less than (<) 90.
- d. Patients with Stage 1 Hypertension BP, defined as => 140/90 and <160/100, i.e., the mean systolic value is equal to or greater than (=>) 140 and less than (<) 160 *and* the mean diastolic value is equal to or greater than (=>) 90 and less than (<) 100.

e. Patients with Stage 2 Hypertension BP, defined as => 160/100, i.e., the mean systolic value is equal to or greater than (=>) 160 and the mean diastolic value is equal to or greater than (=>) 100.

### **Logic Description**

Age of the patient is calculated at beginning of the Report Period.

Blood pressure definition: CRS uses mean of last three Blood Pressures documented on non-ER visits in the past two years. If three BPs are not available, uses mean of last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) blood pressures and dividing by three (or two). If the systolic and diastolic values do not both meet the current category, then the value that is least controlled determines the category.

For the BP documented numerator only, if CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on a non-ER visit during the Report Period.

CRS uses the following codes to define the IHD numerator.

	ICD and Other Codes			
Ischemic Heart Disease	<b>V POV</b> : 410.0-412.*, 414.0-414.9, 429.2			

### **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

#### **Patient List Description**

List of Patients => 20 or who have IHD with BP value, if any.

#### **Measure Source**

HP 2020 HDS-5

# Measure Past Performance and Long-Term Targets

Measure	Percent
HP 2020 goal for adults with high blood pressure (140/90)	26.9%

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\*\*\* IHS 2012 Selected Measures with Community Specified Report \*\*\*

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Baseline Pe							
Cardiovascular Disease and E	Blood E	ressure C	ontro	1			
				CHG from PREV YR %			
Active Clinical Patients ages 20 and older 1,068		753			640		
# w/ BPs documented							
w/in 2 yrs 643 A. # w/Normal BP w/ %	60.2	557	74.0	-13.8	478	74.7	-14.5
of Total Screened 128 B. # w/Pre HTN I BP w/ %	19.9	134	24.1	-4.2	121	25.3	-5.4
of Total Screened 107	16.6	115	20.6	-4.0	83	17.4	-0.7
C. # w/Pre HTN II BP w/ % of Total Screened 148 D. # w/Stage 1 HTN BP w/	23.0	114	20.5	+2.6	105	22.0	+1.1
% of Total Screened 173		150	26.9	+0.0	130	27.2	-0.3
E. # w/Stage 2 HTN BP w/ % of Total Screened 37	5.8	39	7.0	-1.2	39	8.2	-2.4
Male Active Clinical Patient ages 20 and older 432	s	293			241		
ages 20 and order 432		293			241		
	53.5	203	69.3	-15.8	177	73.4	-20.0
	3.5	23	11.3	-7.9	22	12.4	-9.0
B. # w/Pre HTN I BP w/ % of Total Screened 27 C. # w/Pre HTN II BP w/ %	11.7	37	18.2	-6.5	22	12.4	-0.7
of Total Screened 64	27.7	46	22.7	+5.0	45	25.4	+2.3
D. # w/Stage 1 HTN BP w/ % of Total Screened 89		79	38.9	-0.4	63	35.6	+2.9
E. # w/Stage 2 HTN BP w/ % of Total Screened 16		17	8.4	-1.4	25	14.1	-7.2
Female Active Clinical Patie ages 20 and older 636	ents	460			399		
# w/ BPs documented							
w/in 2 yrs 412 A. # w/Normal BP w/ %	64.8	354	77.0	-12.2	301	75.4	-10.7
of Total Screened 120 B. # w/Pre HTN I BP w/ %	29.1	111	31.4	-2.2	99	32.9	-3.8
of Total Screened 80 C. # w/Pre HTN II BP w/ %	19.4	78	22.0	-2.6	61	20.3	-0.8
of Total Screened 84	20.4	68	19.2	+1.2	60	19.9	+0.5
D. # w/Stage 1 HTN BP w/ % of Total Screened 84	20.4	71	20.1	+0.3	67	22.3	-1.9
E. # w/Stage 2 HTN BP w/ % of Total Screened 21	5.1	22	6.2	-1.1	14	4.7	+0.4

Figure 2-78: Sample Report, CVD and Blood Pressure Control

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Cardiovascular Disease and Blood Pressure Control: List of Patients => 20 or who have IHD with BP value, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR -----\_\_\_\_\_ PATIENT1,SANDRA KAY 000001 COMMUNITY #1 F 21 UP,AC PATIENT2, EVELYN 000002 COMMUNITY #1 F 21 /3080F UP /3080F
PATIENT3, MICHELLE 000003 COMMUNITY #1 F 22
UP, AC 125/67 PRE HTN 1
PATIENT4, CAITLYN 000004 COMMUNITY #1 F 22
UP, AC, IHD 131/67 PRE HTN 2
PATIENT5, BRITNEY JANE 000005 COMMUNITY #1 F 22
UP AC 102/56 NORMAL UP,AC 102/56 NORMAL
PATIENT6,KATHRYN ANNE 000006 COMMUNITY #1 F 22
UP,AC 161/90 STG 2 HTN
PATIENT7,RHONDA 000007 COMMUNITY #1 F 22
UP,AC 152/05 CTG 1 VTV 153/85 STG 1 HTN UP,AC

Figure 2-79: Sample Patient List, CVD and Blood Pressure Control

# 2.8.8 Controlling High Blood Pressure

#### **Denominator**

Active Clinical patients ages 18 through 85 diagnosed with hypertension and no documented history of ESRD. Broken down by gender and age groups (18–45 and 46-85).

#### **Numerators**

Number of patients with Blood Pressure value documented during the Report Period.

- f. Patients with *normal blood pressure*, defined as < 120/80; that is, the mean systolic value is less than (<) 120 *and* the mean diastolic value is less than (<) 80.
- g. Patients with *Pre Hypertension I BP*, defined as => 120/80 and less than < 130/80, that is, the mean systolic value is equal to or greater than (=>) 120 and less than (<) 130 *and* the mean diastolic value is equal to 80.
- h. Patients with *Pre Hypertension II BP*, defined as => 130/80 and <140/90; that is, the mean systolic value is equal to or greater than (=>) 130 and less than (<) 140 *and* the mean diastolic value is equal to or greater than (=>) 80 and less than (<) 90.

- i. Patients with *Stage 1 Hypertension* Blood Pressure (BP), defined as => 140/90 and <160/100; that is, the mean systolic value is equal to or greater than (=>) 140 and less than (<) 160 *and* the mean diastolic value is equal to or greater than (=>) 90 and less than (<) 100.
- j. Patients with *Stage 2 Hypertension BP*, defined as => 160/100; that is, the mean systolic value is equal to or greater than (=>) 160 *and* the mean diastolic value is equal to or greater than (=>) 100.

# **Logic Description**

Age of the patient is calculated at beginning of the Report Period.

Blood pressure definition: CRS uses mean of last three BPs documented on non-ER visits during the Report Period. If three BPs are not available, uses mean of last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) blood pressures and dividing by three (or two). If the systolic and diastolic values do not both meet the current category, then the value that is least controlled determines the category.

For the BP documented numerator only, if CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on non-ER visit during the Report Period.

CRS uses the following codes to define ESRD and hypertension.

	CPT Codes	ICD and Other Codes
ESRD	36145, 36147, 36800, 36810, 36815, 36818, 36819, 36820, 36821, 36831-36833, 50300, 50320, 50340, 50360, 50365, 50370, 50380, 90951-90970 or old codes 90918-90925, 90935, 90937, 90939 (old code), 90940, 90945, 90947, 90989, 90993, 90997, 90999, 99512, G0257, G0308-G0327 (old codes), G0392 (old code), G0393 (old code), or \$9339	<b>V POV</b> : 585.5, 585.6, V42.0, V45.1, (old code), V45.11, V45.12, or V56.* <b>V Procedure</b> : 38.95, 39.27, 39.42, 39.43, 39.53, 39.93-39.95, 54.98, or 55.6*
Hypertension		V POV or Problem List Prior to the Report Period and at Least One Hypertension POV during Report Period: 401.*

# **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

# **Patient List Description**

List of patients with hypertension and BP value, if any.

### **Measure Source**

HP 2020 HDS-5, HDS-12

# **Measure Past Performance and Long-Term Targets**

Measure	Percent
HP 2020 goal for adults with high blood pressure (140/90)	26.9%
HP 2020 goal for adults with high blood pressure and whose blood pressure is controlled	61.2%

DU November 25, 2012 Page 200  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Controlling High Blood P	ress	ure							
					CHG from PREV YR %				
Active Clinical Pts 18-85 w/HTN dx	108		101			91			
# w/ BPs documented A. # w/Normal BP w/ %	96	88.9	92	91.1	-2.2	85	93.4	-4.5	
of Total Screened B. # w/Pre HTN I BP w/ %		5.2	6	6.5	-1.3	3	3.5	+1.7	
of Total Screened C. # w/Pre HTN II BP w/		8.3	13	14.1	-5.8	8	9.4	-1.1	
% of Total Screened D. # w/Stage 1 HTN BP w/		24.0	17	18.5	+5.5	19	22.4	+1.6	
% of Total Screened E. # w/Stage 2 HTN BP w/	36	37.5	41	44.6	-7.1	42	49.4	-11.9	
% of Total Screened		10.4	15	16.3	-5.9	13	15.3	-4.9	
A. Active Clinical Pts 18-45 w/HTN dx	23		19			13			
# w/ BPs documented	17	73.9	16	84.2	-10.3	11	84.6	-10.7	

A. # w/Normal BP w/ %									
of Total Screened B. # w/Pre HTN I BP w/ %	2	11.8	2	12.5	-0.7	0	0.0	+11.8	
of Total Screened	1	5.9	1	6.3	-0.4	1	9.1	-3.2	
C. # w/Pre HTN II BP w/	2	15.6	0	10 5	. = 1	0	10.0	0 5	
% of Total Screened D. # w/Stage 1 HTN BP w/	3	17.6	2	12.5	+5.1	2	18.2	-0.5	
% of Total Screened	7	41.2	6	37.5	+3.7	7	63.6	-22.5	
E. # w/Stage 2 HTN BP w/ % of Total Screened	2	11.8	_	21 2	-19.5	1	9.1	+2.7	
% of focal Screened	۷	11.0	5	31.3	-19.5		9.1	+4.7	
B. Active Clinical Pts									
46-85 w/HTN dx	85		82			78			
# w/ BPs									
	79	92.9	76	92.7	+0.3	74	94.9	-1.9	
A. # w/Normal BP w/ % of Total Screened	3	3.8	4	5.3	-1.5	3	4.1	-0.3	
B. # w/Pre HTN I BP w/ %	3	3.0	-	3.3	1.3	3		0.3	
of Total Screened	7	8.9	12	15.8	-6.9	7	9.5	-0.6	
C. # w/Pre HTN II BP w/ % of Total Screened	20	25.3	15	19.7	+5.6	17	23.0	+2.3	
D. # w/Stage 1 HTN BP w/	_ •			• ·	3.0				
% of Total Screened	29	36.7	35	46.1	-9.3	35	47.3	-10.6	
E. # w/Stage 2 HTN BP w/ % of Total Screened	8	10.1	10	13.2	-3.0	12	16.2	-6.1	
- 12 10001 001001100	Ŭ				5.0		,_	· · ·	

Figure 2-80: Sample Report, Controlling High Blood Pressure

Figure 2-81: Sample Patient List, Controlling High Blood Pressure

# 2.8.9 Comprehensive CVD-Related Assessment

### **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 40.6% for the proportion of atrisk patients who have a comprehensive assessment.

### **Denominators**

Active IHD patients ages 22 and older, defined as all Active Clinical patients diagnosed with IHD prior to the Report Period, and at least two visits during the Report Period, and two IHD-related visits ever. (GPRA Denominator)

- a. Active IHD patients ages 22 and older who are not Active Diabetic
- b. Active IHD patients ages 22 and older who are Active Diabetic

#### **Numerators**

*BP Assessed:* Patients with Blood Pressure value documented at least twice in prior two years.

LDL Assessed: Patients with LDL completed in past five years, regardless of result.

*Tobacco Use Assessed:* Patients who have been screened for tobacco use during the Current Report period.

BMI Available: Patients for whom a BMI could be calculated.

**Note**: This numerator does *not* include refusals.

*Lifestyle Counseling:* Patients who have received any lifestyle adaptation counseling, including medical nutrition therapy, or nutrition, exercise or other lifestyle education during the Report Period.

Patients with *comprehensive CVD assessment*, defined as having BP, LDL, and tobacco use assessed, BMI calculated, and lifestyle counseling.

**Note**: This does *not* include depression screening and *does not* include refusals of BMI. (GPRA Numerator)

*Refusal of BMI:* Patients who refused a height or weight measurement and for whom a BMI could not be calculated.

Depression Screening: Patients screened for depression or diagnosed with a mood disorder at any time during the Report Period.

**Note**: This numerator does *not* include refusals.

### **Logic Description**

Age of the patient is calculated at beginning of the Report Period.

Diabetes defined as: Diagnosed with diabetes (first POV in V POV with 250.00–250.93) prior to the Current Report Period, and at least two visits during the Current Report Period, and two DM-related visits ever. Patients not meeting these criteria are considered non-diabetics.

*IHD diagnosis defined as*: 410.0–412.\*, 414.0–414.9, 428.\* or 429.2 recorded in the V POV file.

*Blood pressure definition:* Having a minimum of two BPs documented on non-ER visits during the Report Period in past two years. If CRS does not find two BPs, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on non-ER visit during the past two years.

LDL definition: Finds the most recent test done in the last five years, regardless of the results of the measurement.

BMI definition: CRS calculates when the report is run, using NHANES II. For 19 through 50, height and weight must be recorded within last five years, not required to be on the same day. For over 50, height and weight within last two years; not required to be recorded on same day.

*BMI Refusal definition:* Refusals of a height and/or weight measurement include REF, NMI, and UAS and must be documented during the past year. For ages 19 and older, the height and the weight must be refused during the past year and are not required to be on the same visit.

CRS uses the following codes and taxonomies to define the numerators.

Test	CPT Codes	LOINC Codes	Taxonomy
	80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F	Yes	DM AUDIT LDL CHOLESTEROL TAX

Test	CPT Codes		LOINC Codes	Taxonomy
Tobacco Screening	D1320, 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, 1036F, 1000F, G8455- G8457 (old codes), G8402 (old code) or G8453 (old code)	Any health factor for category Tobacco, TOBACCO (SMOKING), TOBACCO (SMOKELESS – CHEWING/DIP), or TOBACCO (EXPOSURE) (see table on next page) V POV or current Active Problem List: 305.1, 305.1* (old codes), 649.00-649.04, V15.82 Patient education codes: containing "TO-", "-TO", "-SHS", 305.1, 305.1* (old codes), 649.00-649.04, V15.82, D1320, 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, 1036F, 1000F, G8455- G8457 (old codes), G8402 (old code) or G8453 (old code) Dental code: 1320		
Medical Nutrition	97802-97804,	Primary or secondary provider		
Therapy	G0270, G0271	<b>codes</b> : 07, 29, 97, 99 <b>Clinic codes</b> : 67 (dietary) or 36 (WIC)		
Nutrition Education		V POV: V65.3 dietary surveillance and counseling Patient education codes: ending "-N" (nutrition) or "-MNT" (medical nutrition therapy) (or old code "-DT" (diet)) or containing V65.3.		
Exercise Education		V POV: V65.41 exercise counseling Patient education codes: ending "-EX" (exercise) or containing V65.41.		
Related Exercise and Nutrition Education		Patient education codes: ending "-LA" (lifestyle adaptation) or containing "OBS-" (obesity) or 278.00 or 278.01.		

Test	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Depression Screening		V Exam: Exam Code 36 V POV: V79.0 CPT: 1220F BHS Problem Code: 14.1 (Screening for Depression) V Measurement in PCC or BH: PHQ2 or PHQ9		
Mood Disorders		At least 2 visits in PCC or BHS for: Major Depressive Disorder, Dysthymic Disorder, Depressive Disorder NOS, Bipolar I or II Disorder, Cyclothymic Disorder, Bipolar Disorder NOS, Mood Disorder Due to a General Medical Condition, Substance-induced Mood Disorder, or Mood Disorder NOS (see codes below).  V POV: 296.*, 291.89, 292.84, 293.83, 300.4, 301.13, or 311 BHS POV: 14, 15		

All existing national Tobacco Health Factors listed below are counted as tobacco screening.

Health Factor					
Ceremonial	Previous Smokeless				
Cessation-Smokeless	Previous (Former) Smokeless				
Cessation-Smoker	Previous Smoker				
Current Smokeless	Previous (Former) Smoker				
Current Smoker	Smoke Free Home				
Current Smoker, status unknown	Smoker In Home				
Current smoker, every day	Current Smoker & Smokeless				
Current smoker, some day	Exposure To Environmental Tobacco Smoke				
Non-Tobacco User					

# **Key Logic Changes from CRS Version 11.1**

Added POV code V81.1 to BP Documented definition.

# **Patient List Description**

List of patients with assessments received, if any.

### **Measure Source**

Not Available

# **Measure Past Performance Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance (Comprehensive CVD Assessment)	39.8%
IHS FY 2010 Performance (Comprehensive CVD Assessment)	35.0%
IHS FY 2009 Performance (Comprehensive CVD Assessment)	32.0%
IHS FY 2008 Performance (Comprehensive CVD Assessment)	30.0%
IHS FY 2007 Performance (Comprehensive CVD Assessment)	30.0%
IHS FY 2011 Performance (BP Assessed)	97.2%
IHS FY 2010 Performance (BP Assessed)	98.0%
IHS FY 2009 Performance (BP Assessed)	97.0%
IHS FY 2008 Performance (BP Assessed)	98.0%
IHS FY 2011 Performance (LDL Assessed)	92.9%
IHS FY 2010 Performance (LDL Assessed)	92.0%
IHS FY 2009 Performance (LDL Assessed)	91.0%
IHS FY 2008 Performance (LDL Assessed)	90.0%
IHS FY 2011 Performance (Tobacco Assessed)	84.2%
IHS FY 2010 Performance (Tobacco Assessed)	84.0%
IHS FY 2009 Performance (Tobacco Assessed)	83.0%
IHS FY 2008 Performance (Tobacco Assessed)	79.0%
IHS FY 2011 Performance (BMI Assessed)	97.2%
IHS FY 2010 Performance (BMI Assessed)	98.0%
IHS FY 2008 Performance (BMI Assessed or Refused)	85.0%
IHS FY 2011 Performance (Lifestyle Counseling)	45.3%
IHS FY 2010 Performance (Lifestyle Counseling)	41.0%
IHS FY 2009 Performance (Lifestyle Counseling)	39.0%
IHS FY 2008 Performance (Lifestyle Counseling)	38.0%
IHS FY 2011 Performance (Depression Screen)	75.7%
IHS FY 2010 Performance (Depression Screen)	72.0%
IHS FY 2009 Performance (Depression Screen)	62.0%
IHS FY 2008 Performance (Depression Screen)	53.0%

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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000

Comprehensive CVD-Related Assessment
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CRS Clinical Performance Measure Logic Manual for FY 2012 Clinical Measures
December 2011

REF	ORT	<b>ે</b>	PREV YR	%	CHG from	BASE	<b>ે</b>	CHG from	
PEF	RIOD		PERIOD		PREV YR %	PERIOD		BASE %	
Active IHD Pts 22+ (GPRA)	77		45			36			
# w/ BPs documented w/in 2 yrs # w/ LDL done in	72	93.5	45	100.0	-6.5	36	100.0	-6.5	
past 5 yrs # w/Tobacco Screening	60	77.9	38	84.4	-6.5	30	83.3	-5.4	
w/in 1 yr # w/BMI calculated	66	85.7	38	84.4	+1.3	27	75.0	+10.7	
-No Refusals # w/ lifestyle	72	93.5	44	97.8	-4.3	35	97.2	-3.7	
educ w/in 1 yr # w/ BP, LDL, tobacco, BMI and life counseling	43	55.8	22	48.9	+7.0	22	61.1	-5.3	
-No Refusals (GPRA) # w/BMI refusal	35	45.5	19	42.2	+3.2	14	38.9	+6.6	
(No BMI) # w/ Depression screenir	2 1g	2.6	0	0.0	+2.6	0	0.0	+2.6	
or Mood Disorder DX -No Refusals	20	26.0	4	8.9	+17.1	2	5.6	+20.4	
A. Active IHD Pts 22+ and are NOT Active									
Diabetic	41		20			17			
# w/ BPs documented w/in 2 yrs # w/LDL done	36	87.8	20	100.0	-12.2	17	100.0	-12.2	
in past 5 yrs # w/Tobacco Screening	28	68.3	17	85.0	-16.7	13	76.5	-8.2	
w/in 1 yr # w/BMI calculated	33	80.5	15	75.0	+5.5	13	76.5	+4.0	
-No Refusals # w/ lifestyle	38	92.7	20	100.0	-7.3	16	94.1	-1.4	
educ w/in 1 yr # w/ BP, LDL, tobacco,	24	58.5	7	35.0	+23.5	7	41.2	+17.4	
BMI and life counseling -No Refusals # w/BMI refusal	19	46.3	6	30.0	+16.3	4	23.5	+22.8	
(No BMI) # w/ Depression screening	o 1g,	0.0	0	0.0	+0.0	0	0.0	+0.0	
or Mood Disorder DX -No Refusals	13	31.7	1	5.0	+26.7	1	5.9	+25.8	

Figure 2-82: Sample Report, Comprehensive CVD-Related Assessment

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Comprehensive CVD-Related Assessment: List of patients with assessments received, if any.

CRS Clinical Performance Measure Logic Manual for FY 2012 Clinical Measures
December 2011

PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR
PATIENT1, CAITLYN IHD 07/25/12 NEVER SMOKED; PHQ9	000001 COMMUNITY #1  F  22 ALL: BP: 131/67 PRE STG 1; LDL: 06/08/12; TOB: BMI: 25.4; LIFE: 08/15/012 UTI-N SN; DEP: 05/03/12 Meas
IHD 33.7	000002 COMMUNITY #1 F 40 BP: 112/66 NORMAL; TOB: 08/26/12 NEVER SMOKED; BMI:
IHD 03/03/12 Prv 97; DEP:	
IHD,AD 12/30/12 NEVER SMOKED;	000004 COMMUNITY #1  F  68 ALL: BP: 150/82 STG 1 HTN; LDL: 09/13/12; TOB: BMI: 26.8; LIFE: 10/19/12 PM-LA 000005 COMMUNITY #1  F  70
IHD, AD CURRENT SMOKER, STATUS	BP: 149/72 STG 1 HTN; LDL: 10/24/12; TOB: 10/24/12
IHD,AD 09/11/12 Meas PHQ2	BP: 3077F/; TOB: 01/27/12 G8402; BMI: 43.4; DEP:

Figure 2-83: Sample Patient List: Comprehensive CVD-Related Assessment

# 2.8.10 Appropriate Medication Therapy after a Heart Attack

### **Denominator**

Active Clinical patients aged 35 and older discharged for an AMI during the first 51 weeks of the Report Period and were not readmitted for any diagnosis within seven days of discharge. Broken down by gender.

#### **Numerators**

Patients with active prescription for or who have a contraindication/previous adverse reaction to *beta-blockers*.

**Note**: This numerator does *not* include refusals.

Patients with active prescription for or who have a contraindication/previous adverse reaction to ASA (aspirin) or other anti-platelet agent.

**Note**: This numerator does *not* include refusals.

Patients with active prescription for or who have a contraindication/ previous adverse reaction to *ACEIs/ARBs*.

**Note**: This numerator does *not* include refusals.

Patients with active prescription for or who have a contraindication/ previous adverse reaction to *statins*.

**Note**: This numerator does *not* include refusals.

Also included for the numerators above are subnumerators:

- a. Patients with active prescription for the specified medication
- Patients with contraindication/previous adverse reaction to the specified medication

Patients with documented refusal of the specified medication

Patients with active prescriptions for *all post-AMI medications* (i.e., beta-blocker, ASA/anti-platelet, ACEI/ARB, and statin), and/or who have a contraindication/previous adverse reaction.

**Note**: This numerator does *not* include refusals.

# **Logic Description**

Age is calculated at the beginning of the Report Period.

Acute Myocardial Infarction (AMI) definition: POV 410.\*1 (i.e., first eligible episode of an AMI) with Service Category H. If patient has more than one episode of AMI during the first 51 weeks of the Report Period, CRS will include only the first discharge.

**Denominator Exclusions** 

Patients meeting any of the following conditions will be excluded from the denominator.

- 1. Patients with Discharge Type of Irregular (AMA), Transferred, or contains "Death."
- 2. Patients readmitted for any diagnosis within seven days of discharge.
- 3. Patients with a Diagnosis Modifier of C (Consider), D (Doubtful), M (Maybe, Possible, Perhaps), O (Rule Out), P (Probable), R (Resolved), S (Suspect, Suspicious), or T (Status Post).
- 4. Patients with a Provider Narrative beginning with "Consider"; "Doubtful"; "Maybe"; "Possible"; "Perhaps"; "Rule Out"; "R/O"; "Probable"; "Resolved"; "Suspect"; "Suspicious"; or "Status Post."

Numerator Logic

In the logic below, "ever" is defined as anytime through the end of the Report Period.

To be included in the numerators, a patient must meet one of the three conditions below:

- An active prescription (not discontinued as of [discharge date + 7 days] and does
  not have a comment of RETURNED TO STOCK) that was prescribed prior to
  admission, during the inpatient stay, or within seven days after discharge.
  "Active" prescription defined as: Days Prescribed > ((Discharge Date + 7 days) Order Date); or
- 2. A refusal of the medication at least once during hospital stay through seven days after discharge date; or...
- 3. Have a contraindication/previous adverse reaction to the indicated medication.

Refusals and contraindications/previous adverse drug reactions (ADR)/allergies are only counted if a patient did not have a prescription for the indicated medication. Patients without a prescription who have a refusal and/or contraindication/ADR/allergy will be counted in subnumerators B–C. Because a patient may have both a refusal and a contraindication/ADR/allergy, the subnumerator totals of A–C may not add up the to the numerator total.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

## **Beta-Blocker Numerator Logic**

Beta-blocker medication codes defined with medication taxonomy BGP HEDIS BETA BLOCKER MEDS. (Medications are: Noncardioselective Beta Blockers: Carteolol, Carvedilol, Labetalol, Nadolol, Penbutolol, Pindolol, Propranolol, Timolol, Sotalol; Cardioselective Beta Blockers: Acebutolol, Atenolol, Betaxolol, Bisoprolol, Metoprolol, Nebivolol; and Antihypertensive Combinations: Atenolol-chlorthalidone, Bendroflumethiazide-nadolol, Bisoprolol-hydrochlorothiazide, Hydrochlorothiazide-metoprolol, and Hydrochlorothiazide-propranolol.)

Refusal of beta-blocker: REF refusal of any beta-blocker medication in site-populated medication taxonomy BGP HEDIS BETA BLOCKER MEDS at least once during hospital stay through seven days after discharge date.

CRS uses the following codes to define contraindications to beta-blockers.

Contraindication to Beta- Blockers (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Asthma		<b>POV:</b> 2 diagnoses of 493* on different visit dates
Hypotension		POV: 1 diagnosis of 458*
Heart Block >1 Degree		<b>POV:</b> 1 diagnosis of 426.0, 426.12, 426.13, 426.2, 426.3, 426.4, 426.51, 426.52, 426.53, 426.54, or 426.7
Sinus Bradycardia		POV: 1 diagnosis of 427.81
COPD		<b>POV:</b> 2 diagnoses on different visit dates of 491.2*, 496, or 506.4, or a combination of any of these codes, such as 1 visit with 491.20 and 1 with 496
NMI Refusal	G8011 (old code) at least once during hospital stay through 7 days after discharge date	Refusal: NMI (not medically indicated) refusal for any beta-blocker at least once during hospital stay through 7 days after discharge date

CRS uses the following codes to define adverse drug reactions/documented allergies to beta-blockers.

	ICD and Other Codes
Adverse Drug Reaction/Allergy to Beta-Blockers (any of the codes occurring ever)	<b>POV:</b> 995.0-995.3 AND E942.0
	Entry in ART (Patient Allergies File): "beta block*"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "beta block*",
	"bblock*" or "b block*"

# ASA (aspirin)/Other Anti-Platelet Numerator Logic

ASA medication codes defined with medication taxonomy DM AUDIT ASPIRIN DRUGS.

Other anti-platelet medication codes defined with medication taxonomy site-populated DM AUDIT ANTI-PLATELET DRUGS taxonomy.

Refusal of ASA/other anti-platelet: REF refusal of any ASA or anti-platelet medication in site-populated medication taxonomies DM AUDIT ASPIRIN DRUGS or BGP ANTI-PLATELET DRUGS at least once during hospital stay through seven days after discharge date.

CRS uses the following codes to define contraindications to ASA/other anti-platelets.

Contraindication to ASA/Other Anti-Platelets (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Active prescription for Warfarin/Coumadin at time of arrival or prescribed at discharge		Site-Populated Drug Taxonomy: BGP CMS WARFARIN MEDS
Hemorrhage		<b>POV</b> : 459.0
NMI Refusal	G8008 (old code) at least once during hospital stay through 7 days after discharge date	Refusal: NMI (not medically indicated) refusal for any aspirin at least once during hospital stay through 7 days after discharge date

CRS uses the following codes to define adverse drug reactions/documented allergies to ASA/other anti-platelets.

	ICD and Other Codes
	<b>POV</b> : 995.0-995.3 AND E935.3
	Entry in ART (Patient Allergies File): "aspirin"
(any of the codes occurring ever)	Entry in Problem List or in Provider Narrative for
	any POV 995.0-995.3 or V14.8: "ASA" or "aspirin"

# **ACEI/ARB Numerator Logic**

Ace Inhibitor (ACEI) medication codes defined with medication taxonomy BGP HEDIS ACEI MEDS. ACEI medications are: Angiotensin Converting Enzyme Inhibitors (Benazepril, Captopril, Enalapril, Fosinopril, Lisinopril, Moexipril, Perindopril, Quinapril, Ramipril, Trandolopril).

Antihypertensive Combinations (Amlodipine-benazepril, Benazepril-hydrochlorothiazide, Captopril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Fosinopril-hydrochlorothiazide, Hydrochlorothiazide-lisinopril, Hydrochlorothiazide-moexipril, Hydrochlorothiazide-quinapril, Trandolapril-verapamil).

*Refusal of ACEI:* REF refusal of any ACE inhibitor medication in site-populated medication taxonomy BGP HEDIS ACEI MEDS at least once during hospital stay through seven days after discharge date.

CRS uses the following codes to define contraindications to ACE inhibitors.

Contraindication to ACE Inhibitors (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the Report Period with V22.0-V23.9, V72.42, 640.*-649.*, or 651.*-676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.
	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI (not medically indicated) refusal for any ACE inhibitor at least once during hospital stay through 7 days after discharge date

CRS uses the following codes to define adverse drug reactions/documented allergies to ACE inhibitors.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to ACE Inhibitors (any of the codes occurring ever)	<b>POV:</b> 995.0-995.3 AND E942.6
	Entry in ART (Patient Allergies File): "ace inhibitor" or "ACEI"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "ace i*" or "ACEI"

ARB (Angiotensin Receptor Blocker) medication codes defined with medication taxonomy BGP HEDIS ARB MEDS. ARB medications: Angiotensin II Inhibitors (Candesartan, Eprosartan, Irbesartan, Losartan, Olmesartan, Telmisartan, Valsartan.)

Antihypertensive Combinations (Aliskiren-valsartan, Amlodipine-hydrochlorothiazide-valsartan, Amlodipine-olmesartan, Candesartan-hydrochlorothiazide, Eprosartan-hydrochlorothiazide, Hydrochlorothiazide-Irbesartan, Hydrochlorothiazide-Losartan, Hydrochlorothiazide-olmesartan, Hydrochlorothiazide-Valsartan).

*Refusal of ARB:* REF refusal of any ARB medication in site-populated medication taxonomy BGP HEDIS ARB MEDS at least once during hospital stay through seven days after discharge date.

CRS uses the following codes to define contraindications to ARBs.

Contraindication to ARBs (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy	Abortion: Any of the	POV or Problem List: At least two visits during the Report Period with V22.0-V23.9, V72.42, 640.*-649.*, or 651.*-676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.  Abortion: Any of the following POVs: 635*,
	following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260-S2267	636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22

Contraindication to ARBs (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
NMI Refusal		<b>Refusal:</b> NMI refusal for any ARB at least once during hospital stay through 7 days after discharge date

CRS uses the following codes to define adverse drug reactions/documented allergies to ARBs.

	ICD and Other Codes
Adverse Drug Reaction/Allergy to	<b>POV:</b> 995.0-995.3 AND E942.6
ARBs (any of the codes occurring ever)	Entry in ART (Patient Allergies File): "Angiotensin Receptor Blocker" or "ARB"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "Angiotensin Receptor Blocker" or "ARB"

# **Statins Numerator Logic**

Statin medication codes defined with medication taxonomy BGP HEDIS STATIN MEDS. Statin medications: Atorvostatin (Lipitor), Fluvastatin (Lescol), Lovastatin (Altocor), Mevacor, Pravastatin (Pravachol), Simvastatin (Zocor), Rosuvastatin (Crestor).

Statin Combination Products: Advicor, Caduet, PraviGard Pac, Vytorin.

*Refusal of Statin:* REF refusal of any statin medication in site-populated medication taxonomy BGP HEDIS STATIN MEDS at least once during hospital stay through seven days after discharge date.

CRS uses the following codes to define contraindications to statins.

Contraindication to Statins (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the Report Period with V22.0–V23.9, V72.42, 640.* –649.*, or 651.* –676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.

Contraindication to Statins (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period
Acute Alcoholic Hepatitis		POV: 571.1 during the Report Period
NMI Refusal		<b>Refusal:</b> NMI refusal for any statin at least once during hospital stay through 7 days after discharge date

CRS uses the following codes to define adverse drug reactions/documented allergies to statins.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to Statins (any of the codes occurring ever unless otherwise noted)	Site-Populated Lab Taxonomy or LOINC Taxonomy: DM AUDIT ALT TAX, with ALT and/or AST > 3x the Upper Limit of Normal (ULN) (i.e., Reference High) on 2 or more consecutive visits during the Report Period
	Site-Populated Lab Taxonomy or LOINC Taxonomy: BGP CREATINE KINASE TAX, with Creatine Kinase (CK) levels > 10x ULN or CK > 10,000 IU/L during the Report Period
	<b>POV:</b> Myopathy/Myalgia, defined as any of the following during the Report Period: 359.0-359.9, 729.1, 710.5, or 074.1
	<b>POV:</b> 995.0-995.3 AND E942.9
	Entry in ART (Patient Allergies File): "statin" or "statins"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "Statin" or "Statins"

# All Medications Numerator Logic

To be included in this numerator, a patient must have a prescription, refusal, or a contraindication for *all* of the four medication classes (i.e., beta-blocker, ASA/other anti-platelet, ACEI/ARB, *and* statin).

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients with AMI, with appropriate medication therapy, if any.

# **Measure Source**

American Heart Association/American College of Cardiology Guidelines for the Treatment of AMI.

# **Measure Past Performance and Long-Term Targets**

None

DU November 25, 2012 Page 216  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000								
Appropriate Medication	n Thera	py af	ter a Heart	. Att	ack			
	REPORT PERIOD	%	PREV YR PERIOD		CHG from BAS: PREV YR % PE			HG from BASE %
Active Clinical Pts 3 hospitalized for AMI	_		0			0		
# w/beta-blocker Rx/contra/ADR -No Refusals A. # w/beta-blocker	27	38.0	0	0.0	+38.0	0	0.0	+38.0
Rx w/ % of Total	5	18.5	0	0.0	+18.5	0	0.0	+18.5
B. # w/contra/ADR w/ % of Total # w/ beta-blocker	22	81.5	0	0.0	+81.5	0	0.0	+81.5
refusal	2	2.8	0	0.0	+2.8	0	0.0	+2.8
# w/ASA Rx/contra/ADR -No Refusals A. # w/ASA		12.7	0	0.0	+12.7	0	0.0	+12.7
Rx w/% of Total B. # w/contra/ADR	3	33.3	0	0.0	+33.3	0	0.0	+33.3
w/ % of Total # w/ ASA refusal		66.7 4.2				0 0	0.0	+66.7 +4.2
# w/ACEI/ARB Rx/contra/ADR -No Refusals	19	26.8	0	0.0	+26.8	0	0.0	+26.8
A. # w/ACEI/ARB Rx w/% of Total	4	21.1	0	0.0	+21.1	0	0.0	+21.1

B. # w/contra/ADR w/ % of Total # w/ ACEI/ARB	15	78.9	0	0.0	+78.9	0	0.0	+78.9	
refusal	2	2.8	0	0.0	+2.8	0	0.0	+2.8	
<pre># w/statin Rx/contra/ADR</pre>									
-No Refusals A. # w/statin	17	23.9	0	0.0	+23.9	0	0.0	+23.9	
Rx w/% of Total B. # w/contra/ADR	5	29.4	0	0.0	+29.4	0	0.0	+29.4	
w/% of Total # w/ Statin	12	70.6	0	0.0	+70.6	0	0.0	+70.6	
Refusal	2	2.8	0	0.0	+2.8	0	0.0	+2.8	
# w/Rx/contra/ADR of AI									
meds-No Refusals	4	5.6	0	0.0	+5.6	0	0.0	+5.6	

Figure 2-84: Sample Report, Appropriate Medication Therapy after a Heart Attack

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Appropriate Medication Therapy after a Heart Attack: List of patients with AMI, with appropriate medication therapy, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, CECELIA 000001 COMMUNITY #1 F 37 AC BETA: 06/30/12 Contra 2/3 heart block POV 426.3 PATIENT2, KATHLEEN 000002 COMMUNITY #1 F 38 ACEI/ARB: Contra pregnant; STATIN: Contra pregnant PATIENT3, KIMBERLY A 000003 COMMUNITY #1 F 49 ASA: 12/16/11 CLOPIDOGREL BISULFATE 75MG TAB; ACEI/ARB: 01/14/12 Contra BF-HC; STATIN: 01/14/12 Contra BF-HC PATIENT4, TIMOTHY JOHN 000004 COMMUNITY #1 M 57 ACEI/ARB: 06/01/12 Contra NMI CAPTOPRIL 25MG TABS PATIENT5, FELIPE 000005 COMMUNITY #1 M 57 PATIENT6, JAMES DALTON 000006 COMMUNITY #1 M 77 ALL MEDS: BETA: 07/23/12 Contra CPT G8011; ASA: 07/23/12 Contra CPT G8008; ACEI/ARB: 07/27/10 Contra POV 396.0; STATIN: 06/05/12 SIMVASTATIN 40MG TAB

Figure 2-85: Sample Patient List: Appropriate Medication Therapy after a Heart Attack

# 2.8.11 Persistence of Appropriate Medication Therapy after a Heart Attack

#### **Denominator**

Active Clinical patients 35 and older diagnosed with an AMI six months prior to the Report Period through the first six months of the Report Period. Broken down by gender.

#### **Numerators**

Patients with a 135-day course of treatment with beta-blockers, or who have a contraindication/previous adverse reaction to beta-blocker therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 135-day course of treatment with ASA (aspirin) or other anti-platelet agent or who have a contraindication/previous adverse reaction to ASA/anti-platelet therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 135-day course of treatment with ACEIs/ARBs or who have a contraindication/previous adverse reaction to ACEI/ARB therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 135-day course of treatment with statins or who have a contraindication/previous adverse reaction to statin therapy.

Note: This numerator does *not* include refusals.

Also included for the numerators above are subnumerators:

- a. Patients with active prescription for the specified medication
- Patients with contraindication/previous adverse reaction to the specified medication

Patients with documented refusal of the specified medication

Patients with a 135-day course of treatment for all post-AMI medications (i.e., beta-blocker, ASA/anti-platelet, ACEI/ARB, and statin) following first discharge date or visit date, including previous active prescriptions; and/or who have a contraindication/previous adverse reaction.

**Note**: This numerator does *not* include refusals.

# **Logic Description**

Age is calculated at the beginning of the Report Period.

Acute Myocardial Infarction (AMI) definition: POV 410.0\*–410.9\* or 412. AMI diagnosis may be made at an inpatient or outpatient visit but must occur between six months prior to beginning of Report Period through first six months of the Report Period. Inpatient visit defined as Service Category of H (Hospitalization). If patient has more than one episode of AMI during the timeframe, CRS will include only the first hospital discharge or ambulatory visit.

#### **Denominator Exclusions**

Patients meeting any of the following conditions will be excluded from the denominator.

- If inpatient visit, patients with Discharge Type of Irregular (AMA), Transferred, or contains "Death."
- Patients with a Diagnosis Modifier of C (Consider), D (Doubtful), M (Maybe, Possible, Perhaps), O (Rule Out), P (Probable), R (Resolved), S (Suspect, Suspicious), or T (Status Post).
- Patients with a Provider Narrative beginning with "Consider," "Doubtful,"
   "Maybe," "Possible," "Perhaps," "Rule Out," "R/O," "Probable," "Resolved,"
   "Suspect," "Suspicious," or "Status Post."

# **Numerator Logic**

In the logic below, "ever" is defined as anytime through the end of the Report Period.

To be included in the numerators, a patient must meet one of the three conditions below.

- 1. A total days' supply >= 135 days in the 180 days following discharge date for inpatient visits or visit date for ambulatory visits. Medications must not have a comment of RETURNED TO STOCK. Prior active prescriptions can be included if the treatment days fall within the 180 days following discharge/visit date. Prior active prescription defined as most recent prescription (see codes below) prior to admission/visit date with the number of days supply equal to or greater than the discharge/visit date minus the prescription date; *or*
- 2. A refusal of the medication at least once at time of diagnosis through the 180 days after AMI; *or*
- 3. Have a contraindication/previous adverse reaction to the indicated medication.

4. Refusals and contraindications/previous ADR/allergies are only counted if a patient did not have a prescription for the indicated medication. Patients without a prescription who have a refusal and/or contraindication/ADR/allergy will be counted in subnumerators B–C. Because a patient may have both a refusal and a contraindication/ADR/allergy, the sub-numerator totals of A–C may not add up the to the numerator total.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Example of patient included in the beta-blocker numerator who has prior active prescription:

- Admission Date: 2/1/2011, Discharge Date: 2/15/2011
- Must have 135 days prescribed by 8/13/2011 (Discharge Date+180)
- Prior Beta-Blocker Rx Date: 1/15/2011
- # Days Prescribed: 60 (treats patient through 3/15/2011)
- Discharge Date minus Rx Date: 2/15/2011-1/15/2011 = 31, 60 is >= 31, prescription is considered Prior Active Rx
- 3/15/2011 is between 2/15 and 8/13/2011, thus remainder of Prior Active Rx can be counted toward 180-day treatment period
- # Remaining Days Prescribed from Prior Active Rx:
- (60-(Discharge Date-Prior Rx Date) = 60-(2/15/2011-1/15/2011) = 60-31 = 29
- Rx #2: 4/1/2011, # Days Prescribed: 90
- Rx #3: 7/10/2011, #Days Prescribed: 90
- Total Days Supply Prescribed between 2/15 and 8/13/2011: 29+90+90=209

## **Beta-Blocker Numerator Logic**

Beta-blocker medication codes defined with medication taxonomy BGP HEDIS BETA BLOCKER MEDS. (Medications are: Noncardioselective Beta Blockers: Carteolol, Carvedilol, Labetalol, Nadolol, Penbutolol, Pindolol, Propranolol, Timolol, Sotalol; Cardioselective Beta Blockers: Acebutolol, Atenolol, Betaxolol, Bisoprolol, Metoprolol, Nebivolol; and Antihypertensive Combinations: Atenolol-chlorthalidone, Bendroflumethiazide-nadolol, Bisoprolol-hydrochlorothiazide, Hydrochlorothiazide-metoprolol and Hydrochlorothiazide-propranolol.)

*Refusal of beta-blocker:* REF refusal of any beta-blocker medication in site-populated medication taxonomy BGP HEDIS BETA BLOCKER MEDS at least once during the period admission/visit date through the 180 days after discharge/visit date.

CRS uses the following codes to define contraindications to beta-blockers.

Contraindication to Beta- Blockers (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
Asthma		<b>POV:</b> 2 diagnoses of 493* on different visit dates
Hypotension		POV: 1 diagnosis of 458*
Heart Block >1 Degree		<b>POV:</b> 1 diagnosis of 426.0, 426.12, 426.13, 426.2, 426.3, 426.4, 426.51, 426.52, 426.53, 426.54, or 426.7
Sinus Bradycardia		POV: 1 diagnosis of 427.81
COPD		<b>POV:</b> 2 diagnoses on different visit dates of 491.2*, 496, or 506.4, or a combination of any of these codes, such as 1 visit with 491.20 and 1 with 496
NMI Refusal	G8011 (old code) at least once during the period admission/visit date through the 180 days after discharge/visit date	Refusal: NMI refusal for any beta-blocker at least once during the period admission/visit date through the 180 days after discharge/visit date

CRS uses the following codes to define adverse drug reactions/documented allergies to beta-blockers.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy	<b>POV:</b> 995.0-995.3 AND E942.0
to Beta-Blockers	Entry in ART (Patient Allergies File): "beta block*"
(any of the codes occurring anytime up to the 180 days after discharge/visit date)	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "beta block*", "bblock*" or "b block*"

# ASA (Aspirin)/Other Anti-Platelet Numerator Logic

ASA medication codes defined with medication taxonomy DM AUDIT ASPIRIN DRUGS.

Other anti-platelet medication codes defined with medication taxonomy site-populated BGP ANTI-PLATELET DRUGS taxonomy.

Refusal of ASA/other anti-platelet: REF refusal of any ASA or anti-platelet medication in site-populated medication taxonomies DM AUDIT ASPIRIN DRUGS or DM AUDIT ANTI-PLATELET DRUGS at least once during the period admission/visit date through the 180 days after discharge/visit date.

CRS uses the following codes to define contraindications to ASA/other anti-platelets.

Contraindication to ASA/Other Anti-Platelets (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
Active prescription for Warfarin/Coumadin during the period admission/visit date through the 180 days after discharge/visit date		Site-Populated Drug Taxonomy: BGP CMS WARFARIN MEDS
Hemorrhage		<b>POV:</b> 459.0
NMI Refusal	G8008 (old code) at least once during the period admission/visit date through the 180 days after discharge/visit date	Refusal: NMI (not medically indicated) refusal for any aspirin/other anti-platelet at least once during the period admission/visit date through the 180 days after discharge/visit date

CRS uses the following codes to define adverse drug reactions/documented allergies to ASA/other anti-platelets.

	ICD and Other Codes
Adverse Drug Reaction/Allergy to	<b>POV:</b> 995.0-995.3 AND E935.3
ASA/Other Anti-Platelets (any of the codes occurring anytime up to the 180 days after	Entry in ART (Patient Allergies File): "aspirin"
discharge/visit date)	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "ASA" or "aspirin"

## **ACEI/ARB Numerator Logic**

Ace Inhibitor (ACEI) medication codes defined with medication taxonomy BGP HEDIS ACEI MEDS. ACEI medications are: Angiotensin Converting Enzyme Inhibitors (Benazepril, Captopril, Enalapril, Fosinopril, Lisinopril, Moexipril, Perindopril, Quinapril, Ramipril, Trandolopril).

Antihypertensive Combinations (Amlodipine-benazepril, Benazepril-hydrochlorothiazide, Captopril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Fosinopril-hydrochlorothiazide, Hydrochlorothiazide-lisinopril, Hydrochlorothiazide-moexipril, Hydrochlorothiazide-quinapril, Trandolapril-verapamil).

Refusal of ACEI: REF refusal of any ACE Inhibitor medication in site-populated medication taxonomy BGP HEDIS ACEI MEDS at least once during the period admission/visit date through the 180 days after discharge/visit date.

CRS uses the following codes to define contraindications to ACE inhibitors.

Contraindication to ACE Inhibitors (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the period admission/visit date through the 180 days after discharge/visit date with V22.0-V23.9, V72.42, 640.*-649.*, or 651.*-676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the period, CRS will use the first two visits in the period.
	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the period admission/visit date through the 180 days after discharge/visit date Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF-MK, or BF-N during the period admission/visit date through the 180 days after discharge/visit date
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI refusal for any ACE inhibitor at least once during the period admission/visit date through the 180 days after discharge/visit date

CRS uses the following codes to define adverse drug reactions/documented allergies to ACE inhibitors.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to ACE Inhibitors (any of the codes occurring anytime up to the 180 days after discharge/visit date)	<b>POV:</b> 995.0-995.3 AND E942.6
	Entry in ART (Patient Allergies File): "ace inhibitor" or "ACEI"
	Contained within or Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "ace i*" or "ACEI"

ARB (Angiotensin Receptor Blocker) medication codes defined with medication taxonomy BGP HEDIS ARB MEDS. ARB medications are: Angiotensin II Inhibitors (Candesartan, Eprosartan, Irbesartan, Losartan, Olmesartan, Telmisartan, Valsartan.).

Antihypertensive Combinations (Aliskiren-valsartan, Amlodipine-hydrochlorothiazide-valsartan, Amlodipine-olmesartan, Candesartan-hydrochlorothiazide, Eprosartan-hydrochlorothiazide, Hydrochlorothiazide-Irbesartan, Hydrochlorothiazide-Losartan, Hydrochlorothiazide-olmesartan, Hydrochlorothiazide-Valsartan).

Refusal of ARB: REF refusal of any ARB medication in site-populated medication taxonomy BGP HEDIS ARB MEDS at least once during the period admission/visit date through the 180 days after discharge/visit date.

CRS uses the following codes to define contraindications to ARBs.

Contraindication to ARBs (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the period admission/visit date through the 180 days after discharge/visit date with V22.0-V23.9, V72.42, 640.* –649.*, or 651.*–676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (clinic code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the period, CRS will use the first two visits in the period.

Contraindication to ARBs (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
	Abortion: Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		<b>POV:</b> V24.1 during the period admission/visit date through the 180 days after discharge/visit date
		Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF-MK, or BF-N during the period admission/visit date through the 180 days after discharge/visit date
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI refusal for any ARB at least once during the period admission/visit date through the 180 days after discharge/visit date

CRS uses the following codes to define adverse drug reactions/documented allergies to ARBs.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy	<b>POV:</b> 995.0-995.3 AND E942.6
to ARBs (any of the codes occurring anytime up to the 180 days after discharge/visit date)	Entry in ART (Patient Allergies File): "Angiotensin Receptor Blocker" or "ARB"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "Angiotensin Receptor Blocker" or "ARB"

# **Statins Numerator Logic**

*Statin medication codes* defined with medication taxonomy BGP HEDIS STATIN MEDS. Statin medications: Atorvostatin (Lipitor), Fluvastatin (Lescol), Lovastatin (Altocor), Mevacor, Pravastatin (Pravachol), Simvastatin (Zocor), Rosuvastatin (Crestor).

Statin Combination Products: Advicor, Caduet, PraviGard Pac, Vytorin.

*Refusal of Statin:* REF refusal of any statin medication in site-populated medication taxonomy BGP HEDIS STATIN MEDS at least once during admission/visit date through the 180 days after discharge/visit date.

CRS uses the following codes to define contraindications to statins.

Contraindication to Statins (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the period admission/visit date through the 180 days after discharge/visit date with V22.0–V23.9, V72.42, 640.*-649.*, or 651.* –676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the period, CRS will use the first two visits in the period.
	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedure:</b> 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the period admission/visit date through the 180 days after discharge/visit date Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF-MK, or BF-N during the period admission/visit date through the 180 days after discharge/visit date
Acute Alcoholic Hepatitis		POV: 571.1 during the period admission/visit date through the 180 days after discharge/visit date
NMI Refusal		<b>Refusal:</b> NMI refusal for any statin at least once during the period admission/visit date through the 180 days after discharge/visit date

CRS uses the following codes to define adverse drug reactions/documented allergies to statins.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to Statins (any of the codes occurring anytime up to the 180 days after discharge/visit date unless otherwise noted)	Site-Populated Lab Taxonomy or LOINC Taxonomy: DM AUDIT ALT TAX, with ALT and/or AST > 3x the Upper Limit of Normal (ULN) (i.e., Reference High) on 2 or more consecutive visits during the period admission/visit date through the 180 days after discharge/visit date
	Site-Populated Lab Taxonomy or LOINC Taxonomy: BGP CREATINE KINASE TAX, with Creatine Kinase (CK) levels > 10x ULN or CK > 10,000 IU/L during the period admission/visit date through the 180 days after discharge/visit date
	<b>POV:</b> Myopathy/Myalgia, defined as any of the following during the period admission/visit date through the 180 days after discharge/visit date: 359.0–359.9, 729.1, 710.5, or 074.1
	<b>POV:</b> 995.0–995.3 AND E942.9
	Entry in ART (Patient Allergies File): "statin" or "statins"
	Entry in Problem List or in Provider Narrative for any POV 995.0–995.3 or V14.8: "Statin" or "Statins"

# **All Medications Numerator Logic**

To be included in this numerator, a patient must have a prescription, refusal, or a contraindication for *all* of the four medication classes (i.e., beta-blocker, ASA/other anti-platelet, ACEI/ARB, *and* statin).

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients with AMI, with persistent medication therapy, if any.

### **Measure Source**

American Heart Association/American College of Cardiology Guidelines for the Treatment of AMI.

## **Measure Past Performance and Long-Term Targets**

None

DU *** IHS 2012	Selected		ember 25, ures with		nity Specif	ied Rep		age 228 **	
DEMO INDIAN HOSPITAL									
Previo	Report Period: Jan 01, 2012 to Dec 31, 2012 Previous Year Period: Jan 01, 2011 to Dec 31, 2011								
Bas	eline Pe 	riod: 	Jan 01,	2000 	to Dec 31, 2	2000 			
			. '	_		7			
Persistence of Appro	priate M	edica	tion Thera	apy ai	ter a Heart	Attac	ζ		
	REPORT PERIOD	%	PREV YR PERIOD		CHG from BA				
Active Clinical Pts									
w/ AMI DX	61		4			4			
# w/135-day beta-blo Rx/contra/ADR	cker								
-No Refusals		37.7	2	50.0	-12.3	3	75.0	-37.3	
A. # w/135-day beta : Rx w/ % of Total		13.0	2	100.0	-87.0	2	66.7	-53.6	
B. # w/contra/ADR		05.0		2 2	.05.0		22.2	.50 6	
<pre>w/ % of Total # w/ beta-blocker</pre>	20	87.0	0	0.0	+87.0	1	33.3	+53.6	
refusal	1	1.6	0	0.0	+1.6	0	0.0	+1.6	
# w/135-day ASA Rx/contra/ADR									
-No Refusals	5	8.2	0	0.0	+8.2	2	50.0	-41.8	
A. # w/135-day ASA Rx w/% of Total	1	20.0	0	0.0	+20.0	2	100.0	-80.0	
B. # w/contra/ADR									
<pre>w/ % of Total # w/ ASA refusal</pre>		80.0		0.0					
		3.3	, and the second	0.0		ŭ	0.0		
# w/135-day ACEI/ARB Rx/contra/ADR									
-No Refusals		21.3	1	25.0	-3.7	1	25.0	-3.7	
A. # w/135-day ACEI/		7.7	1	100.0	-92.3	1	100.0	-92.3	
B. # w/contra/ADR									
<pre>w/ % of Total # w/ ACEI/ARB</pre>	12	92.3	0	0.0	+92.3	0	0.0	+92.3	
refusal	1	1.6	0	0.0	+1.6	0	0.0	+1.6	
# w/135-day statin									
Rx/contra/ADR -No Refusals	6	9.8	2	50.0	-40.2	2	50.0	-40.2	
A. # w/135-day stati:		9.8	2	50.0	-40.2	2	50.0	-40.2	
Rx w/% of Total	2	33.3	2	100.0	-66.7	2	100.0	-66.7	
B. # w/contra/ADR w/ % of Total	4	66.7	0	0.0	+66.7	0	0.0	+66.7	
# w/ Statin refusal	2	3.3		0.0		0	0.0		
# w/Rx/contra/ADR of	ALL								
meds-No Refusals	2	3.3	0	0.0	+3.3	1	25.0	-21.7	

Figure 2-86: Sample Report, Persistence of Appropriate Medication Therapy after a Heart Attack

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Persistence of Appropriate Medication Therapy after a Heart Attack: List of
patients with AMI, with persistent medication therapy, if any
PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR
                       NUMERATOR
------
PATIENT1,RHONDA 000001 COMMUNITY #1 F 35
AC ALL MEDS: BETA: 03/07/10 03/13/10 Contra 2 POV asthma;
ASA: 05/23/12 Contra NMI ASPIRIN 325MG CAP; ACEI/ARB: 07/01/12 ADR Problem List
995.0 ACEI; STATIN: 07/01/12 ADR creat kinase of 5000
PATIENT2, KATHLEEN 000002 COMMUNITY #1 F 38 AC BETA: 06/02/12 ADR PO
                            BETA: 06/02/12 ADR POV V14.8
PATIENT3, KIMBERLY A 000003 COMMUNITY #1 F 49
AC BETA: 03/01/12 Contra
                            BETA: 03/01/12 Contra 2/3 heart block POV 426.12;
ACEI/ARB: Contra pregnant; STATIN: Contra pregnant
PATIENT4, TIMOTHY 000004 COMMUNITY #1 M 57
                      ACEI/ARB: 06/15/12 Contra POV V24.1; STATIN: 06/15/12
Contra POV V24.1
PATIENT5, JOSHUA 000005 COMMUNITY #1 M 63
                             ACEI/ARB: 02/03/12 Contra Aortic POV 395.0
```

Figure 2-87: Sample Patient List: Persistence of Appropriate Medication Therapy after a Heart Attack

# 2.8.12 Appropriate Medication Therapy in High Risk Patients

## **Denominators**

Active IHD patients ages 22 and older; defined as all Active Clinical patients diagnosed with IHD prior to the Report Period, and at least two visits during the Report Period, and two IHD-related visits ever.

- Active IHD patients ages 22 and older who are not Active Diabetic.
- Active IHD patients ages 22 and older who are Active Diabetic

#### **Numerators**

Patients with a 180-day course of treatment with *beta-blockers* during the Report Period, or who have a contraindication/previous adverse reaction to beta-blocker therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 180-day course of treatment with ASA (aspirin) or other anti-platelet agent during the Report Period, or who have a contraindication/previous adverse reaction to ASA/anti-platelet therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 180-day course of treatment with *ACEIs/ARBs* during the Report Period, or who have a contraindication/previous adverse reaction to ACEI/ARB therapy.

**Note**: This numerator does *not* include refusals.

Patients with a 180-day course of treatment with *statins* during the Report Period, or who have a contraindication/previous adverse reaction to statin therapy.

**Note**: This numerator does *not* include refusals.

Also included for the numerators above are subnumerators:

- a. Patients with active prescription for the specified medication
- Patients with contraindication/previous adverse reaction to the specified medication

Patients with documented refusal of the specified medication

Patients with a 180-day course of treatment for *all medications* (i.e., beta-blocker, aspirin/anti-platelet, ACEI/ARB, *and* statin) during the Report Period, and/or who have a contraindication/previous adverse reaction.

**Note**: This numerator does *not* include refusals.

# **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

*IHD* diagnosis defined as: 410.0–412.\*, 414.0–414.9, or 429.2 recorded in the V POV file.

Diabetes defined as: Diagnosed with diabetes (first POV in V POV with 250.00-250.93) prior to the Current Report Period, and at least two visits during the Current Report period, and two DM-related visits ever. Patients not meeting these criteria are considered non-diabetics.

Numerator Logic

In the logic below, "ever" is defined as anytime through the end of the Report Period.

To be included in the numerators, a patient must meet one of the three conditions below:

- 1. Prescription(s) for the indicated medication with a total days supply of 180 days or more during the Report Period. Medications must not have a comment of RETURNED TO STOCK; *or*
- 2. A refusal of the medication during the Report Period; or
- 3. Have a contraindication/previous adverse reaction to the indicated medication

Refusals and contraindications/previous ADR/allergies are only counted if a patient did not have a prescription for the indicated medication. Patients without a prescription who have a refusal and/or contraindication/ADR/ allergy will be counted in sub-numerators B–C. Because a patient may have both a refusal and a contraindication/ADR/allergy, the sub-numerator totals of A–C may not add up the to the numerator total.

For prescriptions, the days supply requirement may be met with a single prescription or from a combination of prescriptions for the indicated medication that were filled during the Report Period and prescriptions filled prior to the Report Period but which are still active (i.e., prior active prescription). Prior active prescriptions can be included if the treatment days fall within the Report Period. Prior active prescription defined as most recent prescription for the indicated medication (see codes below) prior to Report Period Start Date with the number of days supply equal to or greater than the Report Period Start Date minus the prescription date.

**Note:** If a prescription for a medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Example of patient included in the beta-blocker numerator with prior active prescription:

- Report Period: 07/01/2010–06/30/2011
- Must have 180 days supply of indicated medication 6/30/2011 (end of Report Period)
- Prior Beta-Blocker Rx Date: 06/01/2010
- # Days Prescribed: 60 (treats patient through 07/31/2010)
- Report Period Start Date minus Rx Date: 07/01/2010-06/01/2010 = 30; 60 (#Days Prescribed) is >= 30, prescription is considered Prior Active Rx
- 07/31/2010 is between the Report Period of 07/01/2010 and 06/30/2011, thus remainder of Prior Active Rx can be counted toward 180-days supply

- # Remaining Days Prescribed from Prior Active Rx:
- (# Days Prescribed-(Report Period Start Date-Prior Rx Date) = 60-(07/01/2010-06/01/2010) = 60-30 = 30
- Rx #2: 08/05/2010, # Days Prescribed: 90
- Rx #3: 11/10/2010, #Days Prescribed: 90
- Total Days Supply Prescribed between 07/01/2010 and 06/30/2011, including prior active prescription: 30+90+90=210

# **Beta-Blocker Numerator Logic**

Beta-blocker medication codes defined with medication taxonomy BGP HEDIS BETA BLOCKER MEDS. Medications are: Noncardioselective Beta Blockers: Carteolol, Carvedilol, Labetalol, Nadolol, Penbutolol, Pindolol, Propranolol, Timolol, Sotalol; Cardioselective Beta Blockers: Acebutolol, Atenolol, Betaxolol, Bisoprolol, Metoprolol, Nebivolol; and Antihypertensive Combinations: Atenolol-chlorthalidone, Bendroflumethiazide-nadolol, Bisoprolol-hydrochlorothiazide, Hydrochlorothiazide-metoprolol, and Hydrochlorothiazide-propranolol.

Refusal of beta-blocker: REF refusal of any beta-blocker medication in site-populated medication taxonomy BGP HEDIS BETA BLOCKER MEDS at least once during the Report Period.

CRS uses the following codes to define contraindications to beta-blockers.

Contraindication to Beta- Blockers (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
Asthma		<b>POV</b> : 2 diagnoses of 493* on different visit dates
Hypotension		POV: 1 diagnosis of 458*
Heart Block >1 Degree		<b>POV</b> : 1 diagnosis of 426.0, 426.12, 426.13, 426.2, 426.3, 426.4, 426.51, 426.52, 426.53, 426.54, or 426.7
Sinus Bradycardia		POV: 1 diagnosis of 427.81
COPD		<b>POV</b> : 2 diagnoses on different visit dates of 491.2*, 496, or 506.4, or a combination of any of these codes, such as 1 visit with 491.20 and 1 with 496
NMI Refusal	G8011 (old code) at least once during the Report Period	Refusal: NMI refusal for any beta-blocker at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to beta-blockers.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to Beta-Blockers (any of the codes occurring ever)	<b>POV</b> : 995.0-995.3 AND E942.0
	Entry in ART (Patient Allergies File): "beta block*"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "beta block*",
	"bblock*" or "b block*"

# ASA (aspirin)/Other Anti-Platelet Numerator Logic

ASA medication codes defined with medication taxonomy DM AUDIT ASPIRIN DRUGS.

Other anti-platelet medication codes defined with medication taxonomy site-populated BGP ANTI-PLATELET DRUGS taxonomy.

Refusal of ASA/other anti-platelet: REF refusal of any ASA or anti-platelet medication in site-populated medication taxonomies DM AUDIT ASPIRIN DRUGS or BGP ANTI-PLATELET DRUGS at least once during the Report Period.

CRS uses the following codes to define contraindications to ASA/other anti-platelets.

Contraindication to ASA/Other Anti-Platelets (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
180-day course of treatment for Warfarin/Coumadin during the Report Period		Site-Populated Drug Taxonomy: BGP CMS WARFARIN MEDS
Hemorrhage		<b>POV</b> : 459.0
NMI Refusal	G8008 (old code) at least once during the Report Period	Refusal: NMI (not medically indicated) refusal for any aspirin at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to ASA/other anti-platelets.

	ICD and Other Codes
Adverse Drug Reaction/Allergy to	<b>POV</b> : 995.0-995.3 AND E935.3
(any of the codes occurring ever)	Entry in ART (Patient Allergies File): "aspirin"
	Entry in Problem List or in Provider Narrative for
	any POV 995.0-995.3 or V14.8: "ASA" or "aspirin"

# **ACEI/ARB Numerator Logic**

Ace Inhibitor (ACEI) medication codes defined with medication taxonomy BGP HEDIS ACEI MEDS. ACEI medications are: Angiotensin Converting Enzyme Inhibitors (Benazepril, Captopril, Enalapril, Fosinopril, Lisinopril, Moexipril, Perindopril, Quinapril, Ramipril, Trandolopril.)

Antihypertensive Combinations: (Amlodipine-benazepril, Benazepril-hydrochlorothiazide, Captopril-hydrochlorothiazide, Enalapril-hydrochlorothiazide, Fosinopril-hydrochlorothiazide, Hydrochlorothiazide-lisinopril, Hydrochlorothiazide-moexipril, Hydrochlorothiazide-quinapril, Trandolapril-verapamil.)

*Refusal of ACEI:* REF refusal of any ACE Inhibitor medication in site-populated medication taxonomy BGP HEDIS ACEI MEDS at least during the Report Period.

CRS uses the following codes to define contraindications to ACE inhibitors.

Contraindication to ACE Inhibitors (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy	<b>Abortion:</b> Any of the following: 59100, 59120, 59130, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260-S2267	POV or Problem List: At least two visits during the Report Period with V22.0–V23.9, V72.42, 640.* –649.*, or 651.* –676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.  Abortion: Any of the following POVs: 635*, 636*, or 637*  Procedures: 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period

Contraindication to ACE Inhibitors (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI refusal for any ACE inhibitor at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to ACE inhibitors.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy	<b>POV</b> : 995.0-995.3 AND E942.6
to ACE Inhibitors (any of the codes occurring anytime through the end of the Report Period)	Entry in ART (Patient Allergies File): "ace inhibitor" or "ACEI"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "ace i*" or "ACEI"

ARB (Angiotensin Receptor Blocker) medication codes defined with medication taxonomy BGP HEDIS ARB MEDS. ARB medications are: Angiotensin II Inhibitors (Candesartan, Eprosartan, Irbesartan, Losartan, Olmesartan, Telmisartan, Valsartan.)

Antihypertensive Combinations (Aliskiren-valsartan, Amlodipine-hydrochlorothiazide-valsartan, Amlodipine-olmesartan, Candesartan-hydrochlorothiazide, Eprosartan-hydrochlorothiazide, Hydrochlorothiazide-Irbesartan, Hydrochlorothiazide-Losartan, Hydrochlorothiazide-olmesartan, Hydrochlorothiazide-Valsartan.)

*Refusal of ARB:* REF refusal of any ARB medication in site-populated medication taxonomy BGP HEDIS ARB MEDS at least once during the Report Period.

CRS uses the following codes to define contraindications to ARBs.

Contraindication to ARBs (any of the codes occurring ever unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy		POV or Problem List: At least two visits during the Report Period with V22.0–V23.9, V72.42, 640.* –649.*, or 651.* –676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.
	Abortion: Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59856, 59857, S2260-S2267	<b>Abortion:</b> Any of the following POVs: 635*, 636*, or 637* <b>Procedures</b> : 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage:</b> Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage:</b> Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period
Moderate or Severe Aortic Stenosis		<b>POV:</b> 395.0, 395.2, 396.0, 396.2, 396.8, 424.1, 425.1, or 747.22
NMI Refusal		Refusal: NMI (not medically indicated) refusal for any ARB at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to ARBs.

	ICD and Other Codes
Adverse Drug Reaction/ Allergy to ARBs (any of the codes occurring anytime through the end of the Report Period)	<b>POV</b> : 995.0-995.3 AND E942.6
	Entry in ART (Patient Allergies File): "Angiotensin Receptor Blocker" or "ARB"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "Angiotensin Receptor Blocker" or "ARB"

# **Statins Numerator Logic**

Statin medication codes defined with medication taxonomy BGP HEDIS STATIN MEDS. Statin medications are: Atorvostatin (Lipitor), Fluvastatin (Lescol), Lovastatin (Altocor), Mevacor, Pravastatin (Pravachol), Simvastatin (Zocor), Rosuvastatin (Crestor).

Statin Combination Products: Advicor, Caduet, PraviGard Pac, Vytori.

*Refusal of Statin*: REF refusal of any statin medication in site-populated medication taxonomy BGP HEDIS STATIN MEDS at least once during the Report Period.

CRS uses the following codes to define contraindications to statins.

Contraindication to ACE Inhibitors (any of the codes occurring ever, unless otherwise noted)	CPT Codes	ICD and Other Codes
Pregnancy	<b>Abortion</b> : Any of the following: 59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260-S2267	POV or Problem List: At least two visits during the Report Period with V22.0–V23.9, V72.42, 640.* –649.*, 651.* –676.* and with no documented miscarriage or abortion (defined below) occurring after the second pregnancy POV. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the Report Period, CRS will use the first two visits in the Report Period.  Abortion: Any of the following POVs: 635*, 636*, or 637*  Procedure: 69.01, 69.51, 74.91, 96.49
	<b>Miscarriage</b> : Any of the following: 59812, 59820, 59821, 59830	<b>Miscarriage</b> : Any of the following POVs: 630, 631, 632, 633*, or 634*
Breastfeeding		POV: V24.1 during the Report Period Patient Education: BF-BC, BF-BP, BF-CS, BF-EQ, BF-FU, BF-HC, BF-ON, BF-M, BF- MK, or BF-N during the Report Period
Acute Alcoholic Hepatitis		POV: 571.1 during the Report Period
NMI Refusal		Refusal: NMI refusal for any statin at least once during the Report Period

CRS uses the following codes to define adverse drug reactions/documented allergies to statins.

	ICD and Other Codes
Adverse Drug Reaction/Allergy to Statins (any of the codes occurring anytime through the end of the Report Period, unless otherwise noted)	Site-Populated Lab Taxonomy or LOINC Taxonomy: DM AUDIT ALT TAX, with ALT and/or AST > 3x the Upper Limit of Normal (ULN) (i.e., Reference High) on 2 or more consecutive visits during the Report Period
	Site-Populated Lab Taxonomy or LOINC Taxonomy: BGP CREATINE KINASE TAX, with Creatine Kinase (CK) levels > 10x ULN or CK > 10,000 IU/L during the Report Period
	<b>POV</b> : Myopathy/Myalgia, defined as any of the following during the Report Period: 359.0-359.9, 729.1, 710.5, or 074.1
	<b>POV</b> : 995.0-995.3 AND E942.9
	Entry in ART (Patient Allergies File): "statin" or "statins"
	Entry in Problem List or in Provider Narrative for any POV 995.0-995.3 or V14.8: "Statin" or "Statins"

# **All Medications Numerator Logic**

To be included in this numerator, a patient must have a prescription, refusal, or a contraindication for *all* of the four medication classes (i.e., beta-blocker, ASA/other anti-platelet, ACEI/ARB, *and* statin).

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of IHD patients 22+ with 180-day medication therapy during the Report Period, if any.

#### **Measure Source**

American Heart Association/American College of Cardiology Guidelines

# **Measure Past Performance and Long-Term Targets**

None

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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000
```

Appropriate Medicati	on Thera	py in	High Risk	Patie	ents				
	REPORT PERIOD	%	PREV YR PERIOD	%	CHG from PREV YR %		ફ	CHG from BASE %	
Active IHD pts 22+	59		38			30			
# w/180 day beta-blo Rx/contra/ADR	cker								
-No Refusals	32	54.2	22	57.9	-3.7	14	46.7	+7.6	
A. # w/180 day beta-									
Rx w/% of Total B: # w/contra/ADR	18	56.3	14	63.6	-7.4	11	78.6	-22.3	
w/ % of Total # w/ beta-blocker	14	43.8	8	36.4	+7.4	3	21.4	+22.3	
refusal	1	1.7	0	0.0	+1.7	0	0.0	+1.7	
# w/180 day ASA Rx/contra/ADR									
-No Refusals A. # w/180 day ASA	22	37.3	21	55.3	-18.0	23	76.7	-39.4	
Rx w/% of Total B. # w/contra/ADR	17	77.3	18	85.7	-8.4	18	78.3	-1.0	
w/ % of Total	5	22.7	3	14.3	+8.4	5	21.7	+1.0	
# w/ ASA refusal	1	1.7	0	0.0	+1.7	0	0.0	+1.7	
# w/180 day ACEI/ARB Rx/contra/ADR									
-No Refusals A. # w/180 day ACEI/		49.2	15	39.5	+9.7	18	60.0	-10.8	
Rx w/% of Total B. # w/contra/ADR	26	89.7	14	93.3	-3.7	17	94.4	-4.8	
w/ % of Total # w/ ACEI/ARB	3	10.3	1	6.7	+3.7	1	5.6	+4.8	
refusal	1	1.7	0	0.0	+1.7	0	0.0	+1.7	
# w/180 day statin Rx/contra/ADR									
-No Refusals A. # w/180 day stati	30	50.8	22	57.9	-7.0	14	46.7	+4.2	
Rx w/% of Total B. # w/contra/ADR	26	86.7	20	90.9	-4.2	14	100.0	-13.3	
w/ % of Total	4	13.3	2	9.1	+4.2	0	0.0	+13.3	
# w/ Statin refusal	<del>-</del>	1.7	0		+1.7	0	0.0	+1.7	
# w/180 day Rx/contr of ALL meds	a/ADR/								
-No Refusals	16	27.1	8	21.1	+6.1	6	20.0	+7.1	

Figure 2-88: Sample Report, Appropriate Medication Therapy in High Risk Patients

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Appropriate Medication Therapy in High Risk Patients: List of IHD patients 22+ with 180-day medication therapy during the Report Period, if any.

CRS Clinical Performance Measure Logic Manual for FY 2012 Clinical Measures
December 2011

```
PATIENT NAME HRN COMMUNITY SEX AGE

DENOMINATOR NUMERATOR

PATIENT1, CAITLYN 000001 COMMUNITY #1 F 22

IHD BETA: 09/11/12 Refused METOPROLOL TARTRATE 50MG TAB;
ASA: 09/11/12 Refused ASPIRIN 325MG E.C. TAB; ACEI/ARB: 09/11/12 Refused CAPTOPRIL
25MG TABS; STATIN: 09/11/12 Refused ATORVASTATIN 10MG TAB (ICP)
PATIENT2, CARLA 000002 COMMUNITY #1 F 40

IHD
PATIENT3, GENEVA 000003 COMMUNITY #1 F 47

JACKSON, SHERRY LADAWN 100939 BRAGGS F 68

IHD, AD ACEI/ARB: (268 TOTAL DAYS); STATIN: (319 TOTAL DAYS)
PATIENT4, SHERRY LISA 000004 COMMUNITY #1 F 68

IHD BETA: 09/12/12 Contra hypotension POV 458.9; ASA:
aspirin Contra total days WARFARIN: 372; STATIN: (328 TOTAL DAYS)
PATIENT5, PAULINE 000005 COMMUNITY #1 F 70

IHD, AD ALL MEDS; BETA: (305 TOTAL DAYS); ASA: (291 TOTAL
DAYS); ACEI/ARB: (405 TOTAL DAYS); STATIN: (305 TOTAL DAYS)
```

Figure 2-89: Sample Patient List: Appropriate Medication Therapy in High Risk Patients

# 2.8.13 Stroke and Stroke Rehabilitation: Anticoagulant Therapy Prescribed for Atrial Fibrillation at Discharge

#### Denominator

Number of visits for *User Population patients* ages 18 and older who were hospitalized during the Report Period with ischemic stroke or transient ischemic attack (TIA) with documented permanent, persistent, or paroxysmal atrial fibrillation.

## **Numerators**

Number of visits where patients received a prescription for anticoagulant at discharge

Number of visits where patients refused anticoagulant therapy

Number of visits where patients did not receive anticoagulation therapy

## **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

CRS uses the following codes to define ischemic stroke or transient ischemic attack with atrial fibrillation.

	ICD and Other Codes
Ischemic Stroke or TIA with Atrial Fibrillation (Non-CHS inpatient visit - Type not equal to C and Service Category=H)  The patient must be admitted to the hospital during the report period with a condition described here but the discharge may occur after the report period.	<b>V POV:</b> 433.01, 433.11, 433.21, 433.31, 433.81, 433.91, 434.01, 434.11, 434.91, 435.0, 435.1, 435.2, 435.3, 435.8, or 435.9 AND 427.31 (atrial fibrillation)

Anticoagulant Therapy: Patient must meet one of the conditions below to be counted as receiving anticoagulant therapy. For all prescriptions, medications must not have a comment of RETURNED TO STOCK.

1. Active prescription for Warfarin, aspirin, or other anti-platelet as of discharge date. "Active" prescription defined as:

Rx Days Supply >= (Discharge Date - Prescription Date), where the prescription has not been discontinued as of the discharge date.

2. Prescription for Warfarin, aspirin, or other anti-platelet on discharge date.

*Warfarin Medication:* Any medication in site-populated BGP CMS WARFARIN MEDS taxonomy.

Aspirin Medication: Any medication in site-populated DM AUDIT ASPIRIN DRUGS taxonomy.

Other Anti-Platelet/Anticoagulant Medication: Any medication in the site-populated BGP ANTI-PLATELET DRUGS taxonomy, any medication with VA Drug Class BL700.

*Refusal of Anticoagulant Therapy*: Refusal of any of the following documented on discharge date:

- 1. Any medication in site-populated taxonomies BGP CMS WARFARIN MEDS, DM AUDIT ASPIRIN DRUGS, or BGP ANTI-PLATELET DRUGS; or
- 2. Any medication with VA Drug Class BL700.

*No Anticoagulant Therapy:* Patients who did not have an active prescription for anticoagulant therapy at time of discharge and did not receive or refuse anticoagulant therapy at discharge.

## **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of patients with stroke/TIA and atrial fibrillation with anticoagulant therapy, if any.

#### **Measure Source**

None

## **Measure Past Performance and Long-Term Targets**

None

DU November 25, 2012 Page 248  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Stroke and Stroke Rehabilitation: Anticoagulant Therapy Prescribed for Atrial Fibrillation at Discharge									
	REPORT PERIOD				CHG from BA PREV YR % PE				
# Stroke/TIA w/ At Fib Visits for Use Pop Pts 18+			0			0			
# Visits w/ Anticoagulant Rx # Visits w/	5 2	9.4	0	0.0	+29.4	0	0.0	+29.4	
Refusal # Visits w/ No Anticoagulant Therapy	1 11 6	<ol> <li>5.9</li> <li>4.7</li> </ol>			+5.9		0.0	+5.9 +64.7	

Figure 2-90: Sample Report, Stroke and Stroke Rehabilitation: Anticoagulant Therapy Prescribed for Atrial Fibrillation at Discharge

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Stroke and Stroke Rehabilitation: Anticoagulant Therapy Prescribed for Atrial Fibrillation at Discharge: List of patients with stroke/TIA and atrial fibrillation with anticoagulant therapy, if any.

PATIENT NAME HRN COMMUNITY SEX AGE

Figure 2-91: Sample Patient List: Stroke and Stroke Rehabilitation: Anticoagulant Therapy Prescribed for Atrial Fibrillation at Discharge

# 2.8.14 Cholesterol Management for Patients with Cardiovascular Conditions

#### **Denominators**

Active Clinical patients ages 18 to 75 who, during the first ten months of the year prior to the beginning of the Report Period, were diagnosed with AMI, coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) or who were diagnosed with IVD during the Report Period and the year prior to the Report Period. Broken down by gender.

*User Population patients* ages 18 to 75 who, during the first 10 months of the year prior to the beginning of the Report period, were diagnosed with AMI, coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) *or* who were diagnosed with IVD during the Report Period and the year prior to the Report Period. Broken down by gender.

#### **Numerators**

Patients with LDL completed during the Report Period, regardless of result.

- a. Patients with LDL <=100, completed during the Report Period.
- b. Patients with LDL 101-130, completed during the Report Period.
- c. Patients with LDL >130, completed during the Report Period.

## **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

Searches for most recent LDL test with a result during the Report Period. If more than one LDL test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If an LDL test with a result is not found, CRS searches for the most recent LDL test without a result. For numerator LDL =<100, CPT 3048F will count as meeting the measure.

CRS uses the following codes to define the denominator and numerators.

Diagnosis or Test	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
AMI		V POV: 410.*0, 410.*1		
PCI	92980, 92982, 92995, G0290	V Procedure: 00.66, 36.01 (old code), 36.02 (old code), 36.05, (old code), or 36.06- 36.07		
CABG	33510-33514, 33516–33519, 33521–33523, 33533–33536, S2205-S2209	V POV: V45.81 V Procedure: 36.1*, 36.2		
IVD		V POV: 411.*, 413.*, 414.0*, 414.2, 414.8, 414.9, 429.2, 433.*-434.*, 440.1, 440.2*, 440.4, 444.*, or 445.*		
LDL	80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F		Yes	DM AUDIT LDL CHOLESTEROL TAX

## **Key Logic Changes from CRS Version 11.1**

- Added POV code V45.81 to CABG definition.
- Added POV code V45.82 to PCI definition.
- Removed Procedure code 36.09 from PCI definition.
- Removed CPT code 33140 from PCI definition.

#### **Patient List Description**

List of patients with AMI, CABG, PCI, or IVD w/LDL value, if any.

## **Measure Source**

**HEDIS** 

# **Measure Past Performance and Long-Term Targets**

None

Previo	eport Per ous Year	Measu DEMC iod: J Period	INDIAN Han 01, 20	Commun OSPITA 12 to , 2011	AL Dec 31, 20 1 to Dec 33	012 1, 2011		age 250 **	
	Baseline Period: Jan 01, 2000 to Dec 31, 2000								
	REPORT PERIOD				CHG from PREV YR %				
Active Clinical pts with DX of AMI, CABO PCI, or IVD			28			18			
# w/LDL done A. # w/LDL <=100	38	90.5	21	75.0	+15.5	9	50.0	+40.5	
<pre>w/% of Total Screened B. # w/LDL 101-130 w/% of Total</pre>	14	36.8	12	57.1	-20.3	3	33.3	+3.5	
Screened C. # w/LDL >130 w/% of Total	5	13.2	3	14.3	-1.1	2	22.2	-9.1	
Screened		13.2	4	19.0	-5.9	4	44.4	-31.3	
Male Active Clinical 18-75 with DX of AMI PCI, or IVD	-		16			10			
# w/LDL done A. # w/LDL <=100	20	90.9	11	68.8	+22.2	4	40.0	+50.9	
<pre>w/% of Total Screened B. # w/LDL 101-130 w/% of Total</pre>	5	25.0	4	36.4	-11.4	1	25.0	+0.0	
<pre>w/% of Total Screened C. # w/LDL &gt;130 w/% of Total</pre>	2	10.0	3	27.3	-17.3	1	25.0	-15.0	
Screened	4	20.0	2	18.2	+1.8	2	50.0	-30.0	
Female Active Clinic 18-75 with DX of AMI PCI, or IVD			12			8			
# w/LDL done A. # w/LDL <=100	18	90.0	10	83.3	+6.7	5	62.5	+27.5	
<pre>w/% of Total Screened B. # w/LDL 101-130 w/% of Total</pre>	9	50.0	8	80.0	-30.0	2	40.0	+10.0	
w/% of Total Screened	3	16.7	0	0.0	+16.7	1	20.0	-3.3	

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```
C. # w/LDL >130

w/% of Total

Screened 1 5.6 2 20.0 -14.4 2 40.0 -34.4
```

Figure 2-92: Sample Report, Cholesterol Management for Patients with Cardiovascular Conditions

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Cholesterol Management for Patients with Cardiovascular Conditions: List of patients with AMI, CABG, PCI, or IVD w/LDL value, if any. PATIENT NAME HRN COMMUNITY DENOMINATOR NUMERATOR PATIENT1, SHERRY 000001 COMMUNITY #1 F 68 UP,AC 07/15/12 LDL: CPT 304 PATIENT2,CODY JACK 000002 COMMUNITY #1 M 41 07/15/12 LDL: CPT 3048F UP,AC PATIENT3, TIMOTHY ALLEN 000003 COMMUNITY #1 M 43 UP,AC 10/17/12 LDL: 136
PATIENT4,TRACE 000004 COMMUNITY #1 M 47 10/17/12 LDL: 136 PATIENT5, KENNETH 000005 COMMUNITY #1 M 60 08/06/12 LDL: 108 UP,AC

Figure 2-93: Sample Patient List: Cholesterol Management for Patients with Cardiovascular Conditions

#### 2.8.15 Heart Failure and Evaluation of LVS Function

#### **Denominator**

Active Clinical patients ages 18 or older discharged with heart failure during the Report Period.

#### **Numerators**

Patients whose LVS function was evaluated before arrival, during hospitalization, or is planned for after discharge.

#### **Logic Description**

Age of the patient is calculated as of the hospital admission date.

Denominator exclusions are defined as any of the following:

- 1. Patients receiving comfort measures only (i.e., patients who received palliative care and usual interventions were not received because a medical decision was made to limit care).
- 2. Patients with a Discharge Type of Transferred or Irregular or containing "Death."

3. Patients who had a left ventricular assistive device (LVAD) or heart transplant procedure during hospitalization.

CRS uses the following codes to define the denominator and numerators.

#### **Denominator Exclusions**

	CPT Codes	ICD and Other Codes
Comfort Measures		V POV: V66.7 (Encounter for palliative care) documented during hospital stay
LVAD/Heart Transplant		<b>V Procedure</b> : 33.6, 37.41, 37.51–37.54, 37.61–37.66, 37.68 documented during hospital stay

#### **Denominator Definition**

	CPT Codes	ICD and Other Codes
Heart Failure		V POV (Primary Diagnosis only): 398.91, 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, 428.0, 428.1, 428.20, 428.21, 428.22, 428.23, 428.30, 428.31, 428.32, 428.33, 428.40, 428.41, 428.42, 428.43, 428.9, 429.1, or 997.1 and with Service Category H (hospitalization). NOTE: If a patient has multiple admissions matching these criteria during the Report Period, the earliest admission will be used.

# Numerator Definition (Evaluation of LVS Function): Any of the codes listed below

	CPT Codes	ICD and Other Codes
Ejection Fraction (ordered or documented anytime one year prior to discharge date)	78414, 78468, 78472, 78473, 78480, 78481, 78483, 78494, 93303, 93304, 93307, 93308, 93312, 93314–93318, 93350, 93543, 93555	V Measurement: "CEF" V Procedure: 88.53, 88.54
RCIS Order for Cardiovascular Disorders Referral (ordered during the hospital stay but no later than the hospital discharge date)		ICD Diagnostic Category: "Cardiovascular Disorders" AND one of the following: CPT Categories: "Evaluation and/or Management," "Non-surgical Procedures" or "Diagnostic Imaging"

	CPT Codes	ICD and Other Codes
Other Procedures (documented anytime one year prior to discharge date)		Echocardiogram: V Procedure 88.72, 37.28, 00.24; Nuclear Medicine Test: V Procedure 92.2*; Cardiac Catheterization with a Left Ventriculogram: V Procedure 37.22, 37.23, 88.53, 88.54

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of Active Clinical heart failure patients 18+ who received evaluation of LVS function, if any.

## **Measure Source**

CMS HF-2

## **Measure Past Performance and Long-Term Targets**

None

DU November 25, 2012 Page 255  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Heart Failure and Evaluation of LVS Function  REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %									
AC 18+ w/Heart Failure Dx	44		2			1			
Patients w/Eval of LVS Function	14	31.8	0 (	0.0	+31.8	0	0.0	+31.8	

Figure 2-94: Sample Report, Heart Failure and Evaluation of LVS Function

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Heart Failure and Evaluation of LVS Function: List of Active Clinical heart failure patients 18+ who received evaluation of LVS function, if any.

PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR
PATIENT1, JOAN AC PATIENT2, SARAH AC PATIENT3, JOHN AC PATIENT4, ROGER AC PATIENT5, DANIEL	000164 COMMUNITY #1 F 36     Admission: 06/01/12 LVS: NOT DOCUMENTED 000127 COMMUNITY #1 F 35     Admission: 06/01/12 LVS: 06/03/11 Proc 88.72 000151 COMMUNITY #1 M 36     Admission: 05/01/12 LVS: 05/01/12 Meas CEF 40 000125 COMMUNITY #1 M 47     Admission: 06/01/12 LVS: NOT DOCUMENTED 000129 COMMUNITY #1 M 57
AC	Admission: 06/01/12 LVS: 06/01/12 CPT 78468

Figure 2-95: Sample Patient List: Heart Failure and Evaluation of LVS Function

# 2.9 STD-Related Measure Topics

# 2.9.1 HIV Screening

## **GPRA Measure Description**

During FY 2012, achieve the tentative target rate of 81.8% for the proportion of pregnant patients who are screened for HIV.

#### **Denominators**

All *pregnant Active Clinical female User Population patients* with no documented miscarriage or abortion and with no recorded HIV diagnosis ever (GPRA Denominator).

*User Population patients ages 13–64 with no recorded HIV* diagnosis prior to the Report Period. (GPRA Developmental Denominator).

#### **Numerators**

Patients who were screened for HIV during the past 20 months.

**Note**: This numerator does *not* include refusals. (GPRA Numerator).

Patients with documented HIV screening refusal during the past 20 months.

Patients who were screened for HIV during the Report Period.

**Note**: This numerator does *not* include refusals. (GPRA Developmental Numerator)

Patients with documented HIV screening refusal during the Report Period.

No denominator. This measure is a total count only, not a percentage. Number of HIV screens provided to User Population patients during the report period, where the patient was not diagnosed with HIV any time prior to the screen.

**Note**: This numerator does not include refusals. (GPRA Developmental Numerator)

## **Logic Description**

Age of the patient is calculated at the beginning of the Report period.

Pregnancy definition: At least two visits during the past 20 months from the end of the Report Period. Pharmacy-only visits (Clinic Code 39) will not count toward these two visits. If the patient has more than two pregnancy-related visits during the past 20 months, CRS will use the first two visits in the 20-month period. The patient must not have a documented miscarriage or abortion occurring after the second pregnancy-related visit. In addition, the patient must have at least one pregnancy-related visit occurring during the reporting period. The time period is extended to include patients who were pregnant during the Report Period but whose initial diagnosis (and HIV test) were documented prior to Report Period.

*HIV Screening definition*: For the number of HIV screens provided to User Population patients numerator (count only), a maximum of one HIV screen per patient per day will be counted.

- **Note 1:** The time frame for both screening and refusals for the pregnant patients denominator is anytime during the past 20 months and for User Population patients 13–64 is anytime during the Report Period.
- Note 2: Refusals are allowed during the past 20 months for pregnant patients (vs. only during the Report Period) in the event the patient is at the end of her pregnancy at the beginning of the Report Period and refused the HIV test earlier in her pregnancy during the previous year

CRS uses the following codes and taxonomies to define the denominator and numerators.

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Pregnancy (at least 2 visits in past 20 months with 1 during the Report Period)		<b>V POV</b> : V22.0–V23.9, V72.42, 640.* –649.*, 651.* – 676.*		
Miscarriage (after second pregnancy POV in past 20 months)	59812, 59820, 59821, 59830	<b>V POV</b> : 630, 631, 632, 633*, 634*		
Abortion (after second pregnancy POV in past 20 months)	59100, 59120, 59130, 59136, 59150, 59151, 59840, 59841, 59850, 59851, 59852, 59855, 59856, 59857, S2260—S2267	V POV: 635*, 636* 637* V Procedure: 69.01, 69.51, 74.91, 96.49.		
HIV Diagnosis (documented anytime prior to the end of the Report Period)		V POV or Problem List: 042, 042.0–044.9 (old codes), 079.53, V08, or 795.71		
HIV Screening	86689, 86701- 86703, 87390, 87391, 87534- 87539		Yes	BGP HIV TEST TAX
Refusal of HIV lab test in past 20 months				BGP HIV TEST TAX

# **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of pregnant patients or User Population patients with documented HIV test or refusal, if any.

#### **Measure Source**

HP 2020 HIV-14.3

# **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS FY 2011 Performance	80.0%
IHS FY 2010 Performance	78.0%
IHS FY 2009 Performance	76.0%
IHS FY 2008 Performance	75.0%
IHS FY 2007 Performance	74.0%
IHS FY 2006 Performance	65.0%
IHS FY 2005 Performance	54.0%
HP 2020 Goal	74.1%

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Report Period: Jan 01, 2012 to Dec 31, 2012									
	Previous Year Period: Jan 01, 2011 to Dec 31, 2011								
Ba	aseline Pe	riod:	Jan 01,	2000	to Dec 31,	2000			
HIV Screening									
	REPORT	%	PREV YR	%	CHG from	BASE	% (	CHG from	
	PERIOD		PERIOD		PREV YR %	PERIOD	Ε	BASE %	
Pregnant AC Pts w/	no								
HIV ever (GPRA)	39		38			32			
# w/HIV screening									
-No Refusals									
(GPRA)	15	38.5	6	15.8	+22.7	0	0.0	+38.5	
<pre># w/HIV screening   refusal</pre>	1	2.6	0	0.0	+2.6	0	0.0	+2.6	
User Pop Pts 13-64 w/ no HIV									
(GPRA Dev.)	2,024		1,663			1,519			
	,		·			,			
# w/HIV screening -No Refusals									
(GPRA Dev.)	46	2.3	21	1.3	+1.0	0	0.0	+2.3	
# w/HIV screening									
refusal	4	0.2	0	0.0	+0.2	0	0.0	+0.2	
# HIV screens for T	Jser Pop P	ts							
w/ no prior HIV-No									
(GPRA Dev.)	51		21		+30	0		+51	

Figure 2-96: Sample Report, HIV Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

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```
HIV Screening: List of pregnant patients or User Population patients with documented HIV test or refusal, if any.

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

PATIENT1, HELEN MARY 000001 COMMUNITY #1 F 12

UP 03/31/12 Lab; Screen Count: 1

PATIENT2, CECELIA 000002 COMMUNITY #1 F 19

UP
PATIENT15, BRENDA G 000015 COMMUNITY #2 F 30

UP 03/14/12 CPT 87534; 02/14/12 CPT 86689; Screen Count: 2

PATIENT16, ALYSHA 000016 COMMUNITY #2 F 33

UP, AC PREG 08/25/12 Lab; Screen Count: 1
```

Figure 2-97: Sample Patient List, HIV Screening

## 2.9.2 HIV Quality of Care

#### **Denominator**

All User Population patients ages 13 and older with at least two direct care visits (i.e., not Contract/CHS) with HIV diagnosis during the Report Period, including one HIV diagnosis in last six months.

#### **Numerators**

Patients who received CD4 test only (without HIV viral load) during the Report Period

Patients who received HIV viral load only (without CD4) during the Report Period

Patients who received both CD4 and HIV viral load during the Report Period

Total patients receiving any test

#### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

CRS uses the following codes and taxonomies to define the denominator and numerators.

	ICPI COMPS		LOINC Codes	Taxonomy
HIV		042, 042.0-044.9		
		(old codes), 079.53,		
		V08, 795.71		

	CPT COdes	LOINC Codes	Taxonomy
CD4	86359, 86360 86361	Yes	BGP CD4 TAX
HIV Viral Load	87536, 87539	Yes	BGP HIV VIRAL LOAD TAX

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients 13 and older diagnosed with HIV, with CD4 test, if any.

#### **Measure Source**

HP 2010 developmental measure 13-13a Viral Load Testing

## **Measure Past Performance and Long-Term Targets**

Performance	Percent
IHS 2020 goal for viral load testing	Nearly 100%
IHS 2020 baseline for CD4 testing	Nearly 100%

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HIV Quality of Care									
	REPORT PERIOD	%	PREV YR PERIOD		CHG from PREV YR % PREV YR			CHG from BASE %	
User Pop Pts >13 w/ HIV Dx	6		1			2			
# w/CD4 only # w/viral load	1	16.7	0	0.0	+16.7	0	0.0	+16.7	
only	1	16.7	0	0.0	+16.7	0	0.0	+16.7	
# w/both TOTAL # w/	1	16.7	1	100.0	-83.3	2	100.0	-83.3	
any tests	3	50.0	1	100.0	-50.0	2	100.0	-50.0	

Figure 2-98: Sample Report HIV Quality of Care

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Figure 2-99: Sample Patient List, HIV Quality of Care

# 2.9.3 Chlamydia Screening

#### **Denominators**

Female Active Clinical patients ages 16 through 25.

- a. Female Active Clinical 16-20.
- b. Female Active Clinical 21–25.

Female User Population patients ages 16 through 25.

- a. Female User Population 16–20.
- b. Female User Population 21–25.

#### **Numerator**

Patients tested for Chlamydia during the Report Period.

#### **Logic Description**

Age is calculated at beginning of the Report Period. The following codes are used to determine a test for Chlamydia.

	ICPT CORPS		LOINC Codes	Taxonomy
Chlamydia Test	86631, 86632, 87110, 87270, 87320, 87490-92, 87810, 3511F	<b>V POV</b> : V73.88, V73.98	Yes	BGP CHLAMYDIA TESTS TAX

#### **Key Logic Changes from CRS Version 11.1**

None

# **Patient List Description**

List of patients with documented Chlamydia screening, if any.

#### **Measure Source**

HP 2020 STD-4, annual screening for genital Chlamydia—females enrolled in commercial MCOs (aged 25 years and under); STD-3, annual screening for genital Chlamydia—females enrolled in Medicaid MCOs (aged 25 years and under).

## **Measure Past Performance and Long-Term Targets**

Performance	Percent
HP 2020 goal for Females 16-20 with Medicaid (STD-3.1)	57.9%
HP 2020 goal for Females 21-24 with Medicaid (STD-3.2)	65.3%
HP 2020 goal for Females 16-20 with Commercial Health Insurance (STD-4.1)	44.1%
HP 2020 goal for Females 21-24 with Commercial Health Insurance (STD-4.2)	47.9%

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Chlamydia Testing						2000		
Ciliamydia lesting								
	PORT	%			CHG from PREV YR %			CHG from BASE %
Female Active Clinical 16-25	166		143			128		
# w/Chlamydia Screen	53	31.9	49	34.3	-2.3	43	33.6	-1.7
A. Female Active Clinic 16-20	al 70		55			57		
# w/Chlamydia Screen	24	34.3	16	29.1	+5.2	23	40.4	-6.1
B. Female Active Clinical 21-25 96 88 71								
# w/Chlamydia Screen	29	30.2	33	37.5	-7.3	20	28.2	+2.0
Female User Population 16-25	287		255			239		
# w/Chlamydia Screen	69	24.0	58	22.7	+1.3	51	21.3	+2.7

A. Female User Population 16-20 139 118 119									
# w/Chlamydia Screen	32	23.0	19	16.1	+6.9	25	21.0	+2.0	
B. Female User Population 21-25 148 137 120									
# w/Chlamydia Screen	37	25.0	39	28.5	-3.5	26	21.7	+3.3	

Figure 2-100: Sample Report Chlamydia Testing

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Chlamydia Testing: List of patients with documented Chlamydia screening, if
any.
PATIENT NAME
                      HRN COMMUNITY
                                           SEX AGE
DENOMINATOR
                             NUMERATOR
PATIENT1, MELISSA ANNE 000001 COMMUNITY #1 F 16
UP,AC
PATIENT2, LISA MARIE 000002 COMMUNITY #1 F 16
                             04/04/12 Lab test
UP,AC 04/04/12 Lab test PATIENT3,CRYSTAL LEE 000003 COMMUNITY #1 F 17
UP,AC 07/25/12 Lab test
PATIENT4,DANIELLE 000004 COMMUNITY #1 F 18
UP,AC
                          06/01/12 CPT 87490
PATIENT5, KELLYE 000005 COMMUNITY #1 F 19
UP,AC
```

Figure 2-101: Sample Patient List, Chlamydia Testing

# 2.9.4 Sexually Transmitted Infection Screening

#### **Denominators**

Number of key sexually transmitted infections (STI) incidents for Active Clinical patients that occurred during the period 60 days prior to the beginning of the Report Period through the first 300 days of the Report Period. Key STIs defined as Chlamydia, Gonorrhea, HIV/AIDS, and Syphilis. Two or more key STIs on the same visit will be counted once. Broken down by gender.

Chlamydia screenings needed for key STI incidents for Active Clinical patients that occurred during the defined period. Broken down by gender.

Gonorrhea screenings needed for key STI incidents for Active Clinical patients that occurred during the defined period. Broken down by gender.

HIV/AIDS screenings needed for key STI incidents for Active Clinical patients that occurred during the defined period. Broken down by gender.

*Syphilis screenings* needed for key STI incidents for Active Clinical patients that occurred during the defined period. Broken down by gender.

Number of key sexually transmitted infections (STI) incidents for User Population patients that occurred during the period 60 days prior to the beginning of the Report Period through the first 300 days of the Report Period. Key STIs defined as Chlamydia, gonorrhea, HIV/AIDS, and syphilis. Two or more key STIs on the same visit will be counted once. Broken down by gender.

Chlamydia screenings needed for key STI incidents for User Population patients that occurred during the defined period. Broken down by gender.

Gonorrhea screenings needed for key STI incidents for User Population patients that occurred during the defined period. Broken down by gender.

HIV/AIDS screenings needed for key STI incidents for User Population patients that occurred during the defined period. Broken down by gender.

*Syphilis screenings* needed for key STI incidents for User Population patients that occurred during the defined period. Broken down by gender.

#### **Numerators**

No denominator; count only. The total count of *Active Clinical* patients who were diagnosed with one or more key STIs during the period 60 days prior to the Report Period through the first 300 days of the Report Period. Broken down by gender.

No denominator; count only. The total count of separate *key STI incidents for Active Clinical patients* during the defined period. Broken down by gender.

*Number of complete screenings*, defined as all screenings necessary for a specific STI incident(s), performed from one month prior to the date of relevant STI incident through two months after.

**Note**: This numerator does *not* include refusals.

Number of documented screening refusals.

*Number of needed Chlamydia screenings* performed from one month prior to the date of first STI diagnosis of each incident through two months after.

**Note**: This numerator does *not* include refusals.

Number of documented screening refusals.

*Number of needed Gonorrhea screenings* performed from one month prior to the date of first STI diagnosis of each incident through two months after.

**Note**: This numerator does *not* include refusals.

Number of documented screening refusals.

*Number of needed HIV/AIDS screenings* performed from one month prior to the date of first STI diagnosis of each incident through two months after.

**Note**: This numerator does *not* include refusals.

Number of documented screening refusals.

*Number of needed Syphilis screenings* performed from one month prior to the date of first STI diagnosis of each incident through two months after.

**Note**: This numerator does *not* include refusals.

Number of documented screening refusals.

No denominator; count only. Total count of *User Population patients* who were diagnosed with one or more key STIs during the period 60 days prior to the Report Period through the first 300 days of the Report Period. Broken down by gender.

No denominator; count only. Total count only of *separate key STI incidents for User Population patients* during the defined period. Broken down by gender.

## **Logic Description**

Key STIs are Chlamydia, Gonorrhea, HIV/AIDS, and Syphilis. Key STI diagnoses are defined with the following codes.

	ICD and Other Codes
Chlamydia	<b>V POV</b> : 079.88, 079.98, 099.41, 099.50-099.59
Gonorrhea	V POV: 098.0-098.89
HIV/AIDS	<b>V POV</b> : 042, 042.0-044.9, 079.53, 795.71, V08
Syphilis	<b>V POV</b> : 090.0-093.9, 094.1-097.9

Logic for Identifying Patients Diagnosed with Key STI:

Any patient with one or more diagnoses of any of the key STIs defined above during the period 60 days prior to the beginning of the Report Period through the first 300 days of the Report Period.

Logic for Identifying Separate Incidents of Key STIs:

One patient may have one or multiple occurrences of one or multiple STIs during the year, except for HIV. An occurrence of HIV is only counted if it is the initial HIV diagnosis for the patient ever. Incidents of an STI are identified beginning with the date of the first key STI diagnosis (see definition above) occurring between 60 days prior to the beginning of the Report Period through the first 300 days of the Report Period. A second incident of the same STI (other than HIV) is counted if another diagnosis with the same STI occurs two months or more after the initial diagnosis. A different STI diagnosis that occurs during the same 60-day time period as the first STI counts as a separate incident.

Example of Patient with Multiple Incidents of Single STI:

Visit	Total Incidents
08/01/10: Patient screened for Chlamydia	0
08/08/10: Patient diagnosed with Chlamydia	1
10/15/10: Patient diagnosed with Chlamydia	2
10/25/10: Follow-up for Chlamydia	2
11/15/0: Patient diagnosed with Chlamydia	2
03/01/11: Patient diagnosed with Chlamydia	3

Denominator Logic for Needed Screenings:

One patient may need multiple screening tests based on one or more STI incidents occurring during the time period.

To be included in the needed screening tests denominator, the count will be derived from the number of separate STI incidents and the type(s) of screenings recommended for each incident. The recommended screenings for each key STI are listed below.

STI	Screenings Needed
Chlamydia	Gonorrhea, HIV/AIDS, Syphilis
Gonorrhea	Chlamydia, HIV/AIDS, Syphilis
HIV/AIDS	Chlamydia, Gonorrhea, Syphilis
Syphilis	Chlamydia, Gonorrhea, HIV/AIDS

"Needed" screenings are recommended screenings that are further evaluated for contraindications. The following are reasons that a recommended screening is identified as not needed (i.e., contraindicated).

- 1. The patient has a documented STI diagnosis corresponding to the screening type in the same time period. For example, a patient with both a Chlamydia and a gonorrhea diagnosis on the same visit does not need the recommended Chlamydia screening based on the gonorrhea diagnosis.
- 2. Only one screening for each type of STI is needed during the relevant time period, regardless of the number of different STI incidents identified. For example, if a patient is diagnosed with Chlamydia and Gonorrhea on the same visit, only one screening each is needed for HIV/AIDS and Syphilis.
- 3. A patient with HIV/AIDS diagnosis prior to any STI diagnosis that triggers a recommended HIV/AIDS screening does not need the screening ever.

## Numerator Logic:

To be counted in the numerator, each needed screening in the denominator must have a corresponding laboratory test or test refusal documented in the period from one month prior to the relevant STI diagnosis date through two months after the STI incident.

T.7	COTT	•		1 (* 1	• . •	. 1	C 11	•	1
K OX7	C 1 1	coroonings	Oro	datinad	******	tha	talla	TT7110	CODOC
NEV	'J I I	screenings	415	aermea	wiiii	1110	1()11()	willy	COUCS

	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy	
Chlamydia	86631–86632, 87110, 87270, 87320, 87490– 87492, 87810, 3511F	<b>POV</b> : V73.88, V73.98	Yes	BGP CHLAMYDIA TESTS TAX	
Gonorrhea	87590–87592, 87850, 3511F		Yes	BKM GONORRHEA TEST TAX	
HIV/AIDS	86689, 86701– 86703, 87390– 87391, 87534– 87539		Yes	BGP HIV TEST TAX	
Syphilis	86592–86593, 86781, 87285, 3512F		Yes	BKM FTA-ABS TESTS TAX or BKM RPR TESTS TAX	
Refusal of any screening	Any refusal type (REF, NMI, etc.) for any of the four screening tests as defined above during the specified time period.				

Logic Examples

Example of Patient with Single Diagnosis of Single STI:

08/01/10:	Patient screened for Chlamydia
08/08/10:	Patient diagnosed with Chlamydia–3 screens needed: Gonorrhea, HIV/AIDS, Syphilis
08/13/10:	Patient screened for Gonorrhea, HIV/AIDS, Syphilis
Result:	Result: Denominator: 1 key STI incident, Numerator: 1 complete screening.

# Example of Patient with Multiple Diagnoses of Single STI:

08/01/10:	Patient screened for Chlamydia
08/08/10:	Patient diagnosed with Chlamydia (Incident #1)-3 screens needed: Gonorrhea, HIV/AIDS, Syphilis
12/01/10:	Patient screened for Chlamydia
12/08/10:	Patient diagnosed with Chlamydia (Incident #2) –3 screens needed: Gonorrhea, HIV/AIDS, Syphilis
Result:	Result: Denominator: 2 key STI incidents, Numerator: 1 complete screening (1 each of 3 types)

# Example of Patient with Single Diagnosis of Multiple STIs:

10/15/10:	Patient screened for Chlamydia, Gonorrhea, HIV/AIDS, Syphilis
10/18/10:	Patient diagnosed with Chlamydia–3 screens needed: Gonorrhea, HIV/AIDS, Syphilis
10/20/10:	Patient diagnosed with Syphilis-removes needed screen for Syphilis (see above)
Result:	Result: Denominator: 2 key STI incidents, Numerator: 1 complete screening (prior to triggering diagnoses but within timeframe)

# Example of Patient with Multiple Diagnoses of Multiple STIs:

06/15/05:	Patient diagnosed with HIV/AIDS
08/01/10:	Patient screened for Chlamydia and Gonorrhea
08/08/10:	Patient diagnosed with Chlamydia and Gonorrhea (Incident #1)–1 screen needed: Syphilis (HIV/AIDS not needed since prior diagnosis)
08/08/10	Patient screened for HIV/AIDS and Syphilis—since only the Syphilis screen is needed, the HIV/AIDS screen is not counted at all
12/01/10:	Patient screened for Chlamydia
12/08/10	Patient diagnosed with Chlamydia (Incident #2) –2 screens needed: Gonorrhea and Syphilis
12/10/10	Patient screened for Syphilis

Result:	Result: Denominator: 2 key STI incidents, Numerator: 1 complete	
	screening	

#### **Key Logic Changes from CRS Version 11.1**

- Changed measure from "# Screenings Needed / # Needed Screenings" to "# Complete Screenings / # Key STI Incidents".
- Removed refusals from main numerators. Refusal numerator is a separate numerator, and no longer a subnumerator.
- Added CPT code 3511F to Chlamydia and Gonorrhea Screening definitions.
- Added CPT code 3512F to Syphilis Screening definition.
- Removed POV codes 078.8\* from Chlamydia definition.

#### **Patient List Description**

List of patients diagnosed with one or more STIs during the defined time period with related screenings.

#### **Measure Source**

None

## **Measure Past Performance and Long-Term Targets**

None

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Sexually Transmitted In	fection (ST	'I) Screening (	con't)		
	-	PREV YR PERIOD			CHG from BASE
Active Clinical Pts w/ Key STI Dx	41	13	+28	10	+31
Male Active Clinical Pt w/ Key STI Dx	s 10	7	+3	8	+2
Female Active Clinical w/ Key STI Dx	Pts 31	6	+25	2	+29
Total # Key STI Inciden Active Clinical Pts		13	+31	10	+34

Total # Male AC Key STI Incidents	10		7		+3	8		+2	
Total # Female AC Key STI Incidents	34		6		+28	2		+32	
# Key STI Incidents for AC Pts	43		13			10			
# Complete Screens Performed-No									
Refusals # Documented	7	16.3	0	0.0	+16.3	0	0.0	+16.3	
Refusals	3	7.0	0	0.0	+7.0	0	0.0	+7.0	
# Key STI Incidents for Male AC Pts	10		7			8			
# Complete Screens									
Performed-No Refusals	1	10.0	0	0.0	+10.0	0	0.0	+10.0	
# Documented Refusals	0	0.0	0	0.0	+0.0	0	0.0	+0.0	

Figure 2-102: Sample Report Sexually Transmitted Infection (STI) Screening

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Sexually Transmitted Infection (STI) Screening: List of patients diagnosed with one or more STIs during the defined time period with related screenings. HRN PATIENT NAME COMMUNITY DENOMINATOR NUMERATOR PATIENT1, DIANE 000001 COMMUNITY #1 F 15 UP; AC Visit 1) 02/12/12 POV: HIV 042. 1) CHL-N, GC-N, SYP-N PATIENT2, LEIGHANN 000002 COMMUNITY #1 F 18 UP; AC Visit 1) 11/02/11 POV: GC 098.89 1) CHL-N, HIV-Y 12/02/11 CPT [87390], SYP-N PATIENT3, WHITNEY 000003 COMMUNITY #1 F 25 UP; AC Visit 1) 06/17/12 POV: CHL 078.89 1) GC-Y 06/17/12 Lab [HGB], HIV-N, SYP-N PATIENT4, NANCY 000004 COMMUNITY #1 F 29 UP; AC Visit 1) 03/01/12 POV: CHL 079.88 1) GC-Y 02/15/12 CPT [87592], HIV-Contraind Prior DX 04/11/07 POV: HIV [079.53], SYP-Y 04/01/12 CPT [87285] PATIENT5, JOHN 000005 COMMUNITY #1 M 40 UP; AC Visit 1) 06/15/12 POV: GC 098.89; 2) 07/15/12 POV: HIV 042. 1) CHL-N, HIV-N, SYP-N; 2) CHL-N, GC-N, SYP-N PATIENT6, NORMAN 000006 COMMUNITY #1 M 42 UP; AC Visit 1) 10/11/12 POV: CHL 079.98, 10/11/12 POV: GC 098.891) HIV-N, SYP-N

Figure 2-103: Sample Patient List, Sexually Transmitted Infection (STI) Screening

# 2.10 Other Clinical Measures Topics

# 2.10.1 Osteoporosis Management

#### **Denominators**

Female Active Clinical patients ages 67 and older who had a new fracture occurring six months (180 days) prior to the Report Period through the first six months of the Report Period with no osteoporosis screening or treatment in year prior to the fracture.

Female User Population patients ages 67 and older who had a new fracture occurring 6 months (180 days) prior to the Report Period through the first 6 months of the Report Period with no osteoporosis screening or treatment in year prior to the fracture.

#### **Numerator**

Patients treated or tested for osteoporosis after the fracture.

#### **Logic Description**

Age is calculated at the beginning of the Report Period. Fractures do not include fractures of finger, toe, face, or skull. CRS will search for the first (i.e., earliest) fracture during the period six months (180) days prior to the beginning of the Report Period and the first six months of the Report Period. If multiple fractures are present, only the first fracture will be used.

Index Episode Start Date definition: The Index Episode Start Date is the date the fracture was diagnosed. If the fracture was diagnosed at an outpatient visit (Service Category A, S, or O), the Index Episode Start Date is equal to the Visit Date. If diagnosed at an inpatient visit (Service Category H), the Index Episode Start Date is equal to the Discharge Date.

#### **Denominator Exclusions**

- 1. Patients receiving osteoporosis screening or treatment in the year (365 days) prior to the Index Episode Start Date. Osteoporosis screening or treatment is defined as a Bone Mineral Density (BMD) test (see below for codes) or receiving any osteoporosis therapy medication (see below for codes).
- 2. Patients with a fracture diagnosed at an outpatient visit, which also had a fracture within 60 days prior to the Index Episode Start Date.
- 3. Patients with a fracture diagnosed at an inpatient visit, which also had a fracture within 60 days prior to the ADMISSION DATE.

Osteoporosis Treatment and Testing definition: (1) For fractures diagnosed at an outpatient visit: a nondiscontinued prescription within 6 months (180 days) of the Index Episode Start Date (i.e., visit date) or B) a BMD test within 6 months of the Index Episode Start Date. (2) For fractures diagnosed at an inpatient visit, a BMD test performed during the inpatient stay.

CRS uses the following codes to define fracture and BMD test.

	CPT Codes	ICD and Other Codes
Fracture Codes	21800–21825, 22305–22314, 22316–22324, 22520, 22521, 22523, 22524, 23500–23515, 23570–23630, 23665–23680, 24500–24585, 24620, 24635, 24650–24685, 25500–25609, 25611 (old code), 25620 (old code), 25622-25652, 25680, 25685, 27193–27248, 27254, 27500–27514, 27520–27540, 27750–27828, \$2360, \$2362	V POV: 733.1*, 805*–806*, 807.0*– 807.4, 808*–815*, 818*–825*, 827*, 828* V Procedure: 79.01–79.03, 79.05–79.07, 79.11–79.13, 79.15–79.17, 79.21–79.23, 79.25–79.27, 79.31–79.33, 79.35–79.37, 79.61–79.63, 79.65–79.67, 81.65-81.66.
BMD Test Codes	77078, 76070 (old code), 77079, 76071 (old code), 77080, 76075 (old code), 77081, 76076 (old code), 77083, 76078 (old code), 76977, 78350, 78351, G0130	V POV: V82.81 V Procedure: 88.98

Treatment medication codes are defined with medication taxonomy BGP HEDIS OSTEOPOROSIS MEDS. (Medications are: Alendronate, Alendronate-Cholecalciferol (Fosomax Plus D), Ibandronate (Boniva), Risedronate, Calcitonin, Raloxifene, Estrogen, Injectable Estrogens, and Teriparatide.) Medications must not have a comment of RETURNED TO STOCK.

#### **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of female patients with new fracture who have had osteoporosis treatment or testing, if any.

#### **Measure Source**

**HEDIS** 

#### **Measure Past Performance and Long-Term Targets**

None

```
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      *** IHS 2012 Selected Measures with Community Specified Report ***
                         DEMO INDIAN HOSPITAL
               Report Period: Jan 01, 2012 to Dec 31, 2012
            Previous Year Period: Jan 01, 2011 to Dec 31, 2011
             Baseline Period: Jan 01, 2000 to Dec 31, 2000
Osteoporosis Management
                  REPORT % PREV YR % CHG from BASE
                  PERIOD
                            PERIOD
                                        PREV YR % PERIOD
                                                            BASE %
Female Active Clinical Pts
67 and older
w/fracture
                                                       0
# w/osteoporosis treatment
              3 37.5
                                   0 0.0 +37.5 0 0.0
or testing
                                                                +37.5
Female User Pop Pts
67 and older
w/fracture
                                                       0
# w/osteoporosis treatment
           4 44.4
or testing
                                   0 0.0 +44.4
                                                       0.0
                                                                +44.4
```

Figure 2-104: Sample Report Osteoporosis Management

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Osteoporosis Management: List of female patients with new fracture who have had osteoporosis treatment or testing, if any. PATTENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, ALWENA 000001 COMMUNITY #1 F 68 FX: 01/01/12 CPT 2252
PATIENT2, SYBIL 000002 COMMUNITY #1 F 69 UP,AC FX: 01/01/12 CPT 22524 FX: 02/10/12 CPT 22524 TX: 02/10/12 CPT G0130 PATIENT3, ELIZABETH 000003 COMMUNITY #1 F 78 UP,AC FX: 02/15/12 CPT S2362 PATIENT4, KATIE 000004 COMMUNITY #1 F 80 FX: 02/05/12 PROC 81.66 UP.AC PATIENT5, LINDSAY 000005 COMMUNITY #1 F 81 FX: 02/01/12 DX 733.13 TX: 02/15/12 CPT 77081 PATIENT6, ELIZABETH 000006 COMMUNITY #1 F 86 FX: 01/15/11 DX 733.13 TX: 01/31/12 CPT 77081

Figure 2-105: Sample Patient List, Osteoporosis Management

# 2.10.2 Osteoporosis Screening in Women

#### **Denominators**

Female Active Clinical patients ages 65 and older without a documented history of osteoporosis.

Female User Population patients ages 65 and older without a documented history of osteoporosis.

#### **Numerators**

Patients who had osteoporosis screening documented in the past two years.

**Note**: This numerator does *not* include refusals.

Patients with documented refusal in past year.

### **Logic Description**

Age is calculated at the beginning of the Report Period.

Osteoporosis definition: No osteoporosis diagnosis ever (POV 733.\*).

CRS uses the following codes to define osteoporosis screening.

	V Radiology or CPT Codes	ICD and Other Codes
Osteoporosis Screening (any test documented in the past two years)	Central DEXA: 77080, 76075 (old code) Peripheral DEXA: 77081, 76076 (old code) SEXA: G0130 Central CT: 77078, 76070 (old code) Peripheral CT: 77079, 76071 (old code) US Bone Density: 76977	V Procedure: 88.98 (Quantitative CT) V POV: V82.81 Special screening for other conditions, Osteoporosis
Osteoporosis Screening Refusal	Refusal (in past year): CPT or V Radiology: Central DEXA: 77080 or 76075 (old code); Peripheral DEXA: 77081 or 76076 (old code); SEXA: G0130; Central CT: 77078 or 76070 (old code); Peripheral CT: 77079 or 76071 (old code); US Bone Density: 76977	

# **Key Logic Changes from CRS Version 11.1**

Added V Radiology codes to Osteoporosis Screening definition.

### **Patient List Description**

List of female patients ages 65 and older with osteoporosis screening, if any.

#### **Measure Source**

None

## **Measure Past Performance and Long-Term Targets**

None

DU November 25, 2012 Page 279  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000							
Osteoporosis Screening in W	omen						
REPORT PERIOD	-			CHG from BAS			
Female Active Clinical Pts =>65 54		29			29		
<pre># w/osteoporosis screening in past 2 years</pre>							
-No Refusals 6 # w/ Osteoporosis Screening	11.1	0	0.0	+11.1	0	0.0	+11.1
_	3.7	0	0.0	+3.7	0	0.0	+3.7
Female User Pop Pts =>65 111		79			80		
<pre># w/osteoporosis screening in past 2 years</pre>							
	5.4	0	0.0	+5.4	0	0.0	+5.4
# w/ Osteoporosis Screening Refusals 2		0	0.0	+1.8	0	0.0	+1.8

Figure 2-106: Sample Report, Osteoporosis Screening in Women

```
UP,AC
                              05/15/12 Refused Proc 88.98
PATIENT2, APRIL
                        000002 COMMUNITY #1 F 68
UP,AC
                             04/01/12 CPT G0130
                        000003 COMMUNITY #1 F 69
PATIENT3, JACKIE
                             08/21/12 RAD CT, BONE MIN DENSITY, 1/+, APPEND
UP,AC
PATIENT4, PAULINE
                       000004 COMMUNITY #1 F 70
UP,AC
PATIENT5, SHANNON
                       000005 COMMUNITY #1 F 72
UP,AC
                       000006 COMMUNITY #1 F 78
PATIENT6, TINA MARIE
UP,AC
                             04/15/12 CPT 77081
```

Figure 2-107: Sample Patient List, Osteoporosis Screening in Women

## 2.10.3 Rheumatoid Arthritis Medication Monitoring

#### **Denominator**

Active Clinical patients ages 16 and older diagnosed with *rheumatoid arthritis* (RA) prior to the Report Period and with at least two RA-related visits any time during the Report Period who were prescribed maintenance therapy medication chronically during the Report Period.

#### **Numerator**

Patients who received appropriate monitoring of chronic medication during the Report Period.

## **Logic Description**

Age is calculated at the beginning of the Report Period.

RA defined as diagnosis (POV or Problem List) 714.\* prior to the Report Period, and at least two RA POVs during the Report Period.

For all maintenance therapy medications *except* intramuscular gold, each medication must be prescribed within the past 465 days of the end of the Report Period (i.e., the Medication Period) and the sum of the days supply =>348. This means the patient must have been on the medication at least 75% of the Medication Period. Two examples are shown below to illustrate this logic. *All* medications must not have a comment of RETURNED TO STOCK.

*Example of Patient Not on Chronic Medication (not included in Denominator):* 

Report Period: Jan 1-Dec 31, 2011

Medication Period: 465 days from end of Report Period (Dec 31, 2011): Sep 22, 2010–Dec 31, 2011

Medication Prescribed:

Diclofenac: 1st Rx: Oct 15, 2010, Days Supply=90; 2nd Rx: Jan 01, 2011: Days Supply=90; 3rd Rx: Mar 15, 2011: Days Supply=90.

Total Days Supply=270. 270 is not >348. Patient is not considered on chronic medication and is not included in the denominator.

Example of Patient on Chronic Medication (included in Denominator):

Report Period: Jan 1-Dec 31, 2011

Medication Period: 465 days from end of Report Period (Dec 31, 2011): Sep 22, 2010–Dec 31, 2011

Medications Prescribed:

Sulfasalazine: 1st Rx: Sep 30, 2010, Days Supply=90; 2nd Rx: Dec 30, 2010, Days Supply=90; 3rd Rx: Mar 15, 2011 Days Supply=180.

Total Days Supply=360. 360 is >348. Patient is considered on chronic medication and is included in denominator.

The days' supply requirement may be met with a single prescription or from a combination of prescriptions for the same medication that were filled during the Medication Period. However, for all medications, there must be at least one prescription filled during the Report period.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

For intramuscular gold, the patient must have 12 or more prescriptions during the Report Period.

Appropriate monitoring of rheumatoid arthritis medications is defined with lab tests and varies by medication, as shown in the table below. If patient is prescribed two or more types of medications, patient must meet criteria for all of the medications.

Maintenance Therapy Medications Definition

Medications shown in table below. Except for Gold, Intramuscular, all
medications requiring more than one of each type of test during the Report Period,
there must be a minimum of ten days between tests. For example, if a
Sulfasalazine test was performed on March 1, March 7, and March 21, 2011, the
March 7 test will not be counted since it was performed only six days after the
March 1 test.

Medication	Required Monitoring Tests
Gold, Intramuscular	CBC and urine Protein on same day as each injection during Report Period
Azathrioprine or Sulfasalazine	4 CBCs during the Report Period
Leflunomide or Methotrexate	6 each of CBC, Serum Creatinine, and Liver Function Tests during the Report Period
Cyclosporin	CBC, Liver Function Tests, and Potassium within past 180 days from Report Period end date 12 Serum Creatinine tests during the Report Period
Gold, Oral or Penicillamine	4 each of CBC and Urine Protein during the Report Period
Mycophenolate	CBC within past 180 days from Report Period end date

The medications in the above table are defined with medication taxonomies: BGP RA IM GOLD MEDS, BGP RA AZATHIOPRINE MEDS, BGP RA LEFLUNOMIDE MEDS, BGP RA METHOTREXATE MEDS, BGP RA CYCLOSPORINE MEDS, BGP RA ORAL GOLD MEDS, BGP RA MYCOPHENOLATE MEDS, BGP RA PENICILLAMINE MEDS, BGP RA SULFASALAZINE MEDS.

- 1. **NSAID Medications**: All of the following medications must have Creatinine, Liver Function Tests, and CBC during the Report Period: Diclofenac, Etodolac, Indomethacin, Ketorolac, Sulindac, Tolmetin, Meclofenamate, Mefanamic Acid, Nabumetone, Meloxicam, Piroxicam, Fenoprofen, Flurbiprofen, Ibuprofen, Ketoprofen, Naproxen, Oxaprozin, Aspirin, Choline Magnesium Trisalicylate, Diflunisil, Magnesium Salicylate, Celocoxib. All of these medications *except* aspirin are defined with medication taxonomy BGP RA OA NSAID MEDS. Aspirin defined with medication taxonomy DM AUDIT ASPIRIN DRUGS.
- Glucocorticoid Medications: Dexamethasone, Methylprednisolone, Prednisone, Hydrocortisone, Betamethasone, Prednisonolone, Triamcinolone. These medications defined with medication taxonomy BGP RA GLUCOCORTICOIDS MEDS. Glucocorticoids must have a glucose test, which must be performed during the Report Period.

Example of Patient Not Included in Numerator:

Medications Prescribed and Required Monitoring:

- Gold, Oral, last Rx Jun 15, 2011. Requires CBC and Urine Protein within past 90 days of Report Period end date.
- CBC performed on Dec 1, 2011, which is within past 90 days of Report Period end date of Dec 31, 2011. No Urine Protein performed during that period. Patient is not in numerator.

Example of Patient Included in Numerator:

Medications Prescribed and Required Monitoring:

- Diclofenac, last Rx Sep 1, 2011. Requires LFT and CBC during Report Period.
- Mycophenolate, last Rx Mar 10, 2011. Requires CBC within past 180 days from Report Period end date.
- LFT and CBC performed during Report Period. CBC performed Nov 1, 2011, which is within past 180 days of Report Period end date of Dec 31, 2011. Patient is in numerator.

Monitoring Test	CPT Codes	LOINC Codes	Taxonomy
CBC	85025, 85027	Yes	BGP CBC TESTS
Urine Protein		Yes	DM AUDIT URINE PROTEIN TAX
Serum Creatinine	82540, 82565-75	Yes	DM AUDIT CREATININE TAX
Liver Function Tests-ALT	84460	Yes	DM AUDIT ALT TAX
Liver Function Tests-AST	84450	Yes	DM AUDIT AST TAX
Liver Function	80076	Yes	BGP LIVER FUNCTION TESTS
Glucose	82947, 82948, 82950, 82951, 82952, 82962	Yes	DM AUDIT GLUCOSE TESTS TAX
Potassium	84132	Yes	BGP POTASSIUM TESTS

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of RA patients age 16 and older prescribed maintenance therapy medication with monitoring laboratory tests, if any. The numerator values for patients who meet the measure are prefixed with YES and patients who did not meet the measure are prefixed with NO The chronic medications and all laboratory tests the patient *did* have are displayed.

#### **Measure Source**

None

#### **Measure Past Performance and Long-Term Targets**

None

```
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       *** IHS 2012 Selected Measures with Community Specified Report ***
                            DEMO INDIAN HOSPITAL
                 Report Period: Jan 01, 2012 to Dec 31, 2012
             Previous Year Period: Jan 01, 2011 to Dec 31, 2011
                Baseline Period: Jan 01, 2000 to Dec 31, 2000
Rheumatoid Arthritis Medication Monitoring
                            % PREV YR % CHG from BASE
PERIOD PREV YR % PERIO
                    REPORT
                                                                  % CHG from
                                               PREV YR % PERIOD
                                                                     BASE %
                    PERTOD
Active Clinical Pts =>16
w/RA DX and maintenance
                                       0
                                                              0
therapy RX
# w/RA chronic med
                       2 66.7
                                       0 0.0 +66.7 0 0.0 +66.7
monitoring
```

Figure 2-108: Sample Report, Rheumatoid Arthritis Medication Monitoring

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Rheumatoid Arthritis Medication Monitoring: List of RA patients 16 and older prescribed maintenance therapy medication with monitoring lab tests, if any. The numerator values for patients who meet the measure are prefixed with "YES:" and patients who did not meet the measure are prefixed with "NO:". The chronic medications and all lab tests the patient DID have are displayed. HRN COMMUNITY SEX AGE PATTENT NAME DENOMINATOR NUMERATOR PATIENT1, RUTH 000001 COMMUNITY #1 F 64 YES: NSAID: 10/21/12 CREAT, 09/22/12 CBC, 05/21/12 LFT PATIENT2, SHANNON 000002 COMMUNITY #1 F 72 YES: Glucocorticoids: 12/02/12 Glucose PATIENT34, CATHERINE 000034 COMMUNITY #3 F 50 NO: Glucocorticoids: does not have Glucose

Figure 2-109: Sample Patient List, Rheumatoid Arthritis Medication Monitoring

## 2.10.4 Osteoarthritis Medication Monitoring

#### Denominator

Active Clinical patients ages 40 and older diagnosed with osteoarthritis (OA) prior to the Report Period and with at least two OA-related visits any time during the Report Period and prescribed maintenance therapy medication chronically during the Report Period.

#### **Numerator**

Patients who received appropriate monitoring of chronic medication during the Report Period.

#### **Logic Description**

Age is calculated at the beginning of the Report Period.

OA defined as diagnosis (POV or Problem List) 715.\* prior to the Report period, and at least two OA POVs during the Report Period.

For all maintenance therapy medications, each medication must be prescribed within the past 465 days of the end of the Report Period (i.e., the Medication Period) and the sum of the days supply =>348. This means the patient must have been on the medication at least 75% of the Medication Period. Two examples are shown below to illustrate this logic. Medications must not have a comment of RETURNED TO STOCK.

Example of Patient Not on Chronic Medication (not included in Denominator):

- Report Period: Jan 1–Dec 31, 2011
- Medication Period: 465 days from end of Report Period (Dec 31, 2011): Sep 22, 2010–Dec 31, 2011
- Medication Prescribed:
  - Diclofenac: 1st Rx: Oct 15, 2010, Days Supply=90; 2nd Rx: Jan 1, 2011:
     Days Supply=90; 3rd Rx: Mar 15, 2011: Days Supply=90.
  - Total Days Supply=270. 270 is not >348. Patient is not considered on chronic medication and is not included in the denominator.

*Example of Patient on Chronic Medication (included in Denominator):* 

- Report Period: Jan 1–Dec 31, 2011
- Medication Period: 465 days from end of Report Period (Dec 31, 2011): Sep 22, 2010–Dec 31, 2011
- Medication Prescribed:

- Etodolac: 1st Rx: Sep 30, 2010, Days Supply=90; 2nd Rx: Dec 30, 2010,
   Days Supply=90; 3rd Rx: Mar 15, 2011: Days Supply =180.
- Total Days Supply=360. 360 is >348. Patient is considered on chronic medication and is included in denominator.

The days supply requirement may be met with a single prescription or from a combination of prescriptions for the same medication that were filled during the Medication Period. However, for all medications, there must be at least one prescription filled during the Report Period.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Appropriate monitoring of OA medications is defined with laboratory tests and varies by medication, as shown below.

Maintenance Therapy Medications Defined with the Following NSAID Medications: Diclofenac, Etodolac, Indomethacin, Ketorolac, Sulindac, Tolmetin, Meclofenamate, Mefanamic Acid, Nabumetone, Meloxicam, Piroxicam, Fenoprofen, Flurbiprofen, Ibuprofen, Ketoprofen, Naproxen, Oxaprozin, Aspirin, Choline Magnesium Trisalicylate, Diflunisil, Magnesium Salicylate, Celocoxib. All of these medications, except aspirin, are defined with medication taxonomy BGP RA OA NSAID MEDS. Aspirin defined with medication taxonomy DM AUDIT ASPIRIN DRUGS

All NSAID medications must have Creatinine, Liver Function Tests and CBC during the Report Period.

Example of Patient Not Included in Numerator:

- Medication Prescribed and Required Monitoring:
  - Diclofenac, last Rx Jun 15, 2011. Requires Creatinine, LFT and CBC during Report Period. Only the LFT was performed during Report Period. Patient is not in numerator.

Example of Patient Included in Numerator:

- Medications Prescribed and Required Monitoring:
  - Diclofenac, last Rx Sep 1, 2011. Requires Creatinine, LFT and CBC during Report Period. Creatinine, LFT, and CBC performed during Report Period. Patient is in numerator.

CRS uses the following codes to define the monitoring tests.

Monitoring Test	CPL COMPS	LOINC Codes	Taxonomy
Serum Creatinine	82540, 82565-82575	Yes	DM AUDIT CREATININE TAX
CBC	85025, 85027	Yes	BGP CBC TESTS
Liver Function Tests-ALT	84460	Yes	DM AUDIT ALT TAX
Liver Function Tests-AST	84450	Yes	DM AUDIT AST TAX
Liver Function	80076	Yes	BGP LIVER FUNCTION TESTS

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of OA patients 40 and older prescribed maintenance therapy medication with monitoring laboratory tests, if any. The numerator values for patients who meet the measure are prefixed with YES and patients who did not meet the measure are prefixed with NO. All laboratory tests the patient did have are displayed.

#### **Measure Source**

None

## Measure Past Performance and Long-Term Targets

None

DU November 25, 2012 Page 287  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000					
Osteoarthritis Medication Monitoring  REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %					
Active Clinical Pts =>40 w/OA DX and maintenance therapy RX 3 6 4					
# w/OA chronic med monitoring 2 66.7 3 50.0 +16.7 2 50.0 +16.7					

Figure 2-110: Sample Report, Osteoarthritis Medication Monitoring

Figure 2-111: Sample Patient List, Osteoarthritis Medication Monitoring

#### 2.10.5 Asthma

#### **Denominators**

All *Active Clinical patients*. Broken down by age groups (under 14, 15 to 34, 35 to 64, and 65 and older).

Patients who have had two asthma-related visits during the Report Period or with persistent asthma. Broken down by age groups (under 14, 15 to 34, 35 to 64, and 65 and older).

#### **Numerators**

Patients who have had two asthma-related visits during the Report Period or with persistent asthma.

Patients from Numerator 1 who have been hospitalized at any hospital for asthma during the Report Period.

Patients from Numerator 1 who have visited the ER or Urgent Care for asthma during the Report Period.

Patients from Numerator 1 who have a Severity of 1.

Patients from Numerator 1 who have a Severity of 2.

Patients from Numerator 1 who have a Severity of 3.

Patients from Numerator 1 who have a Severity of 4.

Patients from Numerator 1 who have no documented Severity.

## **Logic Description**

Age is calculated at beginning of Report Period.

Asthma visits definition: Diagnosis (POV) 493.\*.

Persistent asthma definition: Any of the following:

- Active entry in PCC Problem List for 493.\* with Severity of 2, 3 or 4 at *any* time before the end of the Report Period, *or*
- Most recent visit-related asthma entry (i.e., V Asthma) with Severity of 2, 3, or 4 documented *any* time before the end of the Report Period.

Severity definition: Severity of 1, 2, 3 or 4 in an active entry in the PCC Problem List for 493.\* or in V Asthma.

Hospitalizations definition: Service Category H with primary POV 493.\*.

ER and Urgent Care definition: Clinic codes 30 or 80 with primary POV 493.\*.

## **Key Logic Changes from CRS Version 11.1**

- Changed age breakdowns to under 14, 15 to 34, 35 to 64, and 65 and older.
- Added new measures and corresponding logic.
- Updated patient list.

## **Patient List Description**

List of patients diagnosed with asthma and any asthma-related hospitalizations.

#### **Measure Source**

HP 2020 RD-2

## **Measure Past Performance and Long-Term Targets**

Measure	Target
HP1998 baseline for hospitalizations for asthma:	
Under 5	45.6 per 10,000
5-64	12.5 per 10,000
65 and older	17.7 per 10,000
HP 2020 goal for hospitalizations for asthma:	
Under 5	18.1 per 10,000
5-64	8.6 per 10,000
65 and older	20.3 per 10,000

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Asthma (con't)					CHG from PREV YR %				
Total Active Clinic	1,646		1,224			1,101			
# w/asthma A. Under 15 B. 15-34 C. 35-64 D. 65 and older	7 15	12.1 25.9	7 8	19.0	-4.1 -4.6 +6.8	17 3 3	12.0 12.0	-19.7 +0.1 +13.9	
# w/asthma	58		42			25			
<pre># w/asthma hospitalization A. Under 15 B. 15-34 C. 35-64 D. 65 and older</pre>	1 0 0	50.0 0.0 0.0	0 1 0		-50.0 +0.0	1 0 1	50.0	+0.0 -50.0	
# w/ ER/UC visit A. Under 15 B. 15-34 C. 35-64 D. 65 and older	2	10.3 33.3 0.0 33.3 33.3	3 2 0	14.3 50.0 33.3 0.0 16.7	-33.3 +33.3	7 1 1	36.0 77.8 11.1 11.1 0.0	-44.4 -11.1 +22.2	
# w/ Severity 3	7 4 6	6.9 10.3	3 0 2	7.1	+4.9 +6.9 +5.6	0 0 0	0.0	+10.3	

Figure 2-112: Sample Report, Asthma

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```
PATIENT3, PAULINE 000003 COMMUNITY #1 F 70

AC 2 Dx PCC: 03/01/12, 03/03/12; ER/UC: 10/01/12;

Severity: 1

PATIENT4, WILLIAM R 000004 COMMUNITY #1 M 7

AC 2 Dx PCC: 05/05/12, 06/06/12; Hospital: 05/05/12;

ER/UC: 06/06/12

PATIENT5, ZACHARY 000005 COMMUNITY #1 M 11

AC 2 Dx PCC: 03/20/12, 08/08/12

PATIENT42, JOSEPHINE 000042 COMMUNITY #2 F 4

AC 2 Dx PCC: 07/01/12, 09/19/12; Severity: 2
```

Figure 2-113: Sample Patient List, Asthma

## 2.10.6 Asthma Assessments

#### **Denominators**

Active Clinical patients ages 5 and older with persistent asthma within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema or chronic obstructive pulmonary disease (COPD). Broken down by age groups (5-14, 15-34, 35-64, and >65).

#### **Numerator**

Patients with asthma management plan during the Report Period.

Patients with severity documented at any time before the end of the Report Period.

Patients with control documented during the Report Period.

Patients who were assessed for number of symptom free days during the Report Period.

Patients with number of symptom free days score of 0-5.

Patients with number of symptom free days score of 6-12.

Patients with number of symptom free days score of 13-14.

Patients who were assessed for number of school/work days missed during the Report Period.

Patients with number of school/work days missed score of 0-2.

Patients with number of school/work days missed score of 3-7.

Patients with number of school/work days missed score of 8-14.

#### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

*Emphysema definition:* Any visit at any time on or before the end of the Report Period with POV codes: 492.\*, 506.4, 518.1, 518.2.

COPD definition: Any visit at any time on or before the end of the Report Period with POV codes: 491.20, 491.21, 491.22, 493.2\*, 496, 506.4.

### Persistent asthma definition:

- 1. Meeting any of the following four criteria below within the year prior to the beginning of the Report Period *and* during the Report Period:
  - a. At least one visit to Clinic Code 30 (Emergency Medicine) with primary diagnosis 493\* (asthma)
  - b. At least one acute inpatient discharge with Primary Diagnosis 493.\*. Acute inpatient discharge defined as Service Category of H
  - c. At least four outpatient visits, defined as Service Categories A, S, or O, with primary or secondary diagnosis of 493.\* *and* at least two asthma medication dispensing events (see definition below)
  - d. At least four asthma medication dispensing events (see definition below). If the sole medication was leukotriene modifiers, then *must* also have at least one visit with POV 493.\* in the same year as the leukotriene modifier (i.e., during the Report Period or within the year prior to the beginning of the Report Period.)
- 2. Meeting any of the following criteria below:
  - a. Active entry in PCC Problem List for 493.\* with Severity of 2, 3 or 4 at any time before the end of the Report Period, or
  - b. Most recent visit-related asthma entry (i.e., V Asthma) with Severity of 2, 3, or 4 documented any time before the end of the Report Period.

Dispensing event definition: One prescription of an amount lasting 30 days or less. For RXs longer than 30 days, divide the days' supply by 30 and round down to convert. For example, a 100-day RX is equal to three dispensing events (100/30 = 3.33, rounded down to 3). Also, two different RXs dispensed on the same day are counted as two different dispensing events. Inhalers should also be counted as one dispensing event.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Asthma medication codes for denominator defined with medication taxonomies: BGP HEDIS ASTHMA MEDS, BGP HEDIS ASTHMA LEUK MEDS, BGP HEDIS ASTHMA INHALED MEDS. Medications are: Antiasthmatic Combinations (Dyphylline-Guaifenesin, Guaifenesin-Theophylline, Potassium Iodide-Theophylline), Antibody Inhibitor (Omalizumab), Inhaled Steroid Combinations (Budesonide-Formoterol, Fluticasone-Salmeterol), Inhaled Corticosteroids (Belclomethasone, Budesonide, Flunisolide, Fluticasone CFC Free, Mometasone, Triamcinolone), Lekotriene Modifiers (Montelukast, Zafirlukast, Zileuton), Long-Acting, Inhaled Beta-2 Agonists (Aformoterol, Formoterol, Salmeterol), Mast Cell Stabilizers (Cromolyn, Nedocromil), Methylxanthines (Aminophylline, Dyphylline, Oxtriphylline, Theophylline), Short-Acting, Inhaled Beta-2 Agonists (Albuterol, Levalbuterol, Metaproterenol, Pirbuterol). Medications must not have a comment of RETURNED TO STOCK.

Asthma management plan definition: Patient Education code ASM-SMP.

Severity documented definition:

Meeting any of the following criteria below:

- 1. Active entry in PCC Problem List for 493.\* with Severity of 2, 3 or 4 at ANY time before the end of the Report Period or
- 2. Most recent visit-related asthma entry (i.e. V Asthma) with Severity of 2, 3, or 4 documented ANY time before the end of the Report Period.

Control documented definition: POV 493.\* with Asthma Control recorded in the V Asthma file.

*Number of symptom free days definition:* The most recent V Measurement documented during the Report Period.

*Number of school/work days missed definition:* The most recent V Measurement documented during the Report Period.

## **Key Logic Changes from CRS Version 11.1**

New topic.

## **Patient List Description**

List of asthmatic patients with assessments, if any.

#### **Measure Source**

None

## **Measure Past Performance and Long-term Targets**

DU November 25, 2012 Page 295  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000										
Asthma Assessments										
	REPORT PERIOD				CHG from BA					
Active Clinical										
Pts =>5 w/ persister	nt									
asthma	29		11			5				
<pre># w/management plan # w/severity</pre>	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
documented # w/control	17	58.6	5	45.5	+13.2	1	20.0	+38.6		
documented	3	10.3	0	0.0	+10.3	0	0.0	+10.3		
# w/# symptom free										
days	3	10.3	0	0.0	+10.3	0	0.0	+10.3		
<pre># w/# symptom free days 0-5</pre>	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
# w/# symptom free	_	3.1	Ü	0.0	. 3. 1	O	0.0	.3.1		
days 6-12	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
<pre># w/# symptom free days 13-14</pre>	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
# w/# school/work	1	3.4	U	0.0	+3.4	U	0.0	+3.4		
days missed	3	10.3	0	0.0	+10.3	0	0.0	+10.3		
# w/# school/work										
days missed 0-2 # w/# school/work	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
days missed 3-7	1	3.4	0	0.0	+3.4	0	0.0	+3.4		
# w/# school/work days missed 8-14	1	3.4	0	0.0	+3.4	0	0.0	+3.4		

Figure 2-114: Sample Report, Asthma Assessments

```
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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012
```

Previous Year Period: Jan 01, 2011 to Dec 31, 2011 Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Asthma Assessments (con't)									
Active Clini			tent Asthma 35-64						
CURRENT REPORT PERIOD Active Clinical Pts =>5									
w/persistent asthma	11	6	6	6					
# w/management plan	1	0	0	0					
% w/managment plan	9.1	0.0	0.0	0.0					
# w/severity documented	7	3	6	1					
% w/severity documented	63.6		100.0	16.7					
<pre>% w/control documented % w/control documented</pre>	0	1 16.7	1 16.7	1 16.7					
<pre># w/# symptom free days % w/# symptom free days</pre>		0.0	0.0	2 33.3					
# w/# symptom free days 0-5	0	0	0	1					
% w/# symptom free days 0-5		0.0		16.7					
# w/# symptom free days 6-12	1	0	0	0					
% w/# symptom free days $6-12$	9.1	0.0	0.0	0.0					
# w/# symptom free days 13-14		0	0	1					
% w/# symptom free days $13-14$	0.0	0.0	0.0	16.7					
# w/# school/work days missed	1	0	0	2					
% w/# school/work days									
missed	9.1	0.0	0.0	33.3					
<pre># w/# school/work days missed 0-2</pre>	1	0	0	0					
% w/# school/work days missed 0-2	0 1								
	9.1	0.0	0.0	0.0					
<pre># w/# school/work days missed 3-7</pre>	0	0	0	1					
% w/# school/work days									
missed 3-7	0.0	0.0	0.0	16.7					
<pre># w/# school/work days missed 8-14</pre>	0	0	0	1					
% w/# school/work days									
missed 8-14	0.0	0.0	0.0	16.7					

Figure 2-115: Sample Age Breakdown Report, Asthma Assessments

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

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```
Asthma Assessments: List of asthmatic patients with assessments, if any.
PATIENT NAME HRN COMMUNITY
DENOMINATOR NUMERATOR
                                        SEX AGE
DENOMINATOR
                     NUMERATOR
PATIENT1, GENEVA 000001 COMMUNITY #1 F 47
UP, AC Severity 4 in V Asthma 02/02/11 Severity: 4; Symptom Free Days: 03/01/12 [12];
Days Missed: 03/01/12 [0]
PATIENT2, JACKIE 000002 COMMUNITY #1 F 69
UP,AC Severity >1 on PL for 493.00 Severity: 2
PATIENT3, PAULINE 000003 COMMUNITY #1 F 70
UP,AC Severity >1 on PL for 493.00 Mgmt Plan: 06/01/12; Severity: 3
PATIENT4, WILLIAM R 000004 COMMUNITY #1 M 7
UP,AC 4 meds
PATIENT5, ZACHARY 000005 COMMUNITY #1 M 11
AC Severity 4 in V Asthma 04/04/12 Severity: 4; Control: 05/06/12
PATIENT42, JOSEPHINE 000042 COMMUNITY #2 F 4
AC DX ON HOSP/OR ER ON 05/05/11 DX ON HOSP/OR ER ON 06/03/12 Control: 10/03/12;
Symptom Free Days: 07/01/12 [3]; Days Missed: 07/01/12 [3]
```

Figure 2-116: Sample Patient List, Asthma Assessments

## 2.10.7 Asthma Quality of Care

#### **Denominators**

Active Clinical patients ages 5–56 with persistent asthma within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema or chronic obstructive pulmonary disease (COPD). Broken down by age groups (5–9, 10–17, and 18–56).

*User Population patients ages 5–56 with persistent asthma* within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema and chronic COPD. Broken down by age groups (5–9, 10–17, and 18–56).

#### **Numerator**

Patients who had at least one dispensed prescription for preferred asthma therapy medication during the Report Period.

#### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

*Emphysema definition:* Any visit at any time on or before the end of the Report Period with POV codes: 492.\*, 506.4, 518.1, 518.2.

COPD definition: Any visit at any time on or before the end of the Report Period with POV codes: 491.20, 491.21, 491.22, 493.2\*, 496, 506.4.

#### Persistent asthma definition:

- 3. Meeting any of the following four criteria below within the year prior to the beginning of the Report Period *and* during the Report Period:
  - a. At least one visit to Clinic Code 30 (Emergency Medicine) with primary diagnosis 493\* (asthma)
  - b. At least one acute inpatient discharge with Primary Diagnosis 493.\*. Acute inpatient discharge defined as Service Category of H
  - c. At least four outpatient visits, defined as Service Categories A, S, or O, with primary or secondary diagnosis of 493.\* *and* at least two asthma medication dispensing events (see definition below)
  - d. At least four asthma medication dispensing events (see definition below). If the sole medication was leukotriene modifiers, then *must* also have at least one visit with POV 493.\* in the same year as the leukotriene modifier (i.e., during the Report Period or within the year prior to the beginning of the Report Period.)
- 4. Meeting any of the following criteria below:
  - a. Active entry in PCC Problem List for 493.\* with Severity of 2, 3 or 4 at any time before the end of the Report Period, or
  - b. Most recent visit-related asthma entry (i.e., V Asthma) with Severity of 2, 3, or 4 documented any time before the end of the Report Period.

Dispensing event definition: One prescription of an amount lasting 30 days or less. For RXs longer than 30 days, divide the days' supply by 30 and round down to convert. For example, a 100-day RX is equal to three dispensing events (100/30 = 3.33, rounded down to 3). Also, two different RXs dispensed on the same day are counted as two different dispensing events. Inhalers should also be counted as one dispensing event.

Note: If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Asthma medication codes for denominator defined with medication taxonomies: BGP HEDIS ASTHMA MEDS, BGP HEDIS ASTHMA LEUK MEDS, BGP HEDIS ASTHMA INHALED MEDS. Medications are: Antiasthmatic Combinations (Dyphylline-Guaifenesin, Guaifenesin-Theophylline, Potassium Iodide-Theophylline), Antibody Inhibitor (Omalizumab), Inhaled Steroid Combinations (Budesonide-Formoterol, Fluticasone-Salmeterol), Inhaled Corticosteroids (Belclomethasone, Budesonide, Flunisolide, Fluticasone CFC Free, Mometasone, Triamcinolone), Lekotriene Modifiers (Montelukast, Zafirlukast, Zileuton), Long-Acting, Inhaled Beta-2 Agonists (Aformoterol, Formoterol, Salmeterol), Mast Cell Stabilizers (Cromolyn, Nedocromil), Methylxanthines (Aminophylline, Dyphylline, Oxtriphylline, Theophylline), Short-Acting, Inhaled Beta-2 Agonists (Albuterol, Levalbuterol, Metaproterenol, Pirbuterol). Medications must not have a comment of RETURNED TO STOCK.

To be included in the numerator, patient must have a non-discontinued prescription for preferred asthma therapy (see list of medications below) during the Report Period.

Preferred asthma therapy medication codes for numerator defined with medication taxonomy: BGP HEDIS PRIMARY ASTHMA MEDS Medications are: Antiasthmatic Combinations (Dyphylline-Guaifenesin, Guaifenesin-Theophylline, Potassium Iodide-Theophylline), Antibody Inhibitor (Omalizumab), Inhaled Steroid Combinations (Budesonide-Formoterol, Fluticasone-Salmeterol), Inhaled Corticosteroids (Belclomethasone, Budesonide, Flunisolide, Fluticasone CFC Free, Mometasone, Triamcinolone), Lekotriene Modifiers (Montelukast, Zafirlukast, Zileuton), Mast Cell Stabilizers (Cromolyn, Nedocromil), Methylxanthines (Aminophylline, Dyphylline, Oxtriphylline, Theophylline). Medications must not have a comment of RETURNED TO STOCK.

## **Key Logic Changes from CRS Version 11.1**

None

#### **Patient List Description**

List of asthmatic patients with preferred asthmatherapy medications, if any.

#### **Measure Source**

# **Measure Past Performance and Long-Term Targets**

None

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Asthma Quality of Care (con't)									
	REPORT PERIOD				CHG from PREV YR %			CHG from BASE %	
Active Clinical Pts w/persistent asthma	5-56		10			5			
<pre># w/ preferred asthm control med</pre>		33.3	5	50.0	-16.7	3	60.0	-26.7	
A. Active Clinical ages 5-9	11		3			1			
# w/ preferred asthm		27.3	1	33.3	-6.1	1	100.0	-72.7	
B. Active Clinical ages 10-17	2		2			1			
# w/ preferred asthm		100.0	2	100.0	+0.0	0	0.0	+100.0	
C. Active Clinical ages 18-56	8		5			3			
# w/ preferred asthm		25.0	2	40.0	-15.0	2	66.7	-41.7	
User Pop Pts 5-56 w/persistent asthma	21		11			6			
# w/ preferred asthm		33.3	5	45.5	-12.1	3	50.0	-16.7	
A. User Pop ages 5-9	11		4			1			
# w/ preferred asthm control med	a 3	27.3	1	25.0	+2.3	1	100.0	-72.7	
B. User Pop ages 10-17	2		2			1			

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# w/ preferred asthma control med	2 100.0	2 100.0 +0.0	0 0.0 +100.0
C. User Pop ages 18-56	8	5	4
# w/ preferred asthma control med	2 25.0	2 40.0 -15.0	2 50.0 -25.0

Figure 2-117: Sample Report, Asthma Quality of Care

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Asthma Quality of Care: List of asthmatic patients with preferred asthma
therapy medications, if any.
DENOMINATOR HRN
                           COMMUNITY SEX AGE
PATIENT NAME
                     NUMERATOR
PATIENT1, ZACHARY 000011 COMMUNITY
                                             F 5
UP, AC, Severity 4 in V Asthma 02/02/11
PATIENT12, TINA DANIELLE 000012 COMMUNITY F 6
UP,AC,Severity >1 on PL for 493.00,NUM: 06/01/12, MOMETASONE/FORMOTEROL 100/5MCG INH
PATIENT13, THERESA LYNN 000013 COMMUNITY M 47
UP,AC,Severity 2 in V Asthma 03/03/12,NUM: 06/01/12, FLUTICASONE PROPIONATE 110MCG
INHALER
PATIENT36, NATHAN BRADLEY 000014 COMMUNITY
                                             M 16
UP,AC,DX ON HOSP/OR ER ON 09/26/11,4 POVS AND 2 MEDSNUM: 09/19/12, FLUTICASONE
PROPIONATE 110MCG INHALER
PATIENT37, JANELLE MARIE 000015 COMMUNITY F 50
UP, AC, Severity 2 in V Asthma 09/04/11
PATIENT38, THOMAS ELLIS 000016 COMMUNITY M 6
UP,AC,4 meds
```

Figure 2-118: Sample Patient List, Asthma Quality of Care

# 2.10.8 Medication Therapy for Persons with Asthma

#### **Denominators**

Active Clinical patients ages 5-50 with persistent asthma within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema or chronic obstructive pulmonary disease (COPD).

Active Clinical patients ages 5 and older with persistent asthma within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema or chronic obstructive pulmonary disease (COPD). Broken down by age groups (5-14, 15-34, 35-64, and >65).

Active Clinical patients ages five and older with persistent asthma within the year prior to the beginning of the Report Period and during the Report Period, without a documented history of emphysema or chronic obstructive pulmonary disease (COPD) who had 2 or more prescriptions for a LABA during the Report Period. Broken down by age groups (5-14, 15-34, 35-64, and >65).

#### **Numerators**

Suboptimal Control: Patients who were dispensed more than three canisters of a short-acting beta2 agonist inhaler during the same 90-day period during the Report Period.

Absence of Controller Therapy: Patients who were dispensed more than three canisters of short acting beta2 agonist inhalers over a 90-day period and who did not receive controller therapy during the same 90-day period.

Patients who were prescribed two or more controller therapy medications during the Report Period.

Patients who were prescribed two or more inhaled corticosteroid medications during the Report Period.

Patients who were not prescribed two or more inhaled corticosteroid medications during the Report Period.

## **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

*Emphysema definition:* Any visit at any time on or before the end of the Report Period with POV codes: 492.\*, 506.4, 518.1, 518.2.

COPD definition: Any visit at any time on or before the end of the Report Period with POV codes: 491.20, 491.21, 491.22, 493.2\*, 496, 506.4.

Persistent asthma definition:

- 5. Meeting any of the following four criteria below within the year prior to the beginning of the Report Period *and* during the Report Period:
  - a. At least one visit to Clinic Code 30 (Emergency Medicine) with primary diagnosis 493\* (asthma)
  - b. At least one acute inpatient discharge with Primary Diagnosis 493.\*. Acute inpatient discharge defined as Service Category of H

- c. At least four outpatient visits, defined as Service Categories A, S, or O, with primary or secondary diagnosis of 493.\* *and* at least two asthma medication dispensing events (see definition below)
- d. At least four asthma medication dispensing events (see definition below). If the sole medication was leukotriene modifiers, then *must* also have at least one visit with POV 493.\* in the same year as the leukotriene modifier (i.e., during the Report Period or within the year prior to the beginning of the Report Period.)
- 6. Meeting any of the following criteria below:
  - a. Active entry in PCC Problem List for 493.\* with Severity of 2, 3 or 4 at any time before the end of the Report Period, or
  - b. Most recent visit-related asthma entry (i.e., V Asthma) with Severity of 2, 3, or 4 documented any time before the end of the Report Period.

Dispensing event definition: One prescription of an amount lasting 30 days or less. For RXs longer than 30 days, divide the days' supply by 30 and round down to convert. For example, a 100-day RX is equal to three dispensing events (100/30 = 3.33, rounded down to 3). Also, two different RXs dispensed on the same day are counted as two different dispensing events. Inhalers should also be counted as one dispensing event.

**Note:** If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

Asthma medication codes for denominator defined with medication taxonomies: BGP HEDIS ASTHMA MEDS, BGP HEDIS ASTHMA LEUK MEDS, BGP HEDIS ASTHMA INHALED MEDS. Medications are: Antiasthmatic Combinations (Dyphylline-Guaifenesin, Guaifenesin-Theophylline, Potassium Iodide-Theophylline), Antibody Inhibitor (Omalizumab), Inhaled Steroid Combinations (Budesonide-Formoterol, Fluticasone-Salmeterol), Inhaled Corticosteroids (Belclomethasone, Budesonide, Flunisolide, Fluticasone CFC Free, Mometasone, Triamcinolone), Lekotriene Modifiers (Montelukast, Zafirlukast, Zileuton), Long-Acting, Inhaled Beta-2 Agonists (Aformoterol, Formoterol, Salmeterol), Mast Cell Stabilizers (Cromolyn, Nedocromil), Methylxanthines (Aminophylline, Dyphylline, Oxtriphylline, Theophylline), Short-Acting, Inhaled Beta-2 Agonists (Albuterol, Levalbuterol, Metaproterenol, Pirbuterol). Medications must not have a comment of RETURNED TO STOCK.

To be included in the Suboptimal Control and Absence of Controller Therapy numerators, patient must have one or more non-discontinued prescriptions for short acting Beta2 Agonist inhalers totaling at least four canisters in one 90 day period. Short acting Beta2 Agonist inhaler medications defined with medication taxonomy BGP PQA SABA MEDS. (Medications are: Albuterol, Levalbuterol, Pirbuterol). Medications must not have a comment of RETURNED TO STOCK.

## Controller Therapy definition:

At least one non-discontinued prescription of controller therapy medications during the same 90 day period.

Controller therapy medications defined with medication taxonomy BGP PQA CONTROLLER MEDS. (Medications are: Beclomethasone, Budesonide, Budesonide-Formoterol, Ciclesonide, Cromolyn, Flunisolide, Fluticasone, Fluticasone-Salmeterol, Formoterol, Mometasone, Mometasone-Formoterol, Montelukast, Nedocromil, Salmeterol, Theophylline, Triamcinolone, Zafirlukast, Zileuton). Medications must not have a comment of RETURNED TO STOCK.

Inhaled corticosteroid medications defined with medication taxonomy BGP ASTHMA INHALED STEROIDS. (Medications are: Mometasone (Asmanex), Beclovent, Qvar, Vancenase, Vanceril, Vanceril DS, Bitolerol (Tornalate), Pulmicort, Pulmicort Respules, Pulmicort Turbohaler, Salmeterol/fluticasone (Advair), Triamcinolone (Azmacort), Fluticasone (Flovent), Budesonide-Formoterol (Symbicort).) Medications must not have a comment of RETURNED TO STOCK.

Long-Acting Beta-2 Agonist (LABA) medications defined with medication taxonomy BGP ASTHMA LABA MEDS. (Medications are: Aformoterol, Formoterol, Salmeterol.) Medications must not have a comment of RETURNED TO STOCK.

## **Key Logic Changes from CRS Version 11.1**

- Changed the denominator to exclude patients with emphysema or COPD and added corresponding logic.
- Changed persistent asthma logic to match that in Asthma Quality of Care.
- Added new measures and corresponding logic.
- Updated patient list.

## **Patient List Description**

List of patients with asthma with suboptimal control and controller therapy, if any.

#### **Measure Source**

PQA (Pharmacy Quality Alliance)

# **Measure Past Performance and Long-term Targets**

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Medication Therapy for Persons with Asthma									
REPORT % PREV YR % CHG from BASE % CHG from PERIOD PREV YR % PERIOD BASE %									
Active Clinical ages 5-50 w/ asthma	19		8			5			
<pre># w/ Suboptimal   Control # w/ Absence of Control!</pre>	_	5.3	0	0.0	+5.3	0	0.0	+5.3	
Therapy		100.0	0	0.0	+100.0	0	0.0	+100.0	
Active Clinical Pts =>5 w/persistent asthma	29		11			5			
<pre># w/ 2 or more controller Rx # w/ 2 or more inhaled</pre>	10	34.5	5	45.5	-11.0	3	60.0	-25.5	
steroid Rx	3	10.3	3	27.3	-16.9	0	0.0	+10.3	
Active Clinical =>5 w/persistent asthma and LABA Rx	4		0			0			
# w/o 2 or more inhaled steroid Rx	3	75.0	0	0.0	+75.0	0	0.0	+75.0	

Figure 2-119: Sample Report, Medication Therapy for Persons with Asthma

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Medication Therapy for Pers	ons with Asthm	na (con't)								
	Active Clinical Pts =>5 w/persistent asthma									
CURRENT REPORT PERIOD	5-14	15-34	35-64	65+						
Active Clinical Pts =>5 w/persistent asthma	11	6	6	6						

#/ 2 mana mambaallan D	C	2	1	1	
<pre># w/ 2 or more controller Rx % w/ 2 or more controller Rx</pre>			<del>-</del>		
# w/ 2 or more inhaled steroid Rx	3	0	0	0	
% w/ 2 or more inhaled					
steroid Rx	27.3	0.0	0.0	0.0	
PREVIOUS REPORT PERIOD					
Active Clinical Pts =>5 w/persistent asthma	4	2	3	1	
w/persistent asthma	4	4	3	Τ	
<pre># w/ 2 or more controller Rx % w/ 2 or more controller Rx</pre>	2	1	1 33.3	0	
% w/ 2 or more controller Rx	50.0	50.0	33.3	0.0	
# w/ 2 or more inhaled					
steroid Rx % w/ 2 or more inhaled	1	1	1	0	
steroid Rx	25.0	50.0	33.3	0.0	
CHANGE FROM PREVIOUS YR %					
# w/ 2 or more controller Rx	+4.5	-16.7	-16.7	+16.7	
<pre># w/ 2 or more inhaled steroid Rx</pre>	-22 7	-50.0	-33 3	+0 0	
Secrota in	22.7	30.0	33.3	10.0	
BASELINE REPORT PERIOD					
Active Clinical Pts =>5					
w/persistent asthma	2	2	1	0	
# w/ 2 or more controller Rx	1	1	1	0	
% w/ 2 or more controller Rx		50.0	100.0	0.0	
# w/ 2 or more inhaled					
steroid Rx	0	0	0	0	
% w/ 2 or more inhaled steroid Rx	0.0	0.0	0.0	0.0	
	3.0			0.0	
CHANGE FROM BASELINE YR % # w/ 2 or more controller Rx	+4 5	-16 7	-83 3	+16 7	
# w/ 2 or more inhaled					
steroid Rx	-22.7	-50.0	-100.0	+0.0	

Figure 2-120: Sample Age Breakdown Report, Medication Therapy for Persons with Asthma

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient

Medication Therapy for Persons with Asthma: List of patients with asthma
with asthma medications, if any.

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

PATIENT1, GWEN 000001 COMMUNITY #1 F 5
AC, Severity 4 in V Asthma 02/02/11

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```
PATIENT2, ALICE 000002 COMMUNITY #1 F 6
AC, Severity >1 on PL for 493.00 SABA: 06/15/12 ALBUTEROL 90MCG/INHALATION MDI(4)
PATIENT3, GENEVA 000003 COMMUNITY #1 F 47
AC, Severity 2 in V Asthma 03/03/12 2+ CONT: 06/01/12 FLUTICASONE PROPIONATE 110MCG
INHALER, 10/15/12 FLUTICASONE PROPIONATE 110MCG INHALER; 2+ STEROID: 06/01/12
FLUTICASONE PROPIONATE 110MCG INHALER, 10/15/12 FLUTICASONE PROPIONATE 110MCG
INHALER
PATIENT22, MELANIE 000022 COMMUNITY #1 F 47
AC, Severity >1 on PL for 493.00
PATIENT27, RANDALL 000027 COMMUNITY #1 M 6
AC, Severity >1 on PL for 493.00 LABA2+ CONT: 03/03/12 MOMETASONE/FORMOTEROL 100/5MCG
INH, 05/01/12 MOMETASONE/FORMOTEROL 100/5MCG INH; 2+ STEROID: 03/03/12
MOMETASONE/FORMOTEROL 100/5MCG INH, 05/01/12 MOMETASONE/FORMOTEROL 100/5MCG INH
```

Figure 2-121: Sample Patient List, Medication Therapy for Persons with Asthma

# 2.10.9 Community-Acquired Pneumonia (CAP): Assessment of Oxygen Saturation

#### **Denominator**

*Number of visits for User Population patients* ages 18 and older diagnosed with community-acquired bacterial pneumonia at an outpatient visit during the Report Period.

#### **Numerators**

Number of visits where patients had oxygen saturation documented and reviewed.

Number of visits where patients refused oxygen saturation assessment.

Number of visits where patients did not have their oxygen saturation documented and reviewed.

## **Logic Description**

Age of the patient is calculated at the beginning of the report period.

If a patient has more than one visit for community-acquired bacterial pneumonia during the report period, each visit will be counted as long as it has been more than 45 days since the date of the prior visit. For example, a patient was diagnosed on January 1, 2011 and 35 days later he was diagnosed again with pneumonia. That second diagnosis does not count as a separate visit. However, if the patient were diagnosed again on February 16, 2011 (46 days after onset), that diagnosis counts as a separate visit. Because RPMS does not store the date of onset, visit date will be used as a surrogate for onset date.

CRS uses the following codes and taxonomies to define the denominator and numerators.

	ICD and Other Codes
Community-Acquired Bacterial Pneumonia (Non-CHS outpatient visit, defined as visit type not equal to "C" and service category of "A" for Ambulatory, "S" for Day Surgery, or "O" for Observation)	V POV: 481, 482.0, 482.1, 482.2, 482.30, 482.31, 482.32, 482.39, 482.40, 482.41, 482.42, 482.49, 482.81, 482.82, 482.83, 482.84, 482.89, 482.9, 483.0, 483.1, 483.8, 485, 486, 487.0

		ICD and Other Codes	LOINC Codes	Taxonomy
Oxygen Saturation Assessment (any of the arterial blood gas (ABG) or pulse oximetry tests performed at the visit)	94760-94762, 82803, 82805, 82810, or 3028F or 3028F, where 3028F has no modifier of 1P, 2P, 3P, or 8P	Saturation	Yes	BGP CMS ABG TESTS

Refusal of Oxygen Saturation Assessment definition: Patients whose oxygen saturation was not assessed due to a patient refusal of assessment on visit date. Refusal is defined as refusal of any of the tests listed above.

*No Assessment definition*: Patients whose oxygen saturation was not assessed or refused.

## **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients with community-acquired bacterial pneumonia, with oxygen saturation assessment or documented reason for no assessment, if any.

#### **Measure Source**

CMS PQRI Measure #57

## Measure Past Performance and Long-Term Targets

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	***	IHS	2012	Selected	Measures	with	Community	Specified	Report	***		

```
DEMO INDIAN HOSPITAL
               Report Period: Jan 01, 2012 to Dec 31, 2012
            Previous Year Period: Jan 01, 2011 to Dec 31, 2011
              Baseline Period: Jan 01, 2000 to Dec 31, 2000
Community-Acquired Pneumonia (CAP): Assessment of Oxygen Saturation (con't)
                  REPORT
                           % PREV YR
                                        % CHG from BASE
                                                             % CHG from
                  PERIOD
                         PERIOD
                                           PREV YR % PERIOD
                                                               BASE %
# Pneumonia Visits for
                                   12
User Pop Pts 18+
                      53
# Visits w/ O2 Sat
                     23 43.4
4 7.5
                                   1 8.3 +35.1
                                                        2 25.0
# Visits w/ Refusal
                                   0.0
                                              +7.5
                                                        0.0
                                                                   +7.5
# Visits w/ No O2
                     26 49.1
                                   11 91.7
                                              -42.6
                                                        6 75.0
                                                                  -25.9
Sat Assmt
```

Figure 2-122: Sample Report, Community-Acquired Pneumonia (CAP): Assessment of Oxygen Saturation

UP-User Pop; AC-Active Clinical; AD-Active Diabetic; AAD-Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Community-Acquired Pneumonia (CAP): Assessment of Oxygen Saturation: List of patients with community-acquired bacterial pneumonia, with oxygen saturation assessment or documented reason for no assessment, if any. SEX AGE PATTENT NAME HRN COMMUNITY DENOMINATOR NUMERATOR PATIENT1, GENEVA 000001 COMMUNITY #1 F 27

UP,1) 11/25/12 482.81 1) 11/25/12 02 SAT: CPT 3028F

PATIENT2, JACKIE 000002 COMMUNITY #1 F 29

UP,1) 06/15/12 482.89 1) 06/15/12 02 SAT: LAB BLOOD GASES

PATIENT3, PAULINE 000003 COMMUNITY #1 F 38

UP,1) 05/31/12 482.0 1) 05/31/12 None

PATIENT4, WILLIAM 000004 COMMUNITY #1 M 38 UP,1) 05/31/12 482.0; 2) 08/01/12 482.49; 3) 09/17/12 487.0 1) 05/31/12 None; 2) 08/01/12 O2 SAT: CPT 82805; 3) 09/17/12 O2 SAT: CPT 82810 PATIENT5, ZACHARY LEE 000005 COMMUNITY #1 M 36 UP,1) 09/01/12 482.83; 2) 12/01/12 482.83 1) 09/01/12 Refused CPT 94761; 2) 12/01/12 Refused O2 SAT MEAS

Figure 2-123: Sample Patient List, Community-Acquired Pneumonia (CAP): Assessment of Oxygen Saturation

## 2.10.10 Chronic Kidney Disease Assessment

#### **Denominators**

Active Clinical patients ages 18 and older with serum creatinine test during the Report Period.

*User Population patients ages 18 and older* with serum creatinine test during the Report Period.

#### **Numerators**

Patients with Estimated GFR.

- a. Patients with GFR less than (<) 60.
- b. Patients with normal GFR (i.e., >=60).

## **Logic Description**

Age is calculated at beginning of the Report Period.

For the GFR <60 numerator, CRS will include GFR results containing a numeric value less than 60 or with a text value of "<60". For the normal GFR (>=60) numerator, CRS will include GFR results containing a numeric value equal to or greater than 60 or with a text value of "60"

CRS uses the following codes and taxonomies to define the denominator and numerators.

		ICD and Other Codes	LOINC Codes	Taxonomy
Creatinine test	82540, 82565-75		Yes	DM AUDIT CREATININE TAX
Estimated GFR test			Yes	BGP GPRA ESTIMATED GFR TAX

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients with Creatinine test, with GFR and value, if any.

#### **Measure Source**

## Measure Past Performance and Long-Term Targets

None

DU November 25, 2012 Page 303  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Chronic Kidney Disea	ase Asses	sment	(con't)						
	REPORT PERIOD		PREV YR PERIOD		CHG from E			HG from BASE %	
Active Clinical Pts => 18 with Serum Creatinine test	271		258			221			
w/Est GFR A. # w/ GFR <60 B. # w/Normal GFR					+57.0 +7.6			+62.5 +12.4	
(>=60) User Pop Pts =>18 with Serum	137	50.6	2	0.8	+49.8	0	0.0	+50.6	
Creatinine	331		311			262			
# w/ Est GFR A. # w/GFR <60 B. # w/Normal GFR		66.8 16.6			+56.2 +7.0			+60.7 +10.9	
(>=60)	165	49.8	2	0.6	+49.2	0	0.0	+49.8	

Figure 2-124: Sample Report, Chronic Kidney Disease Assessment

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Chronic Kidney Disease Assessment: List of patients with Creatinine test, with GFR and value, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR -----PATIENT1, SHERISA 000001 COMMUNITY #1 F 18 08/16/12 GFR: 78 UP,AC PATIENT2, CAITLYN 000002 COMMUNITY #1 F 22 UP,AC PATIENT3, HALEY DEBRA 000003 COMMUNITY #1 F 07/09/12 GFR: >60 PATIENT4, HELENE MARIE 000004 COMMUNITY #1 F 29 UP,AC 11/20/12 GFR: 114 PATIENT5, MARTHA 000005 COMMUNITY #1 F 30

Figure 2-125: Sample Patient List, Chronic Kidney Disease Assessment

## 2.10.11 Prediabetes/Metabolic Syndrome

#### **Denominators**

Active Clinical patients ages 18 and older diagnosed with prediabetes/metabolic syndrome without a documented history of diabetes.

*User Population patients ages 18 and older* diagnosed with prediabetes/metabolic syndrome without a documented history of diabetes.

#### **Numerators**

*All Screenings:* Patients with all screenings (BP, LDL, fasting glucose or A1c, tobacco screening, BMI, lifestyle counseling, and depression screening).

*BP Assessed:* Patients with Blood Pressure documented at least twice during the Report Period.

LDL Assessed: Patients with LDL completed, regardless of result, during the Report Period.

Fasting Glucose or A1c Assessed: Patients with fasting glucose test or A1c assessed, regardless of result, during the Report Period.

Patients with A1c less than (<) 5.7.

Patients with A1c equal to or greater than (=>) 5.7 and less than (<) 6.5.

Patients with A1c equal to or greater than (=>) 6.5.

Patients with no A1c during the Report Period.

*Tobacco Use Assessed:* Patients who have been screened for tobacco use during the Report Period.

BMI Available: Patients for whom a BMI could be calculated.

**Note**: This numerator does *not* include refusals.

*Lifestyle Counseling:* Patients who have received any lifestyle adaptation counseling, including medical nutrition therapy, or nutrition, exercise or other lifestyle education during the Report Period.

Depression Screening: Patients screened for depression or diagnosed with a mood disorder at any time during the Report period, including documented refusals in past year.

## **Logic Description**

Age is calculated at beginning of the Report Period.

Prediabetes/Metabolic Syndrome defined as:

- 1. Diagnosis of prediabetes/metabolic syndrome, defined as: Two visits during the Report Period with POV 277.7, *or*
- 2. One each of at least three different conditions listed below, occurring during the Report Period except as otherwise noted:
  - BMI => 30 *or* Waist Circumference >40 inches for men or >35 inches for women
  - Triglyceride value >=150
  - HDL value <40 for men or <50 for women
  - Patient diagnosed with hypertension *or* mean BP value => 130/85 where systolic is =>130 *or* diastolic is =>85
  - Fasting Glucose value =>100 and <126

**Note:** Waist circumference and fasting glucose values will be checked last.

Definition for patients without diabetes: No diabetes diagnosis ever (POV 250.00–250.93).

*BMI definition*: CRS calculates BMI at the time the report is run, using NHANES II. For 18 and under, a height and weight must be taken on the same day any time during the Report Period. For 19 through 50, height and weight must be recorded within last five years, not required to be on the same day. For over 50, height and weight within last two years, not required to be recorded on same day.

Hemoglobin A1c definition: Searches for most recent A1c test with a result during the Report Period. If more than one A1c test is found on the same day and/or the same visit and one test has a result and the other does not, the test with the result will be used. If an A1c test with a result is not found, CRS searches for the most recent A1c test without a result. Without result is defined as A1c documented but with no value.

BP definition: CRS uses mean of last three BPs documented on non-ER visits during the Report Period. If three BPs are not available, uses mean of last two non-ER BPs. If a visit contains more than one BP, the lowest BP will be used, defined as having the lowest systolic value. The mean Systolic value is calculated by adding the last three (or two) systolic values and dividing by three (or two). The mean Diastolic value is calculated by adding the diastolic values from the last three (or two) blood pressures and dividing by three (or two).

For the BP documented numerator, if CRS is not able to calculate a mean BP, it will search for CPT 0001F, 2000F, 3074F–3080F or POV V81.1 documented on non-ER visit during the Report Period.

Hypertension: Diagnosis of (POV or problem list) 401.\* occurring prior to the Report Period, and at least one hypertension POV during the Report Period.

CRS uses the following codes and taxonomies to define the denominator and numerators.

Test	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Triglyceride (requires a non-null, numeric result)			Yes	DM AUDIT TRIGLYCERIDE TAX
HDL (requires a non- null, numeric result)			Yes	DM AUDIT HDL TAX
Fasting Glucose- Denominator Definition (requires a non-null, numeric result)			Yes	DM AUDIT FASTING GLUCOSE TESTS
Fasting Glucose- Numerator Definition		<b>V POV</b> : 790.21	Yes	DM AUDIT FASTING GLUCOSE TESTS
Hemoglobin A1c	83036, 83037, 3044F-3046F, 3047F (old code)		Yes	DM AUDIT HGB A1C TAX
LDL	80061, 83700, 83701, 83704, 83715 (old code), 83716 (old code), 83721, 3048F, 3049F, 3050F		Yes	DM AUDIT LDL CHOLESTEROL TAX
Estimated GFR			Yes	BGP GPRA ESTIMATED GFR TAX
Quantitative Urinary Protein Assessment	82042, 82043, or 84156		Yes	BGP QUANT URINE PROTEIN

Test	ICPT COMPS	ICD and Other Codes	LOINC Codes	Taxonomy
End Stage Renal Disease	36145, 36147, 36800, 36810, 36815, 36818, 36819, 36820, 36821, 36831- 36833, 50300, 50320, 50340, 50360, 50365, 50370, 50380, 90951-90970 or old codes 90918- 90925, 90935, 90937, 90939 (old code), 90940, 90945, 90947, 90989, 90993, 90997, 90999, 99512, G0257, G0308-G0327 (old codes), G0392 (old code), G0393 (old code), or S9339	V POV: 585.5, 585.6, V42.0, V45.1 (old code), V45.11, V45.12, or V56.* V Procedure: 38.95, 39.27, 39.42, 39.43, 39.53, 39.93-39.95, 54.98, or 55.6*		

Test	ILPI LAMAS	ICD and Other Codes	LOINC Codes	Taxonomy
Tobacco Screening	D1320, 99406, 99407, G0375 (old code), G0376 (old code), 1034F, 1035F, 1036F, 1000F, G8455- G8457 (old codes), G8402 (old code) or G8453 (old code)	Any health factor for category Tobacco, TOBACCO (SMOKING), TOBACCO (SMOKELESS – CHEWING/DIP), or TOBACCO V POV or current Active Problem List: 305.1, 305.1* (old codes), 649.00-649.04, V15.82 Patient education codes: containing "TO-" or "-TO" or "-SHS", 305.1, 305.1* (old codes), 649.00-649.04, V15.82, D1320, 99406, 99407, G0375 (old code), G0376 (old code), G8455-G8457 (old codes), G8402 (old code) or G8453 (old code) Dental code: 1320		
Lifestyle Counseling – Medical Nutrition Counseling	97802-97804, G0270, G0271	Primary or secondary provider codes: 07, 29 Clinic codes: 67 (dietary) or 36 (WIC)		
Lifestyle Counseling – Nutrition Education		V POV: V65.3 dietary surveillance and counseling <i>or</i> Patient education codes: ending "-N" (nutrition) or "-MNT" (medical nutrition therapy) (or old code "-DT" (diet)) or containing V65.3, 97802-97804, G0270, or G0271.		

Test	CPT Codes	ICD and Other Codes	LOINC Codes	Taxonomy
Lifestyle Counseling – Exercise Education		V POV: V65.41 exercise counseling or Patient education codes: ending "-EX" (exercise) or containing V65.41.		
Lifestyle Counseling – Related Exercise and Nutrition Education		Patient education codes: ending "-LA" (lifestyle adaptation) or containing "OBS-" (obesity) or 278.00 or 278.01.		
Depression Screening		V Exam: Exam Code 36 V POV: V79.0 CPT: 1220F BHS Problem Code: 14.1 (Screening for Depression) V Measurement in PCC or BHS: PHQ2 or PHQ9 Refusals: Exam Code 36		
Mood Disorders		At least two visits in PCC or BHS during Report Period for: Major Depressive Disorder, Dysthymic Disorder, Depressive Disorder NOS, Bipolar I or II Disorder, Cyclothymic Disorder NOS, Mood Disorder Due to a General Medical Condition, Substance-induced Mood Disorder, or Mood Disorder NOS (see codes below). V POV: 296.*, 291.89, 292.84, 293.83, 300.4, 301.13, or 311 BHS POV: 14, 15		

## **Key Logic Changes from CRS Version 11.1**

- Removed nephropathy assessment measure from topic and from All Screenings measure.
- Updated fasting glucose measures to fasting glucose *or* A1c.
- Added new A1c measures and corresponding logic.
- Added POV V81.1 to BP Documented definition.

## **Patient List Description**

List of patients age 18 and older with Prediabetes/Metabolic Syndrome with assessments received, if any. The denominator column displays the condition the patient met, either the diagnosis of 277.7 or the three conditions the patient met (e.g. BMI=35,TG=155,HDL=35).

#### **Measure Source**

"IHS Guidelines for Care of Adults with Prediabetes and/or the Metabolic Syndrome in Clinical Settings (April 2005)"

## Measure Past Performance and Long-Term Targets

November 25, 2012 Page 309  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Prediabetes/Metabolio	c Syndro	ome							
	REPORT PERIOD	%	PREV YR PERIOD		CHG from PREV YR %			CHG from BASE %	
Active Clinical Pts = w/PreDiabetes/ Met Syn	=>18		40			29			
# w/ All screenings	1	2.0	0	0.0	+2.0	0	0.0	+2.0	
# w/ BP documented	51	100.0	34	85.0	+15.0	27	93.1	+6.9	
<pre># w/LDL done # w/ fasting</pre>	34	66.7	27	67.5	-0.8	18	62.1	+4.6	
glucose or Alc	12	23.5	8	20.0	+3.5	2	6.9	+16.6	
# w/A1c <5.7 # w/A1c =>5.7	4	7.8	6	15.0	-7.2	1	3.4	+4.4	
and <6.5	5	9.8	1	2.5	+7.3	0	0.0	+9.8	
# w/ A1c =>6.5		3.9		0.0				+0.5	
# w/no Alc	40	78.4	33	82.5	-4.1	27	93.1	-14.7	
# w/Tobacco Screening	3								
w/in 1 yr	48	94.1	33	82.5	+11.6	21	72.4	+21.7	

# w/BMI calculated									
-No Refusals	48	94.1	40	100.0	-5.9	29	100.0	-5.9	
<pre># w/lifestyle adaptation</pre>									
counseling	24	47.1	15	37.5	+9.6	8	27.6	+19.5	
# w/Depression screening,									
DX, or refusal	10	19.6	1	2.5	+17.1	1	3.4	+16.2	

Figure 2-126: Sample Report, Prediabetes/Metabolic Syndrome

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Prediabetes/Metabolic Syndrome: List of patients 18 and older with
Prediabetes/Metabolic Syndrome with assessments received, if any.
PATIENT NAME HRN COMMUNITY
DENOMINATOR NUMERATOR
PATIENT NAME
                            COMMUNITY
                                            SEX AGE
PATIENT1, HALEY DEBRA 000001 COMMUNITY #1 F 25
UP, AC, BMI=33.64; TRIG=271; HDL=45.4; HTN DX: 01/13/12 2 BPs; TOB: 01/13/12 305.1-DP;
BMI: 33.64; LIFE: 12/12/12 97804-HM;
PATIENT2, CYNTHIA 000002 COMMUNITY #1 F 36
UP,AC,TRIG=166; HDL=38.7; BP=131/80 2 BPs; LDL: 11/21/12 125; TOB: 07/09/12 NEVER
SMOKED; BMI: 28.35;
PATIENT3,ABIGAIL 000003 COMMUNITY #1 F 39
UP,BMI=34.97; TRIG=194; BP=149/84 2 BPs; LDL: 06/12/12 CPT 83704; TOB: 05/14/12
CURRENT SMOKER, STATUS UNKNOWN; BMI: 34.97; DEPR: 11/03/12 POV 296.7 + 11/30/12 POV
PATIENT4, ANNA LINDA 000004 COMMUNITY #1 F 44
UP,AC,277.7 on 03/15/12; 04/15/12 2 BPs;
PATIENT5, DARLENE T 000005 COMMUNITY #1 F 54
UP, AC, BMI=35.37; TRIG=276; HDL=35.6 (ALL:) 2 BPs; LDL: 10/03/12 106; A1C: 5.2; TOB:
10/03/12 NEVER SMOKED; BMI: 35.37; LIFE: 06/06/12 Prv 29; DEPR: DEP SCRN 05/03/12
```

Figure 2-127: Sample Patient List, Prediabetes/Metabolic Syndrome

# 2.10.12 Proportion of Days Covered by Medication Therapy

#### **Denominators**

Active Clinical patients ages 18 and older who had two or more prescriptions for beta-blockers during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for ACEI/ARBs during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for calcium channel blockers (CCB) during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for biguanides during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for sulfonylureas during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for thiazolidinediones during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for statins during the Report Period.

Active Clinical patients ages 18 and older who had two or more prescriptions for antiretroviral agents during the Report Period.

#### **Numerators**

Patients with proportion of days covered (PDC) >=80% during the Report Period.

Patients with a gap in medication therapy >=30 days.

Patients with proportion of days covered (PDC) >=90% during the Report Period.

## **Logic Description**

Age is calculated at the beginning of the report period.

To be included in the denominator, patients must have at least two prescriptions for that particular type of medication on two unique dates of service at any time during the Report Period. Medications must not have a comment of RETURNED TO STOCK.

The Index Prescription Start Date is the date when the medication was first dispensed within the Report Period. This date must be greater than 90 days from the end of the Report Period to be counted in the denominator.

The medications in the measures are defined with medication taxonomies: BGP PQA BETA BLOCKER MEDS, BGP PQA ACEI ARB MEDS, BGP PQA CCB MEDS, BGP PQA BIGUANIDE MEDS, BGP PQA SULFONYLUREA MEDS, BGP PQA THIAZOLIDINEDIONE MEDS, BGP PQA STATIN MEDS, BGP PQA ANTIRETROVIRAL MEDS.

For each PDC numerator:

Proportion of days covered = # of days the patient was covered by at least one drug in the class / # of days in the patient's measurement period.

#### Measurement Period definition:

The patient's measurement period is defined as the number of days between the Index Prescription Start Date and the end of the Report Period. When calculating the number of days the patient was covered by at least one drug in the class, if prescriptions for the same drug overlap, the prescription start date for the second prescription will be adjusted to be the day after the previous fill has ended.

Note: If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e. visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4.

### **Example of Proportion of Days Covered:**

- Report Period: Jan 1 Dec 31, 2011
- 1st Rx is Index Rx Start Date: 3/1/11, Days Supply=90
- Rx covers patient through 5/29/11
- 2nd Rx: 5/26/11, Days Supply=90
- Rx covers patient through 8/27/11
- 3rd Rx: 9/11/11, Days Supply=180
- Gap = (9/11/11 8/27/11) = 15 days
- Rx covers patient through 3/8/12
- Patient's measurement period: 3/1/11 through 12/31/11 = 306 Days
- Days patient was covered: 3/1/11 through 8/27/11 + 9/11/11 through 12/31/11 = 292 Days
- PDC = 292 / 306 = 95%

#### For each Gap numerator:

CRS will calculate whether a gap in medication therapy of 30 or more days has occurred between each consecutive medication dispensing event during the Report Period. A gap is calculated as the days not covered by the days supply between consecutive medication fills.

#### **Example of Medication Gap >=30 Days:**

- Report Period: Jan 1 Dec 31, 2011
- 1st Rx: 4/1/11, Days Supply=30

- Rx covers patient through 4/30/11
- 2nd Rx: 7/1/11, Days Supply=90
- Gap #1 = (7/1/11 4/30/11) = 61 days
- Rx covers patient through 9/28/11
- 3rd Rx: 10/1/11, Days Supply=90
- Gap #2 = (10/1/11 9/28/11) = 2 days
- Rx covers patient through 12/29/11
- Gap #1 >=30 days, therefore patient will be included in the numerator for that medication.

## **Key Logic Changes from CRS Version 11.1**

None

## **Patient List Description**

List of patients 18 and older prescribed medication therapy medication with proportion of days covered and gap days.

### **Measure Source**

PQA (Pharmacy Quality Alliance)

## **Measure Past Performance and Long-Term Targets**

DU November 25, 2012 Page 314  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
REI	Proportion of Days Covered by Medication Therapy  REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %								
Active Clinical Pts w/beta-blockers	53	44	37						
	31 58.5 27 50.9	24 54.5 +3.9 23 52.3 -1.3							
Active Clinical Pts w/ACEI/ARBs	97	80	74						

# w/ PDC >=80% # w/ gap >=30 days	60 45	61.9 46.4	55 32	68.8 40.0	-6.9 +6.4		52.7 56.8	+9.2 -10.4	
Active Clinical Pts w/ CCBs	56		48			55			
# w/ PDC >=80% # w/ gap >=30 days	34 30	60.7 53.6		62.5 41.7	-1.8 +11.9	37 22	67.3 40.0		
Active Clinical Pts w/ biguanides	30		26			11			
# w/ PDC >=80% # w/ gap >=30 days		40.0 63.3		69.2 38.5	-29.2 +24.9	3 8	27.3 72.7	+12.7 -9.4	
Active Clinical Pts w/ sulfonylureas	7		6			7			
# w/ PDC >=80% # w/ gap >=30 days	3 5	42.9 71.4		16.7 83.3	+26.2 -11.9	4 3	57.1 42.9	-14.3 +28.6	
Active Clinical Pts w/ thiazolidinediones	20		15			4			
# w/ PDC >=80% # w/ gap >=30 days	13	65.0 40.0		66.7 46.7	-1.7 -6.7	2 1	50.0 25.0	+15.0 +15.0	
Active Clinical Pts w/ statins	60		49			37			
# w/ PDC >=80% # w/ gap >=30 days	43 24	71.7 40.0		67.3 38.8	+4.3 +1.2	25 16	67.6 43.2	+4.1 -3.2	
Active Clinical Pts w/ antiretrovial agents		2	0			1			
# w/ PDC >=90%	1	50.0	0	0.0	+50.0	0	0.0	+50.0	

Figure 2-128: Sample Report, Proportion of Days Covered by Medication Therapy

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Proportion of Days Covered by Medication Therapy: List of patients 18 and older prescribed medication therapy medication with proportion of days covered and gap days. HRN COMMUNITY PATIENT NAME SEX AGE DENOMINATOR NUMERATOR Patient75,PAULA KAY 000075 COMMUNITY #1 F 34 CCB: IXRD: 03/24/12 [282] Days=228 >80 GAP=52 AC CCB: IXRD: 03/24/12 [2 Patient76, CRSCT 000076 COMMUNITY #1 F 36 CCB: IXRD: 06/01/12 [213] Days=180 >80 GAP=31 Patient77, CRSAC 000077 COMMUNITY #1 F 44 ACEI/ARB: IXRD: 06/05/12 [209] Days=60 <80 GAP=118

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```
Patient78, DEBORA ELLEN 000078 COMMUNITY #1 F 45

AC BB: IXRD: 07/23/12 [161] Days=126 <80; ACEI/ARB: IXRD: 01/29/12 [336] Days=272 >80

Patient79, STELLA LYNN 000079 COMMUNITY #1 F 46

AC BB: IXRD: 01/21/12 [344] Days=299 >80 GAP=38; CCB: IXRD: 01/21/12 [350] Days=218 <80 GAP=70; ACEI/ARB: IXRD: 01/15/12 [350] Days=314 >80; CCB: IXRD: 01/15/12 [350] Days=218 <80 GAP=103

Patient81, CRSNK 000081 COMMUNITY #1 F 51

AC SULF: IXRD: 09/01/12 [121] Days=120 >80
```

Figure 2-129: Sample Patient List, Proportion of Days Covered by Medication Therapy

### 2.10.13 Medications Education

#### **Denominators**

Active Clinical patients with medications dispensed at their facility during the Report Period.

All *User Population patients with medications* dispensed at their facility during the Report Period.

#### **Numerator**

Patients who were provided patient education about their medications in any location.

Patients who refused patient education about their medications in any location.

## **Logic Description**

Patients receiving medications at their facility are identified by any entry in the VMed file for your facility. The purpose of this definition is to ensure that sites are not being held responsible for educating patients about medications received elsewhere that may be recorded in RPMS. CRS assumes that the appropriate facility is the one the user has logged onto to run the report.

**Note:** If a site's system identifier, i.e., ASUFAC code, has changed during the period between the Baseline start date and the Current Year end date, due to compacting/contracting or other reasons, your report may display zeros (0s) or very low counts for some time periods.

CRS uses the following patient education codes to define the numerator:

Medication Education	Any Patient Education code containing "M-" or "-M" (medication)				
	or				
	DMC-IN (Diabetes Medicine–Insulin)				
	FP-DPO (Family Planning–Depot Medroxyprogesterone Injections				
	FP-OC (Family Planning-Oral Contraceptives)				
	FP-TD (Family Planning-Transdermal (Patch))				
	*-NEB (*Nebulizer)				
	*-MDI (*Metered Dose Inhalers)				
Refusal	Any refusal in the past year with Patient Education codes containing "M-" or "-M" or PFE codes DMC-IN, FP-DPO, FP-OC, *-NEB, *-MDI, or FP-TD				
	In the past year, any Patient Education code containing "M-" or "-M" or PFE codes DMC-IN, FP-DPO, FP-OC, *-NEB, *-MDI, or FP-TD with a level of understanding of "refused".				

# **Key Logic Changes from CRS Version 11.1**

Changed the logic to include all –NEB and –MDI codes.

### **Patient List Description**

List of patients receiving medications with med education, if any.

### **Measure Source**

None

### Measure Past Performance and Long-Term Targets

Measure	Target
IHS 2020 Goal	75.0%

```
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                Report Period: Jan 01, 2012 to Dec 31, 2012
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               Baseline Period: Jan 01, 2000 to Dec 31, 2000
Medications Education (con't)
                   REPORT % PREV YR % CHG from BASE % CHG from
                                            PREV YR % PERIOD
                              PERIOD
                                                                   BASE %
                   PERIOD
Active Clinical Pts receiving
medications
                                    631
                                                          592
```

<pre># receiving   medication educ # refusals</pre>	490 3	64.7 0.4	268 0	42.5	+22.3	81 0	13.7	+51.0 +0.4	
User Pop Pts receiving medications	981		800			753			
<pre># receiving   medication educ # refusals</pre>	599 3	61.1	306 0	38.3	+22.8 +0.3	87 0	11.6	+49.5 +0.3	

Figure 2-130: Sample Report, Medications Education

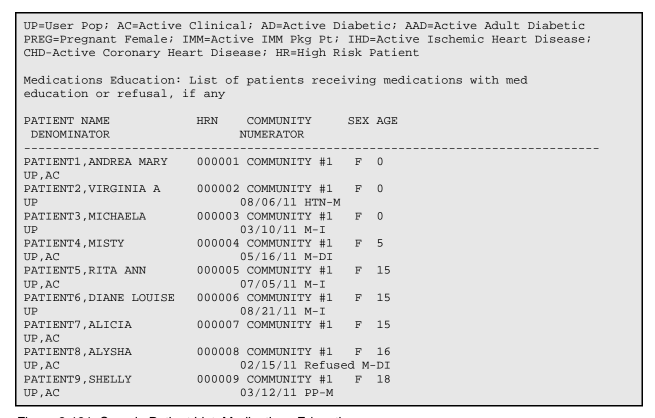


Figure 2-131: Sample Patient List, Medications Education

# 2.10.14 Medication Therapy Management Services

### **Denominators**

Active Clinical patients =>18 with medications dispensed at their facility during the Report Period.

#### **Numerator**

Patients who received medication therapy management (MTM) during the Report Period.

# **Logic Description**

Age is calculated at the beginning of the report period.

Patients receiving medications at their facility are identified by any entry in the VMed file for your facility.

*Medication Therapy Management (MTM) definition*: 1) CPT codes 99605-99607 or 2) Clinic codes: D1, D2.

### **Key Logic Changes from CRS Version 11.1**

New topic.

### **Patient List Description**

List of patients >=18 receiving medications with medication therapy management, if any.

### **Measure Source**

None

# **Measure Past Performance and Long-Term Targets**

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                           DEMO INDIAN HOSPITAL
                Report Period: Jan 01, 2012 to Dec 31, 2012
            Previous Year Period: Jan 01, 2011 to Dec 31, 2011
              Baseline Period: Jan 01, 2000 to Dec 31, 2000
Medications Therapy Management Services (con't)
                   REPORT
                            % PREV YR % CHG from BASE
                                                               % CHG from
                   PERIOD PERIOD PREV YR % PERIOD
                                                                  BASE %
Active Clinical Pts
=>18 receiving
medications
                      594
                                    472
                                                         424
# w/MTM
                       5 0.8
                                     0.0
                                                +0.8
                                                         0
                                                              0.0
                                                                     +0.8
```

Figure 2-132: Sample Report, Medications Therapy Management Services

Figure 2-133: Sample Patient List, Medications Therapy Management Services

# 2.10.15 Self Management (Confidence)

#### **Denominators**

Active Clinical patients assessed for confidence in managing their health problems during the Report Period.

#### **Numerator**

Patients who are very confident in managing their health problems during the Report Period.

### **Logic Description**

Confidence in managing health problems definition: Any health factor for category CONFIDENCE IN MANAGING HEALTH PROBLEMS.

*Very confident definition*: The most recent health factor in the CONFIDENCE IN MANAGING HEALTH PROBLEMS category of VERY SURE.

### **Key Logic Changes from CRS Version 11.1**

New topic.

### **Patient List Description**

List of patients who are confident in managing their health problems.

#### **Measure Source**

None

### **Measure Past Performance and Long-Term Targets**

Measure	Target
IHS 2020 Goal	75.0%

```
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Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000

Self Management (Confidence) (con't)

REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %

Active Clinical Pts assessed for confidence 8 1 0

# very confident 4 50.0 0 0.0 +50.0 0 0.0 +50.0
```

Figure 2-134: Sample Report, Medications Education

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Self Management (Confidence): List of patients who are confident in managing their health problems. DENOMINATOR PATIENT NAME HRN COMMUNITY SEX AGE NUMERATOR PATIENT5, RITA ANN 000005 COMMUNITY #1 F 18 06/01/12 PATIENT6, DIANE LOUISE 000006 COMMUNITY #1 F 20 07/28/12 PATIENT7, ALICIA 000007 COMMUNITY #1 F 21 PATIENT8, ALYSHA 000008 COMMUNITY #1 F 25 AC

Figure 2-135: Sample Patient List, Medications Education

# 2.10.16 Public Health Nursing

#### **Patient-Related Measures**

#### **Denominator**

All User Population patients.

#### **Numerators**

For User Population only, the number of patients in the denominator served by Public Health Nurses (PHNs) in any setting, including Home.

For User Population only, the number of patients in the denominator served by a PHN driver/interpreter in any setting.

For User Population only, the number of patients in the denominator served by PHNs in a HOME setting.

For User Population only, the number of patients in the denominator served by a PHN driver/interpreter in a HOME Setting.

### **Visit-Related Measures**

#### **Denominators**

Number of visits to User Population patients by PHNs in any setting, including Home

- Number of visits to patients ages 0–28 days (Neonate) in any setting.
- Number of visits to patients ages 29 days—12 months (infants) in any setting.
- Number of visits to patients ages 1–64 years in any setting
- Number of visits to patients ages 65 and older (Elders) in any setting
- Number of PHN driver/interpreter (Provider Code 91) visits

Number of visits to User Population patients by PHNs in Home setting

- Number of Home visits to patients age 0–28 days (Neonate)
- Number of Home visits to patients age 29 days to 12 months (Infants)
- Number of Home visits to patients ages 1–64 years
- Number of Home visits to patients aged 65 and over (Elders).
- Number of PHN driver/interpreter (Provider Code 91) visits in a HOME setting.

### **Numerator**

No numerator: count of visits only

### **Logic Description**

PHN visit is defined as any visit with primary or Secondary Provider Code 13 or 91. Home visit defined as: (1) Clinic 11 and a primary or Secondary Provider Code 13 or 91 or (2) Location Home (as defined in Site Parameters) and a primary or Secondary Provider Code 13 or 91.

# **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients with PHN visits documented.

Numerator codes in patient list: All PHN = Number of PHN visits in any setting; Home = Number of PHN visits in home setting; Driver All = Number of PHN driver/interpreter visits in any setting; Driver Home = Number of PHN driver/interpreter visits in home setting.

### **Measure Source**

None

### Measure Past Performance and Long-Term Targets

	All PHN visits	PHN Home Visits
IHS FY 2005 Performance	438,376	Not Reported
IHS FY 2004 Performance	423,379	192,121
IHS FY 2003 Performance	359,089	160,650
IHS FY 2002 Performance	343,874	156,263

```
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                 Report Period: Jan 01, 2012 to Dec 31, 2012
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                Baseline Period: Jan 01, 2000 to Dec 31, 2000
Public Health Nursing (con't)
                                            % CHG from BASE
                    REPORT
                              % PREV YR
                                                                   % CHG from
                    PERIOD
                                 PERIOD
                                               PREV YR % PERIOD
                                                                     BASE %
All User Population
                     2,896
                                  2,456
patients
                                                          2,346
# served by PHNs in
any Setting
                        13
                             0.4
                                      13 0.5
                                                   -0.1
                                                             13
                                                                  0.6
                                                                          -0.1
# served by PHN drivers/
```

interpreter - in any Setting	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
<pre># served by PHNs in a Home Setting # served by PHN</pre>	1	0.0	1	0.0	+0.0	0	0.0	+0.0	
drivers/interpreters in Home Setting	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Total # PHN Visits - Any Setting	18		16		+2	19		-1	
A. Ages 0-28 days	0		0		+0	0		+0	
B. Ages 29 days - 12 months	1		3		-2	0		+1	
C. Ages 1-64 years	16		13		+3	19		-3	
D. Ages 65+	1		0		+1	0		+1	
E. Driver/Interpreter visits - any setting	0		0		+0	0		+0	
Total # PHN Visits - Home Setting	3		1		+2	0		+3	
A. Ages 0-28 days	0		0		+0	0		+0	
B. Ages 29 days- 12 months	0		1		-1	0		+0	
C. Ages 1-64 years	3		0		+3	0		+3	
D. Ages 65+	0		0		+0	0		+0	
E. Driver/interpreter visits -									
Home Setting	0		0		+0	0		+0	

Figure 2-136: Sample Report, Public Health Nursing

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Public Health Nursing: List of patients with PHN visits documented

PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR NUMERATOR

PATIENT1, HELENE MARIE 000001 COMMUNITY #1 F 29

UP 2 all PHN; 0 home; 0 driver all; 0 driver home PATIENT2, KATHLEEN 000002 COMMUNITY #1 F 38

UP 3 all PHN; 3 home; 0 driver all; 0 driver home

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PATIENT40,ERIKA SUE	000040 COMMUNITY #2 F 37
UP	1 all PHN; 0 home; 0 driver all; 0 driver home
PATIENT41, DANIEL RAY	000041 COMMUNITY #2 M 0
UP	1 all PHN; 0 home; 0 driver all; 0 driver home

Figure 2-137: Sample Patient List, Public Health Nursing

# 2.10.17 Breastfeeding Rates

#### **Denominators**

Active Clinical patients who are 30-394 days old

Active Clinical patients who are 30–394 days old who were screened for infant feeding choice at the age of *two months* (45–89 days) (PART Denominator)

Active Clinical patients who are 30–394 days old who were screened for infant feeding choice at the age of six months (165–209 days)

Active Clinical patients who are 30–394 days old who were screened for infant feeding choice at the age of *nine months* (255–299 days)

Active Clinical patients who are 30–394 days old who were screened for infant feeding choice at the age of *one year* (350–394 days)

#### **Numerators**

Patients who were screened for infant feeding choice at least once

Patients who were screened for infant feeding choice at the age of *two months* (45–89 days)

Patients who were screened for infant feeding choice at the age of *six months* (165–209 days)

Patients who were screened for infant feeding choice at the age of *nine months* (255–299 days)

Patients who were screened for infant feeding choice at the age of *one year* (350–394 days)

Patients who, at the age of *two months* (45–89 days), were either exclusively or mostly breastfed (PART Numerator)

Patients who, at the age of *six months* (165–209 days), were either exclusively or mostly breastfed

Patients who, at the age of *nine months* (255–299 days), were either exclusively or mostly breastfed

Patients who, at the age of *one year* (350–394 days), were either exclusively or mostly breastfed

### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

Infant feeding choice definition: The documented feeding choice from the file V Infant Feeding Choice that is closest to the exact age that is being assessed will be used. For example, if a patient was assessed at 45 days old as half breastfed and half formula and assessed again at 65 days old as mostly breastfed, the mostly breastfed value will be used since it is closer to the exact age of two months (i.e., 60 days). Another example is a patient who was assessed at 67 days as mostly breastfed and again at 80 days as mostly formula. In this case, the 67 days value of mostly breastfed will be used. The other exact ages are 180 days for six months, 270 days for nine months, and 365 days for one year.

In order to be included in the age-specific screening numerators, the patient must have been screened at the specific age range. For example, if a patient was screened at six months and was exclusively breastfeeding but was not screened at two months, then the patient will only be counted in the six months numerator.

# **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients 30–394 days old, with infant feeding choice value, if any.

Note:	"DO" represents "Days Old."	

### **Measure Source**

HP 2020, MICH-21.4 Exclusive breastfeeding-through three months, MICH-21.5 Exclusive breastfeeding-through six months.

### Measure Past Performance and Long-Term Targets

Performance	Percent
IHS FY 2011 Performance	26.7%
IHS FY 2010 Performance	33%
IHS FY 2008 Performance	28%
HP 2020 goal for breastfeeding through 3 months of age	44.3%

Performance	Percent
HP 2020 goal for breastfeeding through 6 months of age	23.7%

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	Previous Year Period: Jan 01, 2011 to Dec 31, 2011 Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Breastfeeding Rates										
	REPORT PERIOD				CHG from PREV YR %					
Active Clinical Pts 30-394 days	46		30			34				
<pre># w/infant feeding   choice screening # w/screening @</pre>	11	23.9	0	0.0	+23.9	1	2.9	+21.0		
2 mos # w/screening @	4	8.7	0				2.9			
6 mos # w/screening @	3	6.5	0				0.0			
9 mos # w/screening @ 1 yr	3	8.7 6.5	0		+8.7		0.0			
AC Pts 30-394 days screened @ 2 mos (PART)	4	0.3	0	0.0	.0.3	1	0.0	.0.3		
# @ 2 mos exclusive/ mostly breastfed (PART)		100.0	0	0.0	+100.0	1	100.0	+0.0		
AC Pts 30-394 days screened @ 6 mos	3		0			0				
# @ 6 mos exclusive/mostly breastfed	2	66.7	0	0.0	+66.7	0	0.0	+66.7		
AC Pts 30-394 days screened at 9 mos	4		0			0				
# @ 9 mos exclusive/mostly breastfed	3	75.0	0	0.0	+75.0	0	0.0	+75.0		
AC Pts 30-394 days screened @ 1 yr	3		0			0				
# @ 1 year exclusive/mostly										

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breastfed	2	66.7	0	0.0	+66.7	0	0.0	+66.7	
-----------	---	------	---	-----	-------	---	-----	-------	--

Figure 2-138: Sample Report, Breastfeeding Rates

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Breastfeeding Rates: List of patients 30-394 days old, with infant feeding
choice value, if any.
                HRN COMMUNITY SEX AGE
PATIENT NAME
 DENOMINATOR
                          NUMERATOR
PATIENT1, AMANDA DEBRA 000001 COMMUNITY #1 F 0
PATIENTI, APRILIDITA

AC

Scrn: 33 DU, 02/05/11

PATIENT2, LEROY JAMES

000002 COMMUNITY #1 M 1

Scrn: 2 MOS: 48 DO, 01/20/11 EXCLUSIVE BREASTFEEDING;

Scrn: 2 MOS: 48 DO, 09/05/11 EXCLUSIVE
BREASTFEEDING; 1 YR: 382 DO, 12/20/11 MOSTLY BREASTFEEDING
PATIENT3, TERRY SCOTT 000003 COMMUNITY #1 M 0
PATIENT4, ROBERT 000004 COMMUNITY #1 M 0
AC Scrn: 6 MOS: 187 DO,
                           Scrn: 6 MOS: 187 DO, 08/11/11 EXCLUSIVE BREASTFEEDING
PATIENT11, STEVEN CODY 000011 COMMUNITY #2 M 0
                    Scrn: 2 MOS: 60 DO, 11/03/11 MOSTLY BREASTFEEDING
```

Figure 2-139: Sample Patient List, Breastfeeding Rates

# 2.10.18 Use of High-Risk Medications in the Elderly

### **Denominators**

Active Clinical patients ages 65 and older. Broken down by gender and age groups (65-74, 75-84, >85).

*User Population* patients ages 65 and older. Broken down by gender.

### **Numerators**

Patients who received at least one high-risk medication for the elderly during the Report Period. (GPRA Developmental Numerator)

Patients who received at least two different high-risk medications for the elderly during the Report Period. (GPRA Developmental Numerator)

### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

*Medication definitions*: High-risk medications for the elderly (i.e., potentially harmful drugs) defined with medication taxonomies:

- BGP HEDIS ANTIANXIETY MEDS (Includes combination drugs) (Aspirin-Meprobamate, Meprobamate)
- BGP HEDIS ANTIEMETIC MEDS ((Scopolamine, Trimethobenzamide)
- BGP HEDIS ANALGESIC MEDS (Includes combination drugs)
   (Acetaminophen-diphenhydramine, diphenhydramine-magnesium salicylate, Ketorolac)
- BGP HEDIS ANTIHISTAMINE MEDS (Includes combination drugs) (APAP/dextromethorphan/diphenhydramine,

APAP/diphenhydramine/phenylephrine,

APAP/diphenhydramine/pseudoephedrine, Acetaminophen-diphenhydramine, Atropine/CPM/hyoscyamine/PE/PPA/scopolamine,

Carbetapentane/diphenhydramine/phenylephrine,

Codeine/phenylephrine/promethazine, Codeine-promethazine, Cyproheptadine,

Dexchlorpheniramine, Dexchlorpheniramine/dextromethorphan/PSE,

Dexchlorpheniramine/guaifenesin/PSE,

Dexchlorpheniramine/hydrocodone/phenylephrine,

Dexchlorpheniramine/methscopolamine/PSE, Dexchlorpheniramine-pseudoephedrine, Dextromethorphan-promethazine, Diphenhydramine, Diphenhydramine/hydrocodone/phenylephrine, Diphenhydramine-magnesium salicylate, Diphenhydramine-phenylephrine, Diphenhydramine-pseudoephedrine, Hydroxyzine hydrochloride, Hydroxyzine pamoate, Phenylephrine-promethazine, Promethazine, Tripelennamine)

- BGP HEDIS ANTIPSYCHOTIC MEDS (Thioridazine, Mesoridazine)
- BGP HEDIS AMPHETAMINE MEDS (Amphetamine-destroamphetamine, Benzphetamine, Dexmethylphenidate, Dextroamphetamine, Diethylproprion, Methamphetamine, Methylphenidate, Pemoline, Phendimetrazine, Phenteramine)
- BGP HEDIS BARBITURATE MEDS (Amobarbital, Butabarbital, Mephobarbital, Pentobarbital, Phenobarbital, Secobarbital)
- BGP HEDIS BENZODIAZEPINE MEDS (Includes combination drugs)
   (Amitriptyline-Chlordiazepoxide, Chlordiazepoxide, Chlordiazepoxide-clidinium, Diazepam, Flurazepam)
- BGP HEDIS CALCIUM CHANNEL MEDS (Nifedipine short acting only)
- BGP HEDIS GASTRO ANTISPASM MED (Dicyclomine, Propantheline)

- BGP HEDIS BELLADONNA ALKA MEDS (Includes combination drugs)
   (Atropine, Atropine/CPM/hyoscyamine/PE/scopolamine,
   Atropine/hyoscyamine/PB/scopolamine, Atropine-difenoxin, Atropine-diphenoxylate, Atropine-edrophonium, Belladonna,
   Belladonna/caffeine/ergotamine/pentobarbital,
   Belladonna/ergotamine/phenobarbital,
   Butabarbital/hyoscyamine/phenazopyridine, Digestive enzymes/hyoscyamine/phenyltoloxamine, Hyoscyamine, Hyoscyamine/methenam/m-blue/phenyl salicyl, Hyoscyamine-phenobarbital)
- BGP HEDIS SKL MUSCLE RELAX MED (Includes combination drugs)
   (ASA/caffeine/orphenadrine, ASA/carisoprodol/codeine, Aspirin-carisoprodol,
   Aspirin-meprobamate, Aspirin-methocarbamol, Carisoprodol, Chlorzoxazone,
   Cyclobenzaprine, Metaxalone, Methocarbamol, Orphenadrine)
- BGP HEDIS ORAL ESTROGEN MEDS (Includes combination drugs) (Conjugated estrogen, Conjugated estrogen-medroxyprogesterone, Esterified estrogen, Esterified estrogen-methyltestosterone, Estropipate)
- BGP HEDIS ORAL HYPOGLYCEMIC RX (Chlorpropamide)
- BGP HEDIS NARCOTIC MEDS (Includes combination drugs)
   (ASA/caffeine/propoxyphene, Acetaminophen-pentazocine, Acetaminophen-propoxyphene, Belladonna-opium, Meperidine, Meperidine-promethazine, Naloxone-pentazocine, Pentazocine, Propoxyphene hydrochloride, Propoxyphene napsylate)
- BGP HEDIS VASODILATOR MEDS (Cyclandelate, Dipyridamole-short acting only, Ergot mesyloid, Isoxsuprine)
- BGP HEDIS OTHER MEDS AVOID ELD (Includes androgens and anabolic steroids, thyroid drugs, and urinary anti-infectives) (Methyltestosterone, Nitrofurantoin, Nitrofurantoin macrocrystals, Nitrofurantoin macrocrystalsmonohydrate, Thyroid desiccated).

Note: For each medication, the days supply must be >0. If the medication was started and then discontinued, CRS will recalculate the # Days Prescribed by subtracting the prescription date (i.e., visit date) from the V Medication Discontinued Date. Example: Rx Date=11/15/2011, Discontinued Date=11/19/2011, Recalculated # Days Prescribed=4. Medications must not have a comment of RETURNED TO STOCK.

# **Key Logic Changes from CRS Version 11.1**

# **Patient List Description**

List of patients 65 and older with at least one prescription for a potentially harmful drug.

### **Measure Source**

**HEDIS** 

# **Measure Past Performance and Long-Term Targets**

None

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Use of High-Risk Medications in the Elderly										
	REPORT % PREV YR % CHG from BASE % CHG from PERIOD PERIOD PREV YR % PERIOD BASE %									
Active Clinical Pts =>65	116		64			65				
<pre># w/exposure to at   least 1 high-risk med   (GPRA Dev.) # w/exposure to   multiple high-risk</pre>	23	19.8	12	18.8	+1.1	18	27.7	-7.9		
meds (GPRA Dev.)	11	9.5	1	1.6	+7.9	8	12.3	-2.8		
Male Active Clinical =>65	53		28			27				
<pre># w/exposure to at   least 1 high-risk med   (GPRA Dev.) # w/exposure to   multiple high-risk</pre>	10	18.9	4	14.3	+4.6	6	22.2	-3.4		
meds (GPRA Dev.)	4	7.5	0	0.0	+7.5	2	7.4	+0.1		
Female Active Clinical =>65	63		36			38				
<pre># w/exposure to at   least 1 high-risk   med (GPRA Dev.) # w/exposure to   multiple high-risk</pre>		20.6			-1.6					
meds (GPRA Dev.) User Pop Pts	7	11.1	1	2.8	+8.3	6	15.8	-4.7		

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=>65	222		149			143			
<pre># w/exposure to at   least 1 high-risk   med # w/exposure to</pre>	25	11.3	13	8.7	+2.5	18	12.6	-1.3	
multiple high-risk meds	11	5.0	2	1.3	+3.6	8	5.6	-0.6	

Figure 2-140: Sample Report, Drugs to be Avoided in the Elderly

Previous Year	November 25, Measures with DEMO INDIAN F iod: Jan 01, 20 Period: Jan 01, riod: Jan 01,	Community S HOSPITAL D12 to Dec 3 L, 2011 to D	31, 2012 Dec 31, 2011	Page 326 rt ***							
Use of High-Risk Medications in the Elderly (con't)											
ACTIVE CLINICAL PATIENTS 65+											
		Distributi									
	65-74	75-84	85+								
CURRENT REPORT PERIOD											
AC Patients 65+	70	36	10								
<pre># w/exposure to at least   1 high-risk med % w/exposure to at least</pre>	15	6	2								
1 high-risk med	21.4	16.7	20.0								
<pre># w/exposure to multiple   high-risk meds % w/exposure to multiple</pre>	8	2	1								
high-risk meds	11.4	5.6	10.0								
PREVIOUS YEAR PERIOD AC Patients 65+	39	19	6								
# w/exposure to at least 1 high risk med	7	4	1								
% w/exposure to at least 1 high-risk med	17.9	21.1	16.7								
<pre># w/exposure to multiple   high-risk meds % w/exposure to multiple</pre>	1	0	0								
high-risk meds	2.6	0.0	0.0								
CHANGE FROM PREV YR % w/exposure to at least											
1 high risk med w/exposure to multiple	+3.5	-4.4	+3.3								
high-risk meds	+8.9	+5.6	+10.0								

Figure 2-141: Sample Report, Drugs to be Avoided in the Elderly

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Use of High Risk Medications in the Elderly: List of patients 65 and older
with at least one high-risk medication.
               HRN COMMUNITY SEX AGE
PATIENT NAME DENOMINATOR
                    NUMERATOR
_____
PATIENT1, JONELLE 000001 COMMUNITY #1 F 69
UP,AC 2 drugs: 08/04/12 ESTERIFIED ESTROGENS 0.625MG TAB (ORAL ESTROGEN);
08/04/12 PROPOXYPHENE-N 100MG/APAP 650MG TAB (NARCOTIC)
PATIENT2, PAULINE 000002 COMMUNITY #1 F 70
UP,AC 1 drug:11/02/12 PROPOXYPHENE-N 100MG/APAP 650MG TAB (NARCOTIC)
PATIENT3, NADINE 000003 COMMUNITY #1 F 82
UP,AC 2 drugs: 09/25/12 DIAZEPAM 5MG TAB (BENZODIAZEPINE); 09/25/12
PROPOXYPHENE-N 100MG/APAP 650MG TAB (NARCOTIC)
PATIENT4, JESSE NATHAN 000004 COMMUNITY #1 M 77
UP,AC 1 drug:08/27/122 CYCLOBENZAPRINE HCL 10MG TAB (SKL MUSCLE)
```

Figure 2-142: Sample Patient List, Drugs to be Avoided in the Elderly

### 2.10.19 Functional Status Assessment in Elders

### **Denominator**

Active Clinical patients ages 55 and older. Broken down by gender.

#### **Numerator**

Patients screened for functional status at any time during the Report Period.

### **Logic Description**

Age is calculated at the beginning of the Report Period.

Functional status screening definition: Any non-null values in V Elder Care for (1) at least one of the following ADL fields: toileting, bathing, dressing, transfers, feeding, or continence and (2) at least one of the following IADL fields: finances, cooking, shopping, housework/chores, medications or transportation during the Report Period.

### **Key Logic Changes from CRS Version 11.1**

# **Patient List Description**

List of patients =>55 with functional status codes, if any. The following are the abbreviations used in the Numerator column:

- TLT–Toileting
- BATH–Bathing
- DRES-Dressing
- XFER-Transfers
- FEED-Feeding
- CONT-Continence
- FIN-Finances
- COOK-Cooking
- SHOP-Shopping
- HSWK-Housework/Chores
- MEDS–Medications
- TRNS-Transportation

### **Measure Source**

None

### Measure Past Performance and Long-Term Targets

Previous	ort Perio	Measun DEMO od: Ja eriod	INDIAN HO an 01, 201 : Jan 01,	ommur SPITA 2 to 2011		)12 l, 2011	_	e 328	
Functional Status Asse	essment	in Elo	ders						
	REPORT		PREV YR PERIOD		CHG from PREV YR %			HG from ASE %	
Active Clinical Pts =>55	271		162			127			
# w/functional status screening	2	0.7	0	0.0	+0.7	0	0.0	+0.7	
Male Active Clinical =>55	135		77			60			

# w/functional status screening	1	0.7	0	0.0	+0.7	0	0.0	+0.7
Female Active Clinical =>55	136		85			67		
# w/functional status screening	1	0.7	0	0.0	+0.7	0	0.0	+0.7

Figure 2-143: Sample Report, Functional Status Assessment in Elders

```
UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic
PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease;
CHD-Active Coronary Heart Disease; HR=High Risk Patient
Functional Status Assessment in Elders: List of patients => 55 with functional
status codes, if any.
                 HRN COMMUNITY SEX AGE
PATIENT NAME
DENOMINATOR
                            NUMERATOR
PATIENT1, GLENDA JOYCE 000001 COMMUNITY #1 F 57
PATIENT2, NADINE 000002 COMMUNITY #1 F 61
PATIENT3, CHARLOTTE MAE 000003 COMMUNITY #1 F 64
                       YES: 02/24/12 BATH, CONT, COOK, DRES, FEED, FIN, HSWK,
MEDS, SHOP, TLT, TRNS, XFER
PATIENT4, KATHERINE ANN 000004 COMMUNITY #1 F 66
AC YES: 07/11/12 BATH, F PATIENT5,ANNA MARIE 000005 COMMUNITY #1 F 66
                           YES: 07/11/12 BATH, FIN
AC
PATIENT6, DIANA 000006 COMMUNITY #1 F 67
PATIENT7, PEGGY ANN 000007 COMMUNITY #1 F 70
                           NO: 05/20/12 FIN
```

Figure 2-144: Sample Patient List, Functional Status Assessment in Elders

### 2.10.20 Fall Risk Assessment in Elders

#### **Denominators**

Active Clinical patients ages 65 and older. Broken down by gender.

User Population patients ages 65 and older. Broken down by gender.

#### **Numerators**

Patients who have been screened for fall risk or with a fall-related diagnosis in the past year.

**Note**: This numerator does *not* include refusals.

- a. Patients who have been screened for fall risk in the past year
- b. Patients with a documented history of falling in the past year
- c. Patients with a fall-related injury diagnosis in the past year
- d. Patients with abnormality of gait/balance or mobility diagnosis in the past year

Patients with a documented refusal of fall risk screening exam in the past year

### **Logic Description**

Age of the patient is calculated at the beginning of the Report Period.

Fall risk screening/fall related diagnosis is defined as any of the codes in the table below.

	ICD and Other Codes	Exam Code	E Codes (Injury)
Fall Risk Exam	<b>CPT</b> : 1100F, 1101F, 3288F	V Exam: 37 (Fall Risk)	
History of Falling	V POV: V15.88 (Personal History of Fall)		
Fall-related Injury			V POV (Cause Codes #1-3): E880.*, E881.*, E883.*, E884.*, E885.*, E886.*, E888.*
Abnormality of Gait/Balance or Mobility	V POV: 781.2, 781.3, 719.7, 719.70 (old code), 719.75–719.77 (old codes), 438.84, 333.99, 443.9		
Refusal		V Exam: 37 (Fall Risk)	

### **Key Logic Changes from CRS Version 11.1**

None

### **Patient List Description**

List of patients 65 years or older with fall risk assessment, if any.

#### **Measure Source**

HP 2010 15–28 Reduce hip fractures among older adults.

### Measure Past Performance and Long-Term Targets

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Fall Risk Assessment in Elders (con't)											
Active Clinical Pts 65+	116		64			65					
# w/ fall risk screen/											
Dx-No Refusals A. # w/ fall risk	14	12.1	8	12.5	-0.4	8	12.3	-0.2			
screen B. # w/ history	5	4.3	0	0.0	+4.3	0	0.0	+4.3			
of fall	1	0.9	0	0.0	+0.9	0	0.0	+0.9			
C. # w/ fall injury D. # w/ abnormal	2	1.7	1	1.6	+0.2	3	4.6	-2.9			
gait # w/ refusal		5.2 2.6		10.9	-5.8 +2.6	5 0	7.7 0.0	-2.5 +2.6			
Male Active Clinical											
65+	53		28			27					
# w/ fall risk screen/											
Dx-No Refusals A. # w/ fall risk	6	11.3	3	10.7	+0.6	2	7.4	+3.9			
screen B. # w/ history	2	3.8	0	0.0	+3.8	0	0.0	+3.8			
of fall	1	1.9	0	0.0	+1.9	0	0.0	+1.9			
C. # w/ fall injury D. # w/ abnormal	0	0.0	0	0.0	+0.0	1	3.7	-3.7			
gait # w/ refusal	3 1	5.7 1.9		10.7	-5.1 +1.9	1 0	3.7 0.0	+2.0 +1.9			
Female Active Clinical											
65+	63		36			38					
# w/ fall risk screen/ Dx-No Refusals A. # w/ fall risk	8	12.7	5	13.9	-1.2	6	15.8	-3.1			
screen	3	4.8	0	0.0	+4.8	0	0.0	+4.8			
B. # w/ history of fall	0	0.0	0	0.0	+0.0	0	0.0	+0.0			
C. # w/ fall injury D. # w/ abnormal		3.2	1	2.8	+0.4	2		-2.1			
gait # w/ refusal	3 2	4.8	4		-6.3 +3.2	4 0	10.5	-5.8 +3.2			

Figure 2-145: Sample Report, Fall Risk Assessment in Elders

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient

Fall Risk Assessment in Elders: List of patients 65 years or older with fall risk assessment, if any.

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PATIENT NAME DENOMINATOR	HRN COMMUNITY SEX AGE NUMERATOR
PATIENT1, SHERRY UP, AC	000001 COMMUNITY #1 F 68
PATIENT2, LORETTA LYNN	000002 COMMUNITY #1 F 78
UP,AC	Refused 03/03/12 Ex 37
PATIENT17, NICOLE	000017 COMMUNITY #2 F 71
UP,AC	Abnormal Gait: 11/25/12 POV 443.9
PATIENT18, VERONICA	000018 COMMUNITY #2 F 72
UP	Screen: 03/03/12 CPT 1100F
PATIENT19,STEPHANIE	000019 COMMUNITY #2 F 76
UP,AC	Fall Injury: 11/10/12 E-CODE E883.9

Figure 2-146: Sample Patient List, Fall Risk Assessment in Elders

### 2.10.21 Palliative Care

### **Denominators**

Active Clinical patients ages 18 and older with two or more types of cancer documented during the Report Period. Broken down by gender and age groups (18-54, >55).

### **Numerators**

No denominator; count only. The total number of *Active Clinical patients* with at least one palliative care visit during the Report Period. Broken down by age groups (<18, 18–54, >55).

No denominator; count only. The total number of palliative care visits for *Active Clinical patients* during the Report Period. Broken down by age groups (<18, 18-54, >55).

Patients with at least two palliative care visits during the Report Period.

# **Logic Description**

Age is calculated at the beginning of the Report Period.

Palliative care visit definition: POV V66.7.

Cancer Types:

	ICD Codes
Melanoma	<b>V POV</b> : 172*
Breast	<b>V POV</b> : 174*, 175*, 239.3
Colon	<b>V POV</b> : 153*, 154*, 235.2

	ICD Codes
Gyn	<b>V POV</b> : 180*, 182*, 183*, 184*, 236.1, 236.2
Prostate	<b>V POV</b> : 185* 236.5
Testes/Male GU	<b>V POV</b> : 186*, 187.3, 187.4, 187.9, 236.4, 236.6
Head and Neck	<b>V POV</b> : 140–149.9, 160*, 161*, 162*, 195.0
Urinary Tract	<b>V POV</b> : 188*, 189*, 236.7, 236.91, 239.4, 239.5
Non-melanomatous skin cancer	<b>V POV</b> : 173*, 238.2
Non-colon GI	<b>V POV</b> : 150–152.9, 155–159.9, 235*, 239.0
Lung	<b>V POV</b> : 162*, 235.9, 239.1
Brain	<b>V POV</b> : 190–192.9, 237.5, 237.6, 239.6
Bones/soft tissue	<b>V POV</b> : 170*, 171*, 238.1, 238.2
Endocrine	<b>V POV</b> : 193, 194*, 237.0, 237.4, 239.7
Pleura/mediastinum	<b>V POV</b> : 163*, 164*
Non-specific site	<b>V POV</b> : 195*, 199*, 238.8, 238.9, 239.8, 239.9
Lymph node spread	<b>V POV</b> : 196*
Secondary cancer	<b>V POV</b> : 196*, 197*

### **Patient List Description**

List of patients with a palliative care visit.

# **Key Logic Changes from CRS Version 11.1**

None

### **Measure Source**

None

# **Measure Past Performance and Long-Term Targets**

```
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*** IHS 2012 Selected Measures with Community Specified Report ***

DEMO INDIAN HOSPITAL

Report Period: Jan 01, 2012 to Dec 31, 2012

Previous Year Period: Jan 01, 2011 to Dec 31, 2011

Baseline Period: Jan 01, 2000 to Dec 31, 2000

Palliative Care

REPORT PREV YR CHG from BASE CHG from PERIOD PREV YR PERIOD BASE

Total # of Patients

w/At Least 1 Palliative
```

Care Visit	40	0	+40	0	+40
A. Total # of Patients <18 w/At Least 1 Pallia Care Visit	tive 1	0	+1	0	+1
B. Total # of Patients 1 w/At Least 1 Palliative Care Visit		0	+25	0	+25
C. Total # of Patients 5 w/At Least 1 Palliative Care Visit		0	+14	0	+14
Total # of Palliative Care Visits	62	0	+62	0	+62
A. Total # of Palliative Care Visits-Pts <18	2	0	+2	0	+2
B. Total # of Palliative Care Visits-Pts 18-54	38	0	+38	0	+38
C. Total # of Palliative Care Visits-Pts 55+	22	0	+22	0	+22
Active Clinical Pts 18+ w/ 2+ types of cancer	28	3		3	
# w/ 2+ Palliative Care Visits	14 50.0	0 0.0	+50.0	0 0.0	+50.0

Figure 2-147: Sample Report, Palliative Care

UP-User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Palliative Care: List of patients with a palliative care visit, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR PATIENT1, JOHN 000012 Community #1 M 57 AC 1 visit: 05/01/12 PATIENT2, ROBERT 000013 Community #1 M 59 2 visits: 01/25/12, 05/10/12 AC, CAN AC, CAN 2 visits: 01/25/12, 0 PATIENT3, JAMES 000014 Community #2 M 67 AC, CAN 0 visits: 000015 Community #3 F 78 PATIENT4, TONYA 1 visit: 06/01/12 PATIENT5, RITA ANN 000016 Community #3 F 96 2 visits: 06/01/12; 06/07/12 PATIENT6, Clifford 000017 Community #3 M 24

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AC,CAN 3 visits: 01/25/12, 05/10/12, 08/01/12

Figure 2-148: Sample Patient List, Palliative Care

### 2.10.22 Annual Wellness Visit

### **Denominators**

Active Clinical patients ages 65 and older. Broken down by gender.

### **Numerators**

Patients with at least one Annual Wellness Exam in the past 15 months.

### **Logic Description**

Age is calculated at the beginning of the Report Period.

Annual Wellness Exam: CPT G0438, G0439, G0402.

### **Patient List Description**

List of patients with an annual wellness visit in the past 15 months.

### **Key Logic Changes from CRS Version 11.1**

None

### **Measure Source**

None

# **Measure Past Performance and Long-Term Targets**

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Annual Wellness Visit	REPORT PERIOD		CV YR		CHG from PREV YR			CHG from BASE		
Active Clinical Pts 65+	116		64			65				
# w/ Annual Wellness Exam	2	1.7	0	0.0	+1.7	0	0.0	+1.7		

Male Active Clinical Pts 65+	53		28			27			
# w/ Annual Wellness Exam	0	0.0	0	0.0	+0.0	0	0.0	+0.0	
Female Active Clinical Pts 65+	63		36			38			
# w/ Annual Wellness Exam	2	3.2	0	0.0	+3.2	0	0.0	+3.2	

Figure 2-149: Sample Report, Annual Wellness Visit

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Annual Wellness Visit: List of patients with an annual wellness visit in the past 15 months, if any. PATIENT NAME HRN COMMUNITY SEX AGE DENOMINATOR NUMERATOR Patient1, DENISE 000001 Community #1 F 65 12/31/11 G0402 Patient2, MELISSA GAYLE 000002 Community #1 F 66 Patient3, JESSICA DAWN 000003 Community #1 F 67 AC 02/22/12 G0438 Patient4, RUTH ALICE 000004 Community #1 F 69 Patient5, BRYSON DEWAY 000005 Community #1 F 72 Patient6, BRITTNEY ANN 000006 Community #1 F 73 05/31/12 G0439 Patient7,MARK 000007 Community #1 M 67 Patient8, HOWIE 000008 Community #1 M 72 10/01/11 G0402

Figure 2-150: Sample Patient List, Annual Wellness Visit

# 2.10.23 Goal Setting

#### **Denominators**

User Population patients who received patient education during the report period.

### **Numerators**

Number of patients who set at least one goal during the Report Period.

Number of patients who met at least one goal during the Report Period.

### **Logic Description**

Patient education codes must be the standard national patient education codes, which are included in the Patient and Family Education Protocols and Codes (PEPC) manual published each year. If codes are found that are not in the table, they will not be reported on (i.e. locally-developed codes).

### Numerator Logic:

For Goal Set, the patient education code must have a "GS" value documented during the Report Period.

For Goal Met, the patient education code must have a "GM" value documented during the Report Period but the patient is not required to have set a goal during the Report Period.

# **Patient List Description**

List of User Population patients who received patient education during the Report Period with goal setting information.

# **Key Logic Changes from CRS Version 11.1**

None

### **Measure Source**

None

# **Measure Past Performance and Long-Term Targets**

DU November 25, 2012 Page 338  *** IHS 2012 Selected Measures with Community Specified Report ***  DEMO INDIAN HOSPITAL  Report Period: Jan 01, 2012 to Dec 31, 2012  Previous Year Period: Jan 01, 2011 to Dec 31, 2011  Baseline Period: Jan 01, 2000 to Dec 31, 2000									
Goal Setting									
	REPORT PERIOD		V YR IOD		CHG from PREV YR		_	HG from	
# User Pop w/ Pat Ed	1,024		858			644			
# w/ goal set # w/ goal met		0.8		0.0		0	0.0		

Figure 2-151: Sample Report, Goal Setting

UP=User Pop; AC=Active Clinical; AD=Active Diabetic; AAD=Active Adult Diabetic PREG=Pregnant Female; IMM=Active IMM Pkg Pt; IHD=Active Ischemic Heart Disease; CHD-Active Coronary Heart Disease; HR=High Risk Patient Goal Setting: List of User Population patients who received patient education during the Report Period with goal setting information. PATIENT NAME HRN COMMUNITY SEX AGE
DENOMINATOR PATIENT NAME \_\_\_\_\_ Patient1, Paula 000001 Community #1 F 34 GS: 11/17/12 Patient2, PENNY 000002 Community #1 F 43 Patient3, RITA 000003 Community #1 F 64 GS: 04/15/12, GM: 06/15/12 Patient4, HARRY 000004 Community #1 M 50
UP GM: 10/30/12 Patient5, ROSS 000005 Community #1 M 55 Patient6, FELIPE 000006 Community #1 M 57 UP GS: 09/10/12 Patient7, MARK 000007 Community #1 M 67 Patient7,MARK Patient8, CATHERINE 000008 Community #2 F 72 GM: 07/30/12

Figure 2-152: Sample Patient List, Goal Setting

# **Contact Information**

If you have any questions or comments regarding this distribution, please contact the OIT Help Desk (IHS).

**Phone:** (505) 248-4371 or (888) 830-7280 (toll free)

**Fax:** (505) 248-4363

Web: <a href="http://www.ihs.gov/GeneralWeb/HelpCenter/Helpdesk/index.cfm">http://www.ihs.gov/GeneralWeb/HelpCenter/Helpdesk/index.cfm</a>

Email: support@ihs.gov