

### 1. General Information

On the 4500 Names Modical Record Number (MPN)	Technical S	pecifications
Seq. #: 1500 Name: Medical Record Number (MRN)	ShortName:	Patient_MRN
Coding Instructions: Indicate the patient's medical record number as assigned by the medical practice.	Parent Seq #:	
Target Value: The value on current encounter	Parent Name:	
	Parent Value:	
Selections: (none)	Missing Data:	Illegal
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (20)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 1510 Name: Encounter Date		Specifications
·	ShortName:	EncounterDate
Coding Instructions: Indicate the date of the patient encounter or visit to the physician office.	Parent Seq #: Parent Name:	
Target Value: The value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Illegal
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 1530 Name: Location ID	Technical S	pecifications
Seq. #: 1930 Name. Location in	ShortName:	LocationID
<b>Coding Instructions:</b> Indicate the Location Identification number assigned for the office location by the ACC-NCDR.	Parent Seq #: Parent Name:	
Target Value: The value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Illegal
Colonial (none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions: (none)	Format:	Integer (3)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	Automatic



### 1. General Information

Seq. #: 1540 Name: Provider Last Name	Technical S	Specifications
Seq. #: 1540 Name. 1 Tovider Last Name	ShortName:	Physician_LastNam
Coding Instructions: This element has been retired effective PINNACLE v1.3.	5	е
Target Value: N/A	Parent Seq #: Parent Name:	
Selections: (none)	Parent Value:	
Supporting Definitions (2000)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (50)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 1541 Name: Provider First Name	Technical S	specifications
·	ShortName:	Physician_FirstNam e
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Donout Com #	е
Target Value: N/A	Parent Seq #: Parent Name:	
Selections: (none)	Parent Value:	
Selections. (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (50)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
The state of the s	Technical S	Specifications
Seq. #: 1542 Name: Provider Middle Name	ShortName:	Physician_MidName
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Parent Seq #:	
Torque Value: N/A	Parent Name:	
Target Value: N/A	Parent Value:	
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (50)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User





#### 1. General Information

Con. #. 4550 Nome. Provider NDI	Technical S	pecifications
Seq. #: 1550 Name: Provider NPI	ShortName:	Physician_NPI
Coding Instructions: Indicate the evaluating provider's National Provider Identifier (NPI).	Parent Seq #:	
Target Value, The value on current encounter	Parent Name:	
Target Value: The value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Illegal
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (10)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 1555 Name: Encounter TIN	Technical S	pecifications
Seq. #: 1999 Name. Encounter Till	ShortName:	EncounterTIN
Coding Instructions: Indicate the practice Tax Identification Number (TIN) to which the Encounter shou billed. If the practice has changed TINs or the provider bills to multiple TINs, be or		
that the TIN recorded for the encounter reflects the appropriate billing TIN at the ti		
the encounter.	Parent Value:	
Target Value: The value on current encounter	Missing Data:	No Action
Selections: (none)	Harvested:	Yes (DCR,PINN)
delections. (none)	Format:	Integer (9)
Supporting Definitions: (none)	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 1560 Name: Patient New to the Practice	Technical S	pecifications
·	ShortName:	PatNew
<b>Coding Instructions:</b> Indicate if this encounter is the first time the patient was treated by the practice.	Parent Seq #:	
Note(s):	Parent Name:	
If the patient was treated at the same practice but a different location, then code '	No'. Parent Value:	
Target Value: The value on current encounter	Missing Data:	Report
Selections: Code Selection Text Definition	Harvested:	Yes (DCR,PINN)
Gelections. Code Selection rext Delimition	Format:	Text (Categorical)
0 No	Default Value:	NULL
1 Yes	Usual Range:	
Supporting Definitions: (none)	Valid Range:	
ouppoining Deminions. (110110)	DataSource:	User





#### 1. General Information

Seq. #: 1565 Name: Primary Reason for Encounter

Coding Instructions: This element has been retired effective v1.4

Target Value: The value on current encounter

_				
Selec	tions:	Code	Selection Text	Definition
	·	1	Atrial Fibrillation related	
		2	Coronary Artery Disease related	
		3	Diabetes related	
		4	Heart Failure related	
		5	Hypertension related	
		6	Other Cardiac related reason	
		7	Non-Cardiac related reason	

Supporting Definitions: (none)



### A. Patient Demographics

Con. #. 2000 Name: Detient Last Name	Technical Specifications
Seq. #: 2000 Name: Patient Last Name	ShortName: LastName
Coding Instructions: Indicate the patient's last name. Hyphenated names should be recorded with a hyphenated names should be recorded with a hyphenated names.	Parent Seq #:
Torget Value. The value on current engagners	Parent Name:
Target Value: The value on current encounter	Parent Value:
Selections: (none)	Missing Data: Report
Supporting Definitions: (none)	Harvested: Yes (DCR,PINN)
	Format: Text (50)
	Default Value: NULL
	Usual Range:
	Valid Range:
	DataSource: User
Seq. #: 2010 Name: Patient First Name	Technical Specifications
Seq. #. 2010 Name. I addit i ist Name	ShortName: FirstName
Coding Instructions: Indicate the patient's first name.	Parent Seq #:
Target Value: The value on current encounter	Parent Name:
	Parent Value:
Selections: (none)	Missing Data: Report
Supporting Definitions: (none)	Harvested: Yes (DCR,PINN)
	Format: Text (50)
	Default Value: NULL
	Usual Range:
	Valid Range:
	DataSource: User
Seq. #: 2020 Name: Patient Middle Name	Technical Specifications
·	ShortName: MidName
Coding Instructions: Indicate the patient's middle name(s).	Parent Seq #:
Note(s):	Parent Name:
If the patient has multiple middle names, enter each middle name separated by a	single Parent Value:
space.	Missing Data: Report
Target Value: The value on current encounter	Harvested: Yes (DCR,PINN)
Target Value: The value on current encounter	Format: Text (50)
Selections: (none)	Default Value: NULL
Supporting Definitions: (none)	Usual Range:
	Valid Range:
	DataSource: User



#### A. Patient Demographics

**Seq. #:** 2030 **Name:** SSN

Coding Instructions: Indicate the patient's United States Social Security Number (SSN).

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: SSN

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (9)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User

Seq. #: 2040 Name: Patient ID

Coding Instructions: Indicate the number created and automatically inserted by the software that uniquely

identifies this patient.

Note(s):

Once assigned to a patient at the participating facility, this number should never be

changed or reassigned to a different patient.

If the patient returns to the same medical practice or for follow-up, they must receive

this same unique patient identifier.

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: PatientID

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Integer (9)

Default Value: NULL Usual Range:

Valid Range: 1-999999999

DataSource: Automatic

Seq. #: 2050 Name: Date of Birth

Coding Instructions: Indicate the patient's date of birth.

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: Do

Parent Seq #:

Parent Name:

Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

User

Default Value: NULL

Usual Range:

Valid Range: DataSource:





### A. Patient Demographics

<b>Seq. #</b> : 2060 <b>Name</b> : Sex				Technical S	pecifications
Seq. #. 2000 Name: OCX				ShortName:	Sex
Coding Instructions:	Indicate the p	patient's sex at birth.		Parent Seq #:	
Target Value:	The value or	n current encounter		Parent Name:	
				Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
	1	Male		Harvested:	Yes (DCR,PINN)
	2	Female		Format:	Text (Categorical)
		· Gillaid		Default Value:	NULL
Supporting Definitions:	(none)			Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 2065 Name:	Patient De	acasad		Technical S	pecifications
Seq. #: 2005 Name.	i allent De	ceaseu		ShortName:	Death_Ind
Coding Instructions:	Indicate if the	e patient died, regardless o	f etiology.	Parent Seq #:	
Target Value	The value or	n current encounter		Parent Name:	
_				Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
	0	No		Harvested:	Yes (DCR,PINN)
	1	Yes		Format:	Text (Categorical)
				Default Value:	No
Supporting Definitions:	(none)			Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 2067 Name:	Death Dat	-Δ		Technical S	pecifications
Seq. #: 2007 Name.	Death Dat	.0		ShortName:	Death_Date
Coding Instructions:	Indicate the p	patient's date of death.		Parent Seq #:	2065
Target Value:	The last valu	ie on current encounter		Parent Name:	Patient Deceased
_		le on carrent encounter		Parent Value:	Yes
Selections:	(none)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (DCR,PINN)
<del>-</del>				Format:	Date (mm/dd/yyyy)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	
				DataSource:	User





#### A. Patient Demographics

Seq. #: 2068 Name: Primary Cause of Death

Coding Instructions: Indicate the patient's PRIMARY cause of death, i.e. the first significant abnormal event

which ultimately led to death.

Target Value: The last value on current encounter

Selections: Code Definition Selection Text Cardiac 1 Neurologic 2 Renal 3 Vascular 4 Infection 5 Valvular 6 Pulmonary 7 Unknown 8

**Technical Specifications** DeathCause ShortName: Parent Seq #: 2065 Parent Name: Patient Deceased Parent Value: Yes Missing Data: Report Yes (DCR) Harvested: Format: Text (Categorical) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User

**Technical Specifications** 

ShortName:

Parent Seq #: Parent Name:

Parent Value:

**Missing Data:** 

**Default Value:** 

**Usual Range:** 

Valid Range: DataSource:

Harvested:

Format:

RaceWhite

Report

User

Yes (DCR,PINN)

Text (Categorical)

Supporting Definitions: (none)

Seq. #: 2070 Name: Race - White

Coding Instructions: Indicate if the patient is White as determined by the patient/family.

Other

Note(s):

9

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: White (Race):

Having origins in any of the original peoples of Europe, the Middle East, or North Africa.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

Effective for Patient Discharges April 17, 2015





#### A. Patient Demographics

Seq. #: 2071 Name: Race - Black/African American

Coding Instructions: Indicate if the patient is Black or African American as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No1 Yes

Supporting Definitions: Black/African American (Race):

Having origins in any of the black racial groups of Africa. Terms such as "Haitian" or

"Negro" can be used in addition to "Black or African American."

Source: U.S. Office of Management and Budget. Classification of Federal Data

on Race and Ethnicity

**Technical Specifications** 

**Technical Specifications** 

ShortName:

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range: DataSource:

Harvested: Format: RaceBlack

Report

User

Yes (DCR,PINN)

Text (Categorical)

ShortName: RaceAsian

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No

Usual Range: Valid Range:

DataSource: User

Seq. #: 2072 Name: Race - Asian

Coding Instructions: Indicate if the patient is Asian as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Asian (Race):

Having origins in any of the original peoples of the Far East, Southeast Asia, or the Indian subcontinent including, for example, Cambodia, China, India, Japan, Korea,

Malaysia, Pakistan, the Philippine Islands, Thailand, and Vietnam.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity





#### A. Patient Demographics

Name: Race - American Indian/Alaskan Native **Seq. #:** 2073

Coding Instructions: Indicate if the patient is American Indian or Alaskan Native as determined by the

patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: American Indian or Alaskan Native (Race):

Having origins in any of the original peoples of North and South America (including Central America), and who maintains tribal affiliation or community attachment.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

**Technical Specifications** 

RaceAmIndian

Yes (DCR,PINN)

Text (Categorical)

Report

User

ShortName:

Parent Seq #:

**Parent Name:** 

Parent Value:

Missing Data:

**Default Value: Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

**Seq. #**: 2074 Name: Race - Native Hawaiian/Pacific Islander Coding Instructions: Indicate if the patient is Native Hawaiian or Pacific Islander as determined by the

patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selection Text Selections: Code Definition

> No 0 Yes 1

Supporting Definitions: Native Hawaiian or Pacific Islander (Race):

Having origins in any of the original peoples of Hawaii, Guam, Samoa, or other Pacific

Islands.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

**Technical Specifications** 

ShortName: RaceNatHaw

Parent Seq #: **Parent Name: Parent Value:** 

Missing Data: Report

> Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

**Default Value: Usual Range:** Valid Range:

DataSource: User



#### A. Patient Demographics

Seq. #: 2076 Name: Hispanic or Latino Ethnicity

Coding Instructions: Indicate if the patient is of Hispanic or Latino ethnicity as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No1 Yes

Supporting Definitions: Hispanic or Latino Ethnicity:

A person of Cuban, Mexican, Puerto Rican, Cuban, South or Central American, or other Spanish culture or origin, regardless of race. The term, "Spanish origin," can be used in

addition to "Hispanic or Latino."

Source: U.S. Office of Management and Budget. Classification of Federal Data

on Race and Ethnicity

Seq. #: 2080 Name: Race - Asian Indian

Coding Instructions: Indicate if the patient is Asian Indian as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Race - Asian Indian:

Having origins in any of the original peoples of India.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity.

lian: DataSource:

Technical Specifications

ShortName: HispOrig

Parent Seq #: Parent Name:

Harvested:

Parent Value:

Missing Data: Report

Format: Text (Categorical)

Yes (DCR,PINN)

Default Value: No

Usual Range:

Valid Range:

DataSource: User

ShortName:

Parent Seq #:

**Parent Name:** 

**Parent Value:** 

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

Harvested:

Format:

**Technical Specifications** 

2072

Yes

Report

User

RaceAsianIndian

Yes (DCR,PINN)

Text (Categorical)

Race - Asian



#### A. Patient Demographics

Seq. #: 2081 Name: Race - Chinese

Coding Instructions: Indicate if the patient is Chinese as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Race - Chinese:

Having origins in any of the original peoples of China.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Seq. #: 2082 Name: Race - Filipino

Coding Instructions: Indicate if the patient is Filipino as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Asian - Filipino:

Having origins in any of the original peoples of the Philippines.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

Missing Data: Report
Harvested: Yes (DCR,PINN)
Format: Text (Categorical)

**Technical Specifications** 

RaceFilipino

Race - Asian

Yes (DCR,PINN)

Text (Categorical)

2072

Yes

Report

User

**Technical Specifications** 

2072

Yes

RaceChinese

Race - Asian

Default Value: No
Usual Range:
Valid Range:

ShortName:

Parent Seq #:

Parent Name:

Parent Value:

DataSource: User

ShortName:

Parent Seq #:

**Parent Name:** 

**Parent Value:** 

Missing Data:

Default Value: Usual Range:

Valid Range:

DataSource:

Harvested:

Format:

Effective for Patient Discharges April 17, 2015



#### A. Patient Demographics

Seq. #: 2083 Name: Race - Japanese

Coding Instructions: Indicate if the patient is Japanese as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Asian - Japanese:

Having origins in any of the original peoples of Japan.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Seq. #: 2084 Name: Race - Korean

Coding Instructions: Indicate if the patient is Koreans as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No1 Yes

Supporting Definitions: Asian - Korean:

Having origins in any of the original peoples of Korea.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

ShortName: RaceJapanese

Parent Seq #: 2072

Parent Name: Race - Asian

Parent Value: Yes

**Technical Specifications** 

Missing Data: Report
Harvested: Yes (DCR,PINN)

**Technical Specifications** 

RaceKorean

Race - Asian

Yes (DCR,PINN)

Text (Categorical)

2072

Yes

Report

User

Format: Text (Categorical)

Default Value: No
Usual Range:
Valid Range:

DataSource: User

ShortName:

Parent Seq #:

**Parent Name:** 

**Parent Value:** 

Missing Data:

Default Value: Usual Range:

Valid Range:

DataSource:

Harvested:

Format:

Effective for Patient Discharges April 17, 2015



#### A. Patient Demographics

Name: Race - Vietnamese **Seq. #**: 2085

Coding Instructions: Indicate if the patient is Vietnamese as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Asian - Vietnamese:

Having origins in any of the original peoples of Viet Nam.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Name: Race - Other Asian **Seq. #:** 2086

Coding Instructions: Indicate if the patient is of other asian ethnicity as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selection Text Selections: Code Definition

> No 0 Yes 1

Supporting Definitions: Asian - Other Asian:

Having origins in any of the original peoples elsewhere in Asia.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

**Technical Specifications** 

RaceVietnamese ShortName:

Parent Seq #: 2072

Parent Name: Race - Asian

Parent Value: Yes Missing Data:

Yes (DCR,PINN) Harvested:

**Technical Specifications** 

2072

Yes

Report

User

RaceAsianOther

Yes (DCR,PINN)

Text (Categorical)

Race - Asian

Format: Text (Categorical)

Report

**Default Value:** 

**Usual Range:** 

Valid Range:

ShortName:

Parent Seq #:

**Parent Name:** 

**Parent Value:** 

Missing Data:

**Default Value: Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

DataSource: User



#### A. Patient Demographics

Seq. #: 2090 Name: Race - Native Hawaiian

Coding Instructions: Indicate if the patient is Native Hawaiian as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Native Hawaiian/Pacific Islander - Native Hawaiian:

Having origins in any of the original peoples of the islands of Hawaii.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Seq. #: 2091 Name: Race - Guamanian or Chamorro

Coding Instructions: Indicate if the patient is Guamanian or Chamorro as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on arrival at this facility

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Native Hawaiian/Pacific Islander - Guamanian or Chamorro:

Having origins in any of the original peoples of the Mariana Islands or the island of

Guam.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

Technical Specifications

**Technical Specifications** 

2074

Yes

Islander

Report

ShortName:

Parent Seq #:

Parent Name:

Parent Value:

Missing Data:

Default Value:

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

RaceNativeHawaii

Race - Native

Hawaiian/Pacific

Yes (DCR,PINN)

Text (Categorical)

ShortName: RaceGuamChamorr

User

0

Parent Seg #: 2074

Parent Name: Race - Native

Hawaiian/Pacific

Islander

Parent Value: Yes

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User



#### A. Patient Demographics

Seq. #: 2092 Name: Race - Samoan

Coding Instructions: Indicate if the patient is Samoan as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Native Hawaiian/Pacific Islander - Samoan:

Having origins in any of the original peoples of the island of the Somoa.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Technical Specifications

**Technical Specifications** 

2074

Islander

Report

Yes

ShortName:

Parent Seq #:

Parent Name:

**Parent Value:** 

Missing Data:

Default Value:

**Usual Range:** 

Valid Range: DataSource:

Harvested:

Format:

RaceSamoan

Race - Native

Hawaiian/Pacific

Yes (DCR,PINN)

Text (Categorical)

ShortName: RacePacificIslandOt

her

User

Parent Seq #: 2074

Parent Name: Race - Native

Hawaiian/Pacific

Islander

Parent Value: Yes

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value:

Usual Range: Valid Range:

DataSource: User

Seq. #: 2093 Name: Race - Other Pacific Islander

Coding Instructions: Indicate if the patient is of other pacific island ethnicity as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No

Yes

Supporting Definitions: Native Hawaiian/Pacific Islander - Other Pacific Island:

Having origins in any of the original peoples of any other island in the Pacific.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity





#### A. Patient Demographics

Name: Hispanic Ethnicity Type - Mexican/Mexican **Seq. #:** 2100

American/Chicano

Coding Instructions: Indicate if the patient is of Mexican/Mexican American/Chicano ethnicity as determined by

the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Hispanic Ethnicity - Mexican/Mexican American/Chicano:

Having origins in any of the original peoples of Mexico.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Name: Hispanic Ethnicity Type - Puerto Rican **Seq.** #: 2101

Coding Instructions: Indicate if the patient is of Puerto Rican ethnicity as determined by the patient/family.

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Hispanic Ethnicity - Puerto Rican:

Having origins in any of the original peoples of Puerto Rico.

Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity

User

**Technical Specifications** 

2076

Yes

Report

HispEthnicityMexica

Hispanic or Latino Ethnicity

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #:

**Parent Name:** 

Parent Value:

Missing Data:

Default Value:

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

**Technical Specifications** 

ShortName: HispEthnicityPuerto

Rico

Parent Seq #: 2076

**Parent Name:** Hispanic or Latino

Ethnicity

Text (Categorical)

**Parent Value:** Yes

Missing Data: Report

> Harvested: Yes (DCR,PINN)

Format: Default Value:

**Usual Range:** 

Valid Range:

DataSource: User



#### A. Patient Demographics

Seq. #: 2102 Name: Hispanic Ethnicity Type - Cuban

Coding Instructions: Indicate if the patient is of Cuban ethnicity as determined by the patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Hispanic Ethnicity - Cuban:

Having origins in any of the original peoples of Cuba.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Name: Hispanic Ethnicity Type - Other Hispanic/Latino/Spanish **Sea.** #: 2103

Coding Instructions: Indicate if the patient is of other hispanic/latino/spanish ethnicity as determined by the

patient/family.

Note(s):

If the patient has multiple race origins, specify them using the other race selections in

addition to this one.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Hispanic Ethnicity - Other Hispanic/Latino/Spanish Origin:

Having origins in any of the originals peoples in other Hispanic, Latino or Spanish

territories.

Source: U.S. Office of Management and Budget. Classification of Federal Data on

Race and Ethnicity

Seq. #: 2200 Name: Patient Zip Code

Coding Instructions: Indicate the patient's United States Postal Service zip code of their primary residence.

Note(s):

The Patient Zip Code will display in the Demographics Section of the data collection form however the coding instructions will remain in the Episode of Care Section in the

data dictionary.

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: HispEthnicityCuban

Parent Seq #:

Parent Name: Hispanic or Latino

Ethnicity

Parent Value:

**Missing Data:** Report

Harvested: Yes (DCR,PINN)

> Format: Text (Categorical)

Default Value: **Usual Range:** 

Valid Range:

DataSource: User

**Technical Specifications** 

ShortName: HispEthnicityOtherO

rigin

2076 Parent Seq #:

**Parent Name:** Hispanic or Latino

Ethnicity

**Parent Value:** Yes

Missing Data: Report

> Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

**Default Value: Usual Range:** 

Valid Range:

DataSource: User

**Technical Specifications** ShortName: ZipCode

Parent Seg #: **Parent Name:** 

Parent Value:

Missing Data: Report

> Yes (DCR,PINN) Harvested:

Format: Text (10) Default Value: NULL

**Usual Range:** Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

Seg. #: 4000 Name: Coronary Artery Disease

Coding Instructions: Indicate if the patient has been diagnosed with Coronary Artery Disease (CAD).

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Coronary Artery Disease:

A history of coronary artery disease (CAD) is evidenced by one of the following:

1. Currently receiving medical treatment for CAD

2. History of Myocardial Infarction

3. Prior CV intervention including, but not limited to, CABG and/or PCI

Source: STS

User

**Technical Specifications** 

Report

Yes (DCR,PINN)

Text (Categorical)

ShortName: CAD

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

Default Value:

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

Seq. #: 4002 Name: Coronary Artery Disease Date

Coding Instructions: Indicate the documented date of diagnosis of coronary artery disease. If no diagnosis

date is recorded, indicate the first encounter date where coronary artery disease was

recorded. If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

<u>Technical Specifications</u>

ShortName: CAD\_Date

Parent Seq #: 4000

Parent Seq #: 4000
Parent Name: Coronary Artery

Disease

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4010 Name: Atrial Fibrillation or Flutter

Coding Instructions: Indicate if the patient has been diagnosed with atrial fibrillation or atrial flutter.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Atrial Fibrillation:

A cardiac arrhythmia arising from the atrium with an atrial rate 300 bpm and an irregularly irregular ventricular response in the presence of conduction. AF can be further characterized as:

- · First diagnosed
- Paroxysmal AF: AF that terminates spontaneously or with intervention within 7 days of onset. Episodes may recur with variable frequency.
- Persistent AF: Continuous AF that is sustained >7 days
- Long-standing Persistent AF: Continuous AF >12 months in duration.
- Permanent AF: The term "permanent AF" is used when the patient and clinician make a joint decision to stop further attempts to restore and/or maintain sinus rhythm.

  Acceptance of AF represents a therapeutic attitude on the part of the patient and clinician rather than an inherent pathophysiological attribute of AF. Acceptance of AF may change as symptoms, efficacy of therapeutic interventions, and patient and clinician preferences evolve.
- Nonvalvular AF: AF in the absence of rheumatic mitral stenosis, a mechanical or bioprosthetic heart valve, or mitral valve repair

Source: 2014 AHA/ACC/HRS Guideline for the Management of Patients With Atrial Fibrillation

	ShortName:	Afib
	Parent Seq #:	
	Parent Name:	
	Parent Value:	
	Missing Data:	Report
	Harvested:	Yes (DCR,PINN)
	Format:	Text (Categorical)
	Default Value:	No
	Usual Range:	
	Valid Range:	
ı	l .	

User

**Technical Specifications** 

Seq. #: 4012 Name: Atrial Fibrillation or Flutter Date

Coding Instructions: Indicate the documented date of diagnosis of atrial fibrillation/flutter. If no diagnosis date

is recorded, indicate the first encounter date where atrial fibrillation/flutter was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: Afib\_Date

Parent Seq #: 4010

DataSource:

Parent Name: Atrial Fibrillation or

Flutter

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4020 Name: Dyslipidemia

Coding Instructions: Indicate if the patient has been diagnosed with dyslipidemia.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Dyslipidemia:

National Cholesterol Education Program criteria and includes documentation of the following:

1. Total cholesterol greater than 200 mg/dL (5.18 mmol/l); or

2. Low-density lipoprotein (LDL) greater than or equal to 130 mg/dL (3.37mmol/l); or

3. High-density lipoprotein (HDL) less than 40 mg/dL (1.04 mmol/l).

For patients with known coronary artery disease, treatment is initiated if LDL is greater than 100 mg/dL (2.59 mmol/l), and this would qualify as hypercholesterolemia.

Source: 2013 ACCF/AHA Key Data Elements and Definitions for Measuring the Clinical Management and Outcomes of Patients With Acute Coronary Syndromes and Coronary Artery Disease

Technical Specifications

**Technical Specifications** 

Dyslipidemia

Report

User

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

ShortName: Dyslipidemia\_Date

Parent Seq #: 4020

Parent Name: Dyslipidemia

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User

Seq. #: 4022 Name: Dyslipidemia Date

Coding Instructions: Indicate the documented date of diagnosis of dyslipidemia. If no diagnosis date is

recorded, indicate the first encounter date where dyslipidemia was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)





#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4030 Name: Hypertension

Coding Instructions: Indicate if the patient has been diagnosed with hypertension.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No1 Yes

Supporting Definitions: Hypertension:

Hypertension is defined by any one of the following:

1. History of hypertension diagnosed and treated with medication, diet and/or exercise.

2. Prior documentation of blood pressure greater than 140 mm Hg systolic and/or 90 mm Hg diastolic for patients without diabetes or chronic kidney disease, or prior documentation of blood pressure greater than 130 mm Hg systolic and/or 80 mm Hg diastolic on at least two occasions for patients with diabetes or chronic kidney disease.

3. Currently on pharmacologic therapy for treatment of hypertension.

Source: 2013 ACCF/AHA Key Data Elements and Definitions for Measuring the Clinical Management and Outcomes of Patients With Acute Coronary Syndromes and Coronary Artery Disease

Parent Seq #:
Parent Name:
Parent Value:
Missing Data: Report

**Technical Specifications** 

ShortName: Hypertension

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No
Usual Range:
Valid Range:

DataSource: User

Seq. #: 4032 Name: Hypertension Date

Coding Instructions: Indicate the documented date of diagnosis of hypertension. If no diagnosis date is

recorded, indicate the first encounter date where hypertension was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

ShortName: Hypertension\_Date
Parent Seq #: 4030
Parent Name: Hypertension
Parent Value: Yes
Missing Data: No Action

**Technical Specifications** 

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

Name: Heart Failure **Seq. #**: 4040

Coding Instructions: Indicate if the patient has been diagnosed with heart failure (HF).

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

> 0 No Yes

Supporting Definitions: Heart Failure:

Seq. #: 4042 Name: Heart Failure Date

Selections: (none)

Supporting Definitions: (none)

Physician documentation or report of any of the following symptoms of heart failure prior to this care encounter described as unusual dyspnea on light exertion, recurrent dyspnea occurring in the supine position, fluid retention, low cardiac output secondary to cardiac dysfunction; or the description of rales, jugular venous distension, or pulmonary edema. A previous hospital admission with principal diagnosis of heart failure is considered evidence of heart failure history.

Date of first onset may be helpful.

Source: ACC/AHA 2005 Key Data Elements and Definitions for Measuring the Clinical Management and Outcomes of Patients With Chronic Heart

recorded, indicate the first encounter date where heart failure was recorded.

Coding Instructions: Indicate the documented date of diagnosis of heart failure. If no diagnosis date is

If multiple diagnosis dates exist indicate the earliest value.

**Technical Specifications** 

HF\_Date

**Technical Specifications** 

Report

User

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #: **Parent Name:** 

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

ShortName: Parent Seg #:

> **Parent Name:** Heart Failure

Parent Value: Yes

Missing Data: No Action

> Yes (DCR,PINN) Harvested:

> > Format: Date (mm/dd/yyyy)

**Default Value:** 

**Usual Range:** Valid Range:

DataSource:

**Seg.** #: 4050 Name: Heart Failure new diagnosis (within 12 months)

Target Value: The first value on current encounter

Coding Instructions: Indicate if the patient has been diagnosed with heart failure (HF) within the last 12 months.

Target Value: Any occurrence between 12 months prior to current encounter and completion of current

encounter

Selections: Code Selection Text Definition

> No 0 Yes

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: HF\_New\_Dia

4040 Parent Seq #:

**Parent Name:** Heart Failure

Parent Value: Yes

Missing Data: Report

> Harvested: Yes (PINN)

Text (Categorical) Format:

**Default Value:** 

**Usual Range:** Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

Name: Stable Angina **Seq. #:** 4060

Coding Instructions: Indicate if the patient has been diagnosed with stable angina.

Note(s):

Angina without a change in frequency or pattern for the 6 weeks prior to this visit.

Angina is controlled by rest and/or oral or transcutaneous medications.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

> No 0

Yes 1

Name: Stable Angina Date

Supporting Definitions: (none)

**Seq. #:** 4062

**Technical Specifications** 

**Technical Specifications** 

StableAngina

Report

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #: **Parent Name:** 

Parent Value:

Missing Data:

**Default Value: Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

Coding Instructions: Indicate the documented date of diagnosis of stable angina. If no diagnosis date is Parent Seq #: 4060

recorded, indicate the first encounter date where stable angina was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

ShortName: StableAngina\_Date

User

**Parent Name:** Stable Angina

Parent Value: Yes

Missing Data: No Action

> Yes (DCR,PINN) Harvested:

Format: Date (mm/dd/yyyy)

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource: User

Name: Stable Angina new diagnosis (within 12 months) **Seq. #**: 4070

Coding Instructions: Indicate if the patient has been diagnosed with stable angina within the past 12 months.

Target Value: Any occurrence between 12 months prior to current encounter and completion of current

encounter

Selections: Code Selection Text Definition

> No 0

Yes

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: StableAngina\_New\_

4060 Parent Seq #:

**Parent Name:** Stable Angina

Parent Value:

Missing Data: Report

> Harvested: Yes (PINN)

> > Format: Text (Categorical)

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource: User





	B. Diagnoses/Conditions/Comorbodities		
Seq. #: 4080 Name:	Unstable Angina	Technical S	pecifications
<b>Seq.</b> #: 4000 Name.	Unstable Angina	ShortName:	UnStableAngina
Coding Instructions:	Indicate if the patient has been diagnosed with unstable angina.	Parent Seq #:	
Target Value:	Any occurrence between birth and completion of current encounter	Parent Name:	
Selections:	Code Selection Text Definition	Parent Value:	
ociodions.	Code Selection reac Definition	Missing Data:	Report
	0 No	Harvested:	Yes (PINN)
	1 Yes	Format:	Text (Categorical)
Supporting Definitions:	(none)	Default Value:	No
3		Usual Range:	
		Valid Range:	
		DataSource:	User
Seq. #: 4082 Name:	Unstable Angina Date		<u>-</u>
Coding Instructions:	Indicate the documented date of diagnosis of unstable angina. If no diagnosis date is	ShortName:	UnstableAngina_Da e
ooding manuchons.	recorded, indicate the first encounter date where unstable angina was recorded.	Parent Seq #:	4080
	If multiple diagnosis dates exist indicate the earliest value.	Parent Name:	Unstable Angina
		Parent Value:	Yes
Target Value	The first value on current encounter	Missing Data:	No Action
-		Harvested:	Yes (PINN)
Selections:	(none)	Format:	Date (mm/dd/yyyy)
Supporting Definitions:	(none)	Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User
Soa #: 4090 Name:	Peripheral Arterial Disease	Technical S	pecifications
•		ShortName:	PAD
Coding Instructions:	Indicate if the patient has been diagnosed with Peripheral Arterial Disease (PAD).	Parent Seq #:	
	For PAD resulting in restriction of both blood flow and oxygen to a certain organ or part of	Parent Name:	
	the body, also code 'Yes' to ischemic vessel disease (IVD).	Parent Value:	
		Missing Data:	Report
Target Value:	Any occurrence between birth and completion of current encounter	Harvested:	Yes (DCR,PINN)
Selections:	Code Selection Text Definition	Format:	Text (Categorical)
		Default Value:	No
	0 No	Usual Range:	
	1 Yes	Valid Range:	
Supporting Definitions:	(none)	DataSource:	User





Seq. #: 4092 Name: Peripheral Arterial Disease Date		Technical S	pecifications
<b>Seq. #:</b> 4092 <b>Name</b> :	Seq. m. 4032 Name. I onphoral Attorial Bloodso Bato		PAD_Date
Coding Instructions:	Indicate the documented date of diagnosis of peripheral artery disease. If no diagnosis date is recorded, indicate the first encounter date where peripheral artery disease was	Parent Seq #:	4090
	recorded.	Parent Name:	Peripheral Arterial Disease
	If multiple diagnosis dates exist indicate the earliest value.	Parent Value:	Yes
		Missing Data:	No Action
Target Value:	The first value on current encounter	Harvested:	Yes (DCR,PINN)
Selections	(none)	Format:	Date (mm/dd/yyyy)
		Default Value:	NULL
Supporting Definitions:	(none)	Usual Range:	
		Valid Range:	
		DataSource:	User
Seg #- 4100 Name	PAD - Acute Limb Ischemia	Technical S	pecifications
•		ShortName:	PADAcuteLimblsch
Coding Instructions:	Indicate if the patient has been diagnosed with Acute Limb Ischemia as a result of Peripheral Arterial Disease (PAD).	Parent Seq #: Parent Name:	
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:	
Selections	Code Selection Text Definition	Missing Data:	Report
Ociocuons.	Code Selection rext Definition	Harvested:	Yes (DCR,PINN)
	0 No	Format:	Text (Categorical)
	1 Yes	Default Value:	No
Supporting Definitions:	(none)	Usual Range:	
		Valid Range:	
		DataSource:	User
Soa # /102 Name:	PAD - Acute Limb Ischemia Date	Technical S	pecifications
•	Indicate the documented date of diagnosis of Acute Limb Ischemia. If no diagnosis date is	ShortName:	PADAcuteLimblsch_ Date
·	recorded, indicate the first encounter date where acute limb ischemia was recorded.	Parent Seq #:	4100
	If multiple diagnosis dates exist indicate the earliest value.	Parent Name:	PAD - Acute Limb Ischemia
		Parent Value:	Yes
Target Value:	The first value on current encounter	Missing Data:	No Action
Selections	(none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions:	(none)	Format:	Date (mm/dd/yyyy)
Supporting Deminions.	, ()	Default Value:	NULL
		Usual Range:	
		Valid Range:	
	DataSource:	User	





Seq. #: 4110 Name: PAD - Claudication					Technical Specifications	
Seq. #. 4110 Name. 1710 Oldudoulon				ShortNam	e: PADClaud	
Coding Instructions: Indicate if the patient has been diagnosed with claudication as a result of peripheral arterial disease (PAD).				heral Parent Seq Parent Name		
Target Value: Any occurrence between birth and encounter				Parent Value	<b>)</b> :	
Selections:	: Code	Selection Text	Definition	Missing Data	: Report	
		GOIOGION TOXE	Dominion	Harvested	: Yes (DCR,PINN)	
	0	No		Forma	t: Text (Categorical)	
	1	Yes		Default Value	: No	
Supporting Definitions:	(none)			Usual Range	<b>:</b> :	
				Valid Range	<b>:</b> :	
				DataSource	: User	
Seq. #: 4112 Name:	· PAD - Cla	udication Date		<u>Technica</u>	al Specifications	
•					e: PADClaud_Date	
Coding Instructions:			osis of claudication. If no diagnosis date ate where claudication was recorded.	Farein Seq		
	If multiple die	annois datas sviet indicate	the corlinatively	Parent Name		
	ir muitipie dia	agnosis dates exist indicate	e the earliest value.	Parent Value		
				Missing Data		
Target Value:	The first valu	e on current encounter		Harvested	, ,	
Selections:	: (none)			Forma	( 33337	
Supporting Definitions	(none)			Default Value		
Supporting Definitions:	(none)			Usual Range		
				Valid Range	<b>:</b> :	
				DataSource		
Seq. #: 4120 Name:	: PAD - Crit	ical Limb Ischemia			al Specifications	
•			and unitable Contained I limb I and a series and a series		e: PADCritLimbIsch	
Coding instructions:		e patient has been diagnos rterial Disease (PAD).	ed with Critical Limb Ischemia as a resu	Parent Seq		
Target Value:	Any occurrer	nce between birth and com	pletion of current encounter	Parent Value	<b>:</b> :	
Selections:	: Code	Selection Text	Definition	Missing Data	: Report	
233333		CO.OGIOTI TOM	25	Harvested	: Yes (DCR,PINN)	
	0	No		Forma	t: Text (Categorical)	
	1	Yes		Default Value	: No	
Supporting Definitions:	Supporting Definitions: (none)			Usual Range	<b>:</b> :	
., 5	. ,			Valid Range	):	
				DataSource	: User	





O #- 4400 Nome-	PAD - Critical Limb Ischemia Date	Technical S	<u>pecifications</u>
Seq. #: 4122 Name:	FAD - Chilcal Limb ischemia Date	ShortName:	PADCritLimblsch_D
	Indicate the documented date of diagnosis of critical limb ischemia. If no diagnosis date is recorded, indicate the first encounter date where critical limb ischemia was recorded.		ate
	,	Parent Seq #: Parent Name:	4120 PAD - Critical Limb
	If multiple diagnosis dates exist indicate the earliest value.	Parent Name:	Ischemia
		Parent Value:	Yes
Target Value:	The first value on current encounter	Missing Data:	No Action
Selections:	(none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions:	(none)	Format:	Date (mm/dd/yyyy)
Supporting Demillions.	(none)	Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User
Soa #: 4130 Name:	PAD - Foot/Leg Cellulitis	Technical S	<u>pecifications</u>
•		ShortName:	PADFootCell
	Indicate if the patient has been diagnosed with foot/leg cellulitis as a result of peripheral arterial disease (PAD).	Parent Seq #: Parent Name:	
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:	
Selections:	Code Selection Text Definition	Missing Data:	Report
	Dominion Dominion	Harvested:	Yes (DCR,PINN)
	0 No	Format:	Text (Categorical)
	1 Yes	Default Value:	No
Supporting Definitions:	(none)	Usual Range:	
•		Valid Range:	
		DataSource:	User
Soa # /132 Name:	PAD - Foot/Leg Cellulitis Date	Technical S	<u>pecifications</u>
3eq. #. 4132 Name.	TAD TOOVEG Cellulitis Date	ShortName:	PADFootCell_Date
_	Indicate the documented date of diagnosis of foot/leg cellulitis. If no diagnosis date is recorded, indicate the first encounter date where foot/leg cellulitis was recorded. If multiple diagnosis dates exist indicate the earliest value.	Parent Seq #: Parent Name:	4130 PAD - Foot/Leg Cellulitis
Target Value:	The first value on current encounter	Parent Value:	Yes
Selections:		Missing Data:	No Action
Sciecuolis.	(none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions:	(none)	Format:	Date (mm/dd/yyyy)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User





#### B. Diagnoses/Conditions/CoMorbodities

Seg. #: 4140 Name:	Technical S	Technical Specifications		
<b>Seq. #:</b> 4140 <b>Name:</b>	ShortName:	PADLowExtOst		
Coding Instructions:	<b>Coding Instructions:</b> Indicate if the patient has been diagnosed with lower extremity Osteomyeleitis as a result of peripheral arterial disease (PAD) with or without limb ischemia.			
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:		
Selections:	Code Selection Text Definition	Missing Data:	Report	
	Solid Solidion Fox	Harvested:	Yes (DCR,PINN)	
	0 No	Format:	Text (Categorical)	
	1 Yes	Default Value:	No	
Supporting Definitions:	(none)	Usual Range:		
		Valid Range:		
		DataSource:	User	
Seg #: 4142 Name:	PAD - Lower Extremity Osteomyeleitis Date	Technical S	Specifications	
•		ShortName:	PADLowExtOst_Dat e	
Coding Instructions:	Indicate the documented date of diagnosis of lower extremity osteomyeleitis. If no diagnosis date is recorded, indicate the first encounter date where lower extremity	Parent Seq #:	4140	
	osteomyeleitis was recorded.	Parent Name:	PAD - Lower	
	If multiple diagnosis dates exist, indicate the earliest value.		Extremity Osteomyeleitis	
		Parent Value:	Yes	
Target Value:	The first value on current encounter	Missing Data:	No Action	
Selections:	(none)	Harvested:	Yes (DCR,PINN)	
Supporting Definitions:	(none)	Format:	Date (mm/dd/yyyy)	
Supporting Deminions.	(None)	Default Value:	NULL	
		Usual Range:		
		Valid Range:		
		DataSource:	User	
Seg #: 4150 Name:	Diabetes Mellitus (any)	Technical S	Specifications	
•		ShortName:	Diabetes	
Coding Instructions:	Indicate if the patient has a history of diabetes mellitus regardless of the type of diabetes mellitus, the duration of disease or need for antidiabetic agents.	Parent Seq #:		
		Parent Name:		
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:	_	
Selections:	Code Selection Text Definition	Missing Data:	Report	
	a Na	. Harvested:	Yes (DCR,PINN)	
	0 No	Format:	Text (Categorical)	
	1 Yes	Default Value:	No	
Supporting Definitions:	Diabetes Mellitus:	Usual Range:		
	The American Diabetes Association criteria (33) include documentation of the following: 1. Hemoglobin A1c 6.5%; or	Valid Range:	Haar	
	DataSource:	User		
	<ol> <li>Fasting plasma glucose 126 mg/dL (7.0 mmol/L); or</li> <li>2-h Plasma glucose 200 mg/dL (11.1 mmol/L) during an oral glucose tolerance test; or</li> <li>In a patient with classic symptoms of hyperglycemia or hyperglycemic crisis, a random plasma glucose 200 mg/dL (11.1 mmol/L) This does not include gestational diabetes.</li> </ol>			

Source: 2013 ACCF/AHA Key Data Elements and Definitions for Measuring the Clinical Management and Outcomes of Patients With Acute Coronary

Syndromes and Coronary Artery Disease





#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4152 Name: Diabetes Mellitus Date

Coding Instructions: Indicate the documented date of diagnosis of diabetes. If no diagnosis date is recorded,

indicate the first encounter date where diabetes was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: Diabetes\_Date

**Parent Seq #:** 4150

Parent Name: Diabetes Mellitus

(any)

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User

Seg. #: 4160 Name: Diabetes Mellitus Type I

Coding Instructions: Indicate if the patient has been diagnosed with Diabetes Mellitus Type I.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Diabetes Mellitus Type I:

Type 1 diabetes is a condition characterized by high blood glucose levels caused by a total lack of insulin. Occurs when the body's immune system attacks the insulin-producing beta cells in the pancreas and destroys them. The pancreas then produces little or no insulin.

Source: American Diabetes Association

**Technical Specifications** 

ShortName: DiabMellTypel

Parent Seq #: 4150

Parent Name: Diabetes Mellitus

(any)

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Text (Categorical)

Default Value: N

Usual Range: Valid Range:

DataSource: User

Seg. #: 4162 Name: Diabetes Mellitus Type I Date

Coding Instructions: Indicate the earliest documented patient diagnosis date of Diabetes Mellitus Type I.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: DiabMellTypeI\_Date

Parent Seq #: 4160

Parent Name: Diabetes Mellitus

Type I

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

	b. Diagnoses/Conditions/Comorbodities			
Con #. 4170 Name.	Diabetes Mellitus Type II	Technical Specifications		
<b>Seq. #</b> : 4170 <b>Name</b> :	Diabetes Mellitus Type II	ShortName:	DiabMellTypeII	
Coding Instructions:	Indicate if the patient has been diagnosed with Diabetes Mellitus Type II.	Parent Seq #:	4150	
Target Value:	Any occurrence between birth and completion of current encounter	Parent Name:	Diabetes Mellitus (any)	
Selections:	Code Selection Text Definition	Parent Value:	Yes	
		Missing Data:	Report	
	0 No	Harvested:	Yes (DCR)	
	1 Yes	Format:	Text (Categorical)	
Supporting Definitions:	Diabetes Mellitus Type II:	Default Value:	No	
	Type 2 diabetes is a condition characterized by high blood glucose levels caused by	Usual Range:		
	either a lack of insulin or the body's inability to use insulin efficiently.	Valid Range:		
	Source: American Diabetes Association	DataSource:	User	
	Technical Specifications			
Seq. #: 4172 Name:	Diabetes Mellitus Type II Date		-	
Coding Instructions:	Indicate the earliest documented patient diagnosis date of Diabetes Mellitus Type II.	Shorthame.	DiabMellTypeII_Dat e	
·	,	Parent Seq #:	4170	
_	The first value on current encounter	Parent Name:	Diabetes Mellitus Type II	
Selections:	(none)	Parent Value:	Yes	
Supporting Definitions:	(none)	Missing Data:	No Action	
		Harvested:	Yes (DCR)	
		Format:	Date (mm/dd/yyyy)	
		Default Value:	NULL	
		Usual Range:		
		Valid Range:		
		DataSource:	User	
Seq. #: 4180 Name:	Pro-diabates	Technical S	pecifications	
Seq. #: 4100 Name:	rie-ulabeles	ShortName:	PreDiabetes	
Coding Instructions:	Indicate if the patient has been diagnosed with pre-diabetes.	Parent Seq #:	4150	
Target Value:	Any occurrence between birth and completion of current encounter	Parent Name:	Diabetes Mellitus (any)	
Selections:	Code Selection Text Definition	Parent Value:	Yes	
		I	_	

Report

User

Yes (DCR)

Text (Categorical)

**Missing Data:** 

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

No

Yes

Source: American Diabetes Association

0

Supporting Definitions: Pre-diabetes:

Pre-diabetes is a condition in which blood glucose levels are higher than normal but are not high enough for a diagnosis of diabetes. People with pre-diabetes are at increased

risk for developing Type 2 diabetes and for heart disease and stroke. Other names for

pre-diabetes are impaired glucose tolerance and impaired fasting glucose.





#### B. Diagnoses/Conditions/CoMorbodities

**Technical Specifications** Name: Pre-diabetes Date Seq. #: 4182 PreDiabetes\_Date ShortName: Coding Instructions: Indicate the earliest documented patient diagnosis date of pre-diabetes. Parent Seq #: 4180 Parent Name: Pre-diabetes Target Value: The first value on current encounter Parent Value: Yes Selections: (none) Missing Data: No Action Yes (DCR) Harvested: Supporting Definitions: (none) Format: Date (mm/dd/yyyy) **Default Value: Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Diabetic Peripheral Neuropathy Sea. #: 4190 ShortName: DiabPheriNeuro Coding Instructions: Indicate if the patient has documented diabetic peripheral neuropathy. Parent Seq #: **Parent Name:** Target Value: Any occurrence between birth and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (DCR) No 0 Format: Text (Categorical) Yes 1 **Default Value:** Supporting Definitions: Diabetic Peripheral Neuropathy: **Usual Range:** Peripheral neuropathy is nerve damage that affects the feet, legs, or hands. Peripheral

Seg. #: 4192 Name: Diabetic Peripheral Neuropathy Date

Coding Instructions: Indicate the documented date of diagnosis of Diabetic Peripheral Neuropathy. If no

neuropathy causes pain, numbness, or a tingling feeling.

Source: American Diabetes Association

diagnosis date is recorded, indicate the first encounter date where hypertension was

recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: DiabPheriNeuro\_Dat

User

Parent Seq #: 4190

Valid Range:

DataSource:

Parent Name: Diabetic Peripheral

Neuropathy

**Parent Value:** Yes

Missing Data: No Action

> Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

**Usual Range:** Valid Range:

DataSource: User





Co. # 4200 Name	eq. #: 4200 Name: Diabetic Autonomic Neuropathy		Technical Specifications		
Seq. #: 4200 Name:			ShortName:	DiabAutoNeuro	
Coding Instructions:	Indicate if the	e patient has documente	ed diabetic autonomic neuropathy.	Parent Seq #:	
Target Value: Any occurrence between birth and completion of current encounter		Parent Name:			
rarget value.	Any occurren	ice between birth and t	completion of current encounter	Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
Supporting Definitions:	0	No		Harvested:	Yes (DCR)
	1	Yes	Format:	Text (Categorical)	
	ı			Default Value:	No
	Diabetic Autonomic Neuropathy:		Usual Range:		
	Autonomic neuropathy is a type of neuropathy affecting the lungs, heart, stomach, intestines, bladder or genitals.		Valid Range:		
	,	ğ	41	DataSource:	User
	Source: American Diabetes Association				

	Source. American Diabetes Association			
• " 1000 N		Technical S	Technical Specifications	
•	Diabetic Autonomic Neuropathy Date  Indicate the documented date of diagnosis of Diabetic Autonomic Neuropathy. It		DiabAutoNeuro_Dat e	
county men actions.	diagnosis date is recorded, indicate the first encounter date where hypertension recorded.  If multiple diagnosis dates exist indicate the earliest value.		4200 Diabetic Autonomic Neuropathy	
	in maniple diagnosis dates exist maleate the edificat value.	Parent Value:	Yes	
Target Value:	The first value on current encounter	Missing Data: Harvested:	No Action Yes (DCR)	
Selections:	(none)	Format:	Date (mm/dd/yyyy)	
Supporting Definitions:	(none)	Default Value:	NULL	
Supporting Demittons.	(none)	Usual Range:		
		Valid Range:		
		DataSource:	User	
0 - 11 4040 Name	" 4040 Names Dighetic Detinenathy		Technical Specifications	
Seq. #: 4210 Name: Diabetic Retinopathy		ShortName:	DiabRetinopathy	
•	Indicate if the patient has documented diabetic retinopathy.	Parent Seq #: Parent Name:		
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:		
Selections:	Code Selection Text Definition	Missing Data:	Report	
	o No	Harvested:	Yes (DCR)	
	1 Yes	Format:	Text (Categorical)	
		Default Value:	No	
Supporting Definitions:	Diabetic Retinopathy:	Usual Range:		
	Diabetic retinopathy or retinopathy is an eye disease that is caused by damage small blood vessels in the retina. Loss of vision may result.	valid Range: DataSource:	User	
	Source: American Diabetes Association	Datassurce.	0301	





#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4212 Name: Diabetic Retinopathy Date

Coding Instructions: Indicate the documented date of diagnosis of Diabetic Retinopathy. If no diagnosis date is

recorded, indicate the first encounter date where hypertension was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: DiabRetino\_Date

Parent Seq #: 4210

Parent Name: Diabetic Retinopathy

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULI

Usual Range: Valid Range:

DataSource: User

Seq. #: 4220 Name: Ischemic Vascular Disease

Coding Instructions: Indicate if the patient has documented ischemic vascular disease.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

1 Yes

Supporting Definitions: Ischemic Vascular Disease:

Ischemic vascular disease entails a clogging of the arteries that results in restriction of both blood flow and oxygen to a certain organ or part of the body. This could result in a

number of problems that are dependent upon the location of the blockage.

Source:

Technical Specifications

ShortName: IVD

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: No
Usual Range:

Valid Range:

DataSource: User

Seg. #: 4222 Name: Ischemic Vascular Disease Date

Coding Instructions: Indicate the documented date of diagnosis of ischemic vascular disease. If no diagnosis

date is recorded, indicate the first encounter date where ischemic vascular disease was

recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: IVD\_Date

Parent Seq #: 4220

Parent Name: Ischemic Vascular

Disease

Parent Value: Yes

Missing Data: No Action
Harvested: Yes (PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User





#### B. Diagnoses/Conditions/CoMorbodities

O #- 4000 Nome	Poriphoral Vaccular Disease	Technical S	Technical Specifications	
<b>Seq. #:</b> 4230 <b>Name:</b>	Peripheral Vascular Disease	ShortName:	PVD	
Coding Instructions:	ndicate if the patient has documented peripheral vascular disease.	Parent Seq #:		
Target Value	Any occurrence between birth and completion of current encounter	Parent Name:		
_	·	Parent Value:		
Selections:	Code Selection Text Definition	Missing Data:	Report	
	o No	Harvested:	Yes (PINN)	
	1 Yes	Format:	Text (Categorical)	
Cumparting Definitions	none)	Default Value:	No	
Supporting Definitions:	ione)	Usual Range:		
		Valid Range:		
		DataSource:	User	
Seq. #: 4232 Name:	Peripheral Vascular Disease Date	<u>Technical S</u>	pecifications	
•	·	ShortName:	PVD_Date	
Coding Instructions:	ndicate the documented date of diagnosis of peripheral vascular disease. ate is recorded, indicate the first encounter date where peripheral vascula ecorded.		4230 Peripheral Vascular	
			Disease	
	multiple diagnosis dates exist indicate the earliest value	Parent Value:	Yes	
		Missing Data:	No Action	
Target Value:	he first value on current encounter	Harvested:	Yes (PINN)	
Selections:	none)	Format:	Date (mm/dd/yyyy)	
Commandina Definitions	nana)	Default Value:	NULL	
Supporting Definitions:	ione)	Usual Range:		
		Valid Range:		
		DataSource:	User	
Seg. #: 4240 Name:	Chronic Kidney Disease	<u>Technical S</u>	pecifications	
•		ShortName:	CKD_History	
-	ndicate if the patient has documented chronic kidney disease.	Parent Seq #: Parent Name:		
Target Value:	Any occurrence between birth and completion of current encounter	Parent Value:		
Selections:	Code Selection Text Definition	Missing Data:	Report	
	n No	Harvested:	Yes (DCR,PINN)	
	.,	Format:	Text (Categorical)	
	•	Default Value:	No	
Supporting Definitions:	Chronic Kidney Disease/Renal Insufficiency:	Usual Range:		
	Chronic kidney disease is defined as either kidney damage or GFR 60 mL for 3 months.	· · · · · · · · · · · · · · · · · · ·		
	Kidney damage is defined as pathologic abnormalities or markers of dama abnormalities in blood or urine tests or imaging studies.	age, including	User	

Indicate the patient's stage of disease:

- \* Stage 0: No known kidney disease
- \* Stage 1: Kidney damage with normal or high GFR 90 mL/min/1.73 m2
- \* Stage 2: Kidney damage with mildly decreased GFR 60 89 mL/min/1.73 m2
- \* Stage 3: Moderately decreased GFR 30 59 mL/min/1.73 m2
- \* Stage 4: Severely decreased GFR 15 29 mL/min/1.73 m2
- \* Stage 5: Kidney failure GFR 15 mL/min/1.73 m2 or on dialysis

Source: 2012 ACCF/AHA/ACR/SCAI/SIR/STS/SVM/SVN/SVS Key Data Elements and Definitions for Peripheral Atherosclerotic Vascular Disease



#### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4242 Name: Chronic Kidney Disease Date

Coding Instructions: Indicate the documented date of diagnosis of chronic kidney disease. If no diagnosis date

is recorded, indicate the first encounter date where chronic kidney disease was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: CKD\_Date

Parent Seq #: 4240

Parent Name: Chronic Kidney

Disease

Parent Value: Yes

Missing Data: No Action

**Harvested:** Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

ShortName:

Parent Seq #: Parent Name:

**Parent Value:** 

Missing Data:

Default Value:

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

DataSource: User

**Technical Specifications** 

CLD\_History

Report

No

User

Yes (DCR,PINN)

Text (Categorical)

Seq. #: 4250 Name: Chronic Liver Disease

Coding Instructions: Indicate if the patient has documented cirrhosis or chronic liver disease.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

Yes

0 No

Supporting Definitions: Chronic Liver Disease/Hepatic Dysfunction:

Hepatic dysfunction is defined as dysfunction of the liver that results in hypoalbuminemia (<2 grams/dL), coagulopathy (PT > 1.5 x upper limits of normal), and hyperbilirubinemia (> 3.0 x upper limits of normal). Code as "Yes" if the patient develops 2 out of these 3

laboratory abnormalities.

Source: STS

**Technical Specifications** 

ShortName: CLD\_Date

Parent Seq #: 4250

Parent Name: Chronic Liver

Disease

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User

Seq. #: 4252 Name: Chronic Liver Disease Date

Coding Instructions: Indicate the documented date of diagnosis of chronic liver disease. If no diagnosis date is

recorded, indicate the first encounter date where chronic liver disease was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)





### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4260 Name: Metabolic Syndrome

Coding Instructions: Indicate if the patient has been diagnosed with metabolic syndrome.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

0 No1 Yes

Supporting Definitions: Metabolic Syndrome:

Metabolic syndrome is a name for a group of risk factors that occur together and increase the risk for coronary artery disease, stroke, and type 2 diabetes.

Metabolic syndrome is present if you have three or more of the following signs:

- Blood pressure equal to or higher than 130/85 mmHg

- Fasting blood sugar (glucose) equal to or higher than 100 mg/dL

- Large waist circumference (length around the waist):

- Men - 40 inches or more

- Women - 35 inches or more

- Low HDL cholesterol:

- Men - under 40 mg/dL- Women - under 50 mg/dL

- Triglycerides equal to or higher than 150 mg/dL

Source: U.S. National Library of Medicine's MedlinePlus

**Technical Specifications** 

ShortName: MetaSyndro

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Text (Categorical)

Default Value: No Usual Range:

Valid Range:

DataSource: User

**Technical Specifications** 

• " 1000 Name	Matabalia	Cum drama Data		<u>recnnicai s</u>	pecifications
<b>Seq.</b> #: 4262 Name:	Metabolic	Syndrome Date		ShortName:	MetaSyndro_Date
Coding Instructions:	Indicate the	earliest documented patie	ent diagnosis date of Metabolic Syndrome.	Parent Seq #:	4260
				Parent Name:	Metabolic Syndrome
Target Value:	The first valu	ue on current encounter		Parent Value:	Yes
Selections:	(none)			Missing Data:	No Action
Supporting Definitions:	(none)			Harvested:	Yes (DCR)
oupporting beamaions.	()			Format:	Date (mm/dd/yyyy)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	
				DataSource:	User
				Technical S	pecifications
Seq. #: 4263 Name:	Systemic	Embolism		ShortName:	Syst_Embo
Coding Instructions:	This element	has been retired effective	e PINNACLE v1.3.	Parent Seg #:	
				Parent Name:	
Target Value:	N/A			Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
	-			Harvested:	Yes (PINN)
	0	No		Format:	Text (Categorical)
	1	Yes		Default Value:	No
Supporting Definitions:	(none)				NO
				Usual Range:	
				Valid Dango:	
				Valid Range:	

User

DataSource:





### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4264 Name:	Prior Stro	ke or TIA	Technical S	Specifications
00q: #: 1201 Hamo!			ShortName:	PriorStrokeCVA
Coding Instructions:	This elemen	t has been retired effective PINNACLE v1.3.	Parent Seq #:	
Target Value:	N/A		Parent Name:	
Selections:	Codo	Selection Text Definition	Parent Value:	
ociections.		Selection rext Definition	Missing Data:	Report
	0	No	Harvested:	Yes (PINN)
	1	Yes	Format:	Text (Categorical)
			Default Value:	No
Supporting Definitions:	(none)		Usual Range:	
			Valid Range:	
			DataSource:	User
O # 4070 Nome.	Contropo	ragio	Technical S	pecifications
<b>Seq. #</b> : 4270 <b>Name</b> :	Gasiropa	6515	ShortName:	Gastroparesis
<b>Coding Instructions:</b>	Indicate if th	e patient has documented gastroparesis.	Parent Seg #:	
			Parent Name:	
•	•	nce between birth and completion of current encounter	Parent Value:	
Selections:	Code	Selection Text Definition	Missing Data:	Report
	0	No	Harvested:	Yes (DCR)
	· ·	Yes	Format:	Text (Categorical)
	1	165	Default Value:	No
Supporting Definitions:	Gastropare	esis:	Usual Range:	
		sis is a form of neuropathy that affects the stomach. Digestion of food may be	Valid Range:	
	control diffic	or delayed, resulting in nausea, vomiting, or bloating, making blood glucose sult.	DataSource:	User

Sea #: 4272	Name:	Gastroparesis	Date
-------------	-------	---------------	------

Coding Instructions: Indicate the documented date of diagnosis of gastroparesis. If no diagnosis date is

Source: American Diabetes Association

recorded, indicate the first encounter date where hypertension was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical S	pecifications
ShortName:	Gastroparesis_Date
Parent Seq #: Parent Name:	4270 Gastroparesis
Parent Value:	Yes
Missing Data:	No Action
Harvested:	Yes (DCR)
Format:	Date (mm/dd/yyyy)
Default Value:	NULL
Usual Range:	
Valid Range:	
DataSource:	User





### B. Diagnoses/Conditions/CoMorbodities

**Technical Specifications** Name: Erectile Dysfunction (men) **Seq. #:** 4280 ShortName: ErectDysfun Coding Instructions: Indicate if the patient has documented erectile dysfunction. Parent Seq #: **Parent Name:** Target Value: Any occurrence between birth and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (DCR) 0 No Format: Text (Categorical) Yes **Default Value:** Supporting Definitions: Erectile Dysfunction (men): **Usual Range:** Impotence of organic origin. Valid Range:

Seq. #: 4282 Name: Erectile Dysfunction Date

Coding Instructions: Indicate the documented date of diagnosis of erectile dysfunction. If no diagnosis date is

recorded, indicate the first encounter date where hypertension was recorded.

If multiple diagnosis dates exist indicate the earliest value.

Source: ICD-9-CM 607.84/ICD-10-CM 607.84

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Seq. #:** 4290 Name: Depression

Coding Instructions: Indicate if the patient has documented depression.

Target Value: Any occurrence between birth and completion of current encounter

Selections: Code Selection Text Definition

> No 0

Yes

Supporting Definitions: Depression:

Depression is characterized by depressed or sad mood, diminished interest in activities which used to be pleasurable, weight gain or loss, psychomotor agitation or retardation, fatigue, inappropriate guilt, difficulties concentrating, as well as recurrent thoughts of

death

Source: CDC

**Technical Specifications** 

ShortName: ErectDysfun\_Date

User

Parent Seq #: 4280

DataSource:

Parent Name: **Erectile Dysfunction** 

(men)

Parent Value: Yes

Missing Data: No Action

> Harvested: Yes (DCR)

Date (mm/dd/yyyy)

Format: **Default Value:** NULL

**Usual Range:** Valid Range:

DataSource: User

**Technical Specifications** 

ShortName: Depression

Parent Seq #:

**Parent Name:** 

Parent Value:

Missing Data: Report

> Harvested: Yes (DCR)

Format: Text (Categorical)

Default Value: No

**Usual Range:** 

Valid Range:

DataSource: User



### B. Diagnoses/Conditions/CoMorbodities

Seq. #: 4292 Name: Depression Date

**Coding Instructions:** Indicate the earliest documented patient diagnosis date of Depression.

Target Value: The first value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

**ShortName:** Depression\_Date

Parent Seq #: 4290

Parent Name: Depression

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User

#### C. Event

Seq. #: 5135 Name: Event ID

Coding Instructions: Indicate all patient's history of cardiac events.

Target Value: Any occurrence between birth and start of current encounter

Selections: (none)

Supporting Definitions: E001 - Myocardial Infarction:

A myocardial infarction is evidenced by any of the following:

1. A rise and fall of cardiac biomarkers (preferably troponin) with at least one of the values in the abnormal range for that laboratory [typically above the 99th percentile of the upper reference limit (URL) for normal subjects] together with at least one of the following manifestations of myocardial ischemia:

a. Ischemic symptoms.

b. ECG changes indicative of new ischemia (new ST-T changes, new left bundle branch block, or loss of R- wave voltage).

c. Development of pathological Q- waves in 2 or more contiguous leads in the ECG (or equivalent findings for true posterior MI).

d. Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.

e. Documentation in the medical record of the diagnosis of acute myocardial infarction based on the cardiac biomarker pattern in the absence of any items enumerated in a-d due to conditions that may mask their appearance (e.g., peri-operative infarct when the patient cannot report ischemic symptoms; baseline left bundle branch block or ventricular pacing).

2. ECG changes associated with prior myocardial infarction can include the following (with or without prior symptoms):

a. Any Q-wave in leads V2-V3 >=0.02 seconds or QS complex in leads V2 and V3. b. Q-wave >=0.03 seconds and >=0.1 mV deep or QS complex in leads I, II, aVL, aVF, or V4-V6 in any two leads of a contiguous lead grouping (I, aVL, V6; V4-V6; II, III, and aVF). c. R-wave >=0.04 seconds in V1-V2 and R/S >=1 with a concordant positive Twave in the absence of a conduction defect.

3. Imaging evidence of a region with new loss of viable myocardium at rest in the absence of a non-ischemic cause. This can be manifest as:

a. Echocardiographic, CT, MR, ventriculographic or nuclear imaging evidence of left ventricular thinning or scarring and failure to contract appropriately (i.e., hypokinesis, akinesis, or dyskinesis).

b. Fixed (non-reversible) perfusion defects on nuclear radioisotope imaging (e.g., MIBI, thallium).

4. Medical record documentation of prior myocardial infarction.

Source: Joint ESC-ACC-AHA-WHF 2007 Task Force Consensus Document"Universal Definition of Myocardial Infarction".

#### E002 - PCI - Bare Metal Stent Impant:

Indicate the date the patient had a Percutaneous Coronary Intervention (PCI) that involved the implantation of a Bare Metal Stent (BMS).

Source

#### E003 - PCI - Drug Eluting Stent Implant:

Indicate the date the patient had a Percutaneous Coronary Intervention (PCI) that involved the implantation of a Drug Eluting Stent (DES).

Source:

#### E004 - PCI - Other (non-stent) Intervention:

Indicate the date the patient had a Percutaneous Coronary Intervention (PCI) that involved balloon angioplasty. This does not include the implant of a Bare Metal or Drug Eluting Stent.

Source:

**Technical Specifications** 

ShortName: EventID

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (4)

Default Value: NUI

Usual Range: Valid Range:

DataSource: User

#### E005 - Systemic Embolism:

Indicate if the patient has been diagnosed with a systemic embolism.

Source:

#### E006 - Minor Hemorrhage:

The patient had a documented minor hemorrhage - regardless of location.

A minor hemorrhage is either clinically overt but not major or occult (e.g., asymptomatic guaiac-positive stoolA major hemorrhage is one which leads to transfusion of at least 2 units of whole blood or erythrocytes, requires hospitalization or surgery, results in permanent disability, or involves a critical anatomic site; or any bleeding event the physician characterizes as "major".

#### Source

#### E007 - Intercranial Hemorrhage:

Indicate if the patient had an intracranial hemorrhage.

Intracranial hemorrhage is defined as bleeding into or around the brain potentially caused by one of the following:

- Hemorrhagic conversion of a primary ischemic stroke
- Subarachnoid hemorrhage
- Intracerebral hemorrhage
- Other (including subdural and epidural hematomas)
- Unknown

#### Note

- If the patient had an intracranial hemorrhage with a loss off brain function, intracranial hemorrhage (event ID E007) and hemorrhagic stroke (event ID E016) should be specified.
- 2. If the patient had an intracranial hemorrhage without loss of brain function only, intracranial hemorrhage (event ID E007).

#### Source

#### E008 - NICM Hemorrhage (location unknown):

Indicate if the patient had a documented major hemorrhage - outside of the cranium.

A major hemorrhage is one which leads to transfusion of at least 2 units of whole blood or erythrocytes, requires hospitalization or surgery, results in permanent disability, or involves a critical anatomic site; or any bleeding event the physician characterizes as "major".

#### Source:

#### E009 - NICM Hemorrhage Location - Intra-articular (Atraumatic):

Indicate if the patient had a documented major hemorrhage within a joint.

Source:

#### E010 - NICM Hemorrhage Location - Intra-ocular:

Bleeding associated with abrupt deterioration of visual acuity.

Source:

#### E011 - NICM Hemorrhage Location - Intra-spinal:

Indicate if the patient had a documented major hemorrhage within the spinal.

Source:

#### **E012 - NICM Hemorrhage Location - Pericardial:**

Indicate if the patient had a documented major hemorrhage around the heart.

Source:

#### E013 - NICM Hemorrhage Location - Retroperitoneal/Abdominal:

Indicate if the patient had a documented major hemorrhage around the abdomen.

#### **E027 - Permanent Pacemaker:**

Indicate if the patient has a permanent pacemaker. Event must not be selected if the patient had the device previously, but the device is no longer in place.

Source:

#### E014 - TIA:

The patient had a transient ischemic attack (TIA).

A transient ischemic attack (TIA) is a brief episode of loss of blood flow to part of the brain resulting in transient stroke-like symptoms. Most symptoms of a TIA disappear within an hour, although they may last for up to 24 hours and include:

- Numbness or weakness, especially on one side of the body
- Confusion or trouble speaking or understanding speech
- Trouble seeing in one or both eyes

Source:

#### E021 - Cardioversion:

The patient received an electrical or pharmacological cardioversion, whether successful or unsuccessful.

Source:

#### E015 - Ischemic Stroke:

The patient had a documented ischemic stroke.

An ischemic stroke is a loss of neurological function caused when a blood vessel that supplies blood to the brain is blocked.

Source

#### E028 - Vascular Comlication (requiring intervention):

Indicate if the patient had a documented vascular complication intervention.

Vascular complications can include, but are not limited to, access site occlusions, peripheral embolizations, dissections, pseudoaneurysms and/or AV fistulas. Any noted vascular complication must have had an intervention such as a fibrin injection, angioplasty, or surgical repair to qualify. Prolonged pressure does not qualify as an intervention, but ultrasonic guided compression after making a diagnosis of pseudoaneurysm does qualify.

Source:

#### E016 - Hemorrhagic Stroke:

The patient had a hemorrhagic stroke.

Hemorrhagic stroke is defined as bleeding into or around the brain that results in transient or permanent neurological deficit.

#### Note:

- 1. If the patient had an intracranial hemorrhage with a loss off brain function, intracranial hemorrhage (event ID E007) and hemorrhagic stroke (event ID E016) should be specified.
- 2. If the patient had an intracranial hemorrhage without loss of brain function only, intracranial hemorrhage (event ID E007).

Source

#### E017 - Coronary Artery Bypass Graft:

The patient had coronary artery bypass graft (CABG) surgery.

#### E026 - PTCA:

Indicate if the patient received percutaneous transluminal coronary angioplasty (PTCA).

PTCA is a minimally invasive procedure to open up blocked coronary arteries, allowing blood to circulate unobstructed to the heart muscle. This angioplasty is a non-stent balloon angioplasty.

Note: If a stent is used during angiography, PTCA (event E026) must not be selected.

Source:

#### E024 - CRT-D:

The patient received a cardiac resynchronization therapy device and defibrillator (CRT-D).

A CRT-D is a biventricular pacemaker that sends electrical signals to both ventricles as well as a defibrillator. It may or may not have an atrial pacing wire.

#### Note:

1. Event must not be selected if the patient had the device previously, but the device is no longer in place.

Source:

#### E018 - Cardiac Valve Surgery:

The patient had cardiac valve surgery.

Source:

#### **E019 - Heart Transplantation:**

The patient had a heart transplantation surgery.

Source:

#### E025 - ICD:

The patient has an implantable cardioverter defibrillator (ICD).

Event must not be selected if the patient had the device previously, but the device is no longer in place.

Source:

#### **E020 - Cardiac Therapeutic Procedure:**

The patient had any procedure to treat a pathologic structural, or pathophysiological functional, disorder of the heart.

Source:

#### E022 - LVAD:

The patient has a left ventricular assist device (LVAD).

An LVAD is a mechanical pump that temporarily and artificially aids the natural pumping action of the left ventricle.

#### Note:

1. Event must not be selected if the patient had the device previously, but the device is no longer in place.

Source:

#### E023 - CRT:

Indicate if the patient received a cardiac resynchronization therapy (CRT) device.

A CRT device is a biventricular pacemaker that sends electrical signals to both ventricles that resynchronizes the heart chambers and helps it pump more effectively. It may or may not have an atrial pacing wire.

#### Note:

1. Event must not be selected if the patient received a cardiac resynchronization therapy device and defibrillator (CRT-D).

#### E039 - Gestational Diabetes Mellitus (GDM):

Gestational Diabetes Mellitus is a type of diabetes mellitus that develops only during pregnancy and usually disappears upon delivery, but increases the risk that the mother will develop diabetes later. GDM is managed with meal planning, activity, and, in some cases, insulin.

Source: American Diabetes Association

#### E040 - Diabetic ketoacidosis (DKA):

Diabetic ketoacidosis (DKA) is an emergency condition in which extremely high blood glucose levels, along with a severe lack of insulin, result in the breakdown of body fat for energy and an accumulation of ketones in the blood and urine. Signs of DKA are nausea and vomiting, stomach pain, fruity breath odor and rapid breathing. Untreated DKA can lead to coma and death.

Source: American Diabetes Association

#### E041 - Hyperosmolar Hyperglycemic Syndrome (HHS):

Hyperosmolar hyperglycemic nonketotic syndrome is an emergency condition in which one's blood glucose level is very high and ketones are not present in the blood or urine. If HHNS is not treated, it can lead to coma or death.

Source: American Diabetes Association

#### E042 - Hypoglycemia (Severe):

Hypoglycemia is a condition that occurs when one's blood glucose is lower than normal, usually less than 70 mg/dL. Signs include hunger, nervousness, shakiness, perspiration, dizziness or light-headedness, sleepiness, and confusion. If left untreated, hypoglycemia may lead to unconsciousness. Hypoglycemia is treated by consuming a carbohydraterich food such as a glucose tablet or juice. It may also be treated with an injection of glucagon if the person is unconscious or unable to swallow. Also called an insulin reaction.

Source: American Diabetes Association

#### **E049 - Acute Pancreatitis:**

Acute pancreatitis is a sudden attack causing inflammation of the pancreas and usually associated with severe upper abdominal pain. The pain may last several days and may be serious

Source:

#### E050 - Bariatric Surgery:

Indicate if the patient has undergone bariatric surgery. Bariatric surgery can include:

- Adjustable gastric banding (AGB)
- Roux-en-Y gastric bypass (RYGB)
- Biliopancreatic diversion with a duodenal switch (BPD-DS)
- Vertical sleeve gastrectomy (VSG)

Source:

#### E051 - Bariatric Surgery - Adjustable Gastric Banding:

A type of bariatric surgery that involves inserting a thin, inflatable ring or gastric band to create a new, smaller stomach pouch.

Unlike conventional gastric bypass surgery, gastric band surgery is: Minimally invasive no cutting, stapling, or re-routing of the intestinal tract. Reversible and adjustable.

Source:

#### E052 - Bariatric Surgery - Biliopancreatic diversion with duodenal switch:

A type of bariatric surgery in which a large portion of the stomach is left intact, including the pyloric valve that regulates the release of contents from the stomach into the small intestine. The duodenum is divided near this valve, and the small intestine divided as well. The portion of the small intestine connected to large intestine is attached to the short duodenal segment next to the stomach. The remaining segment of the duodenum connected to the pancreas and gallbladder is attached to this limb closer to the large intestine

#### E053 - Bariatric Surgery - Roux-en-Y gastric bypass:

A type of bariatric surgery that reduces the size of your stomach to a small pouch – about the size of an egg. It does this by stapling off a section of it. This reduces the amount of food you can take in at meals. The surgeon then attaches this pouch directly to the small intestine, bypassing most of the rest of the stomach and the upper part of the small intestine. This reduces the amount of fat and calories you absorb from the foods you are able to eat for even more weight loss.

RYGB can be done as an open surgery, with a large cut (incision) on your abdomen to reach your stomach. Or it can be done as a laparoscopic RYGB, using a lighted tube with a tiny camera, called a laparoscope.

#### Source:

#### E054 - Bariatric Surgery - Vertical Sleeve gastrectomy:

A type of bariatric surgery that generates weight loss by restricting the amount of food (and therefore calories) that can be eaten by removing 85% or more of the stomach without bypassing the intestines or causing any gastrointestinal malabsorption.

#### Source:

#### E055 - Foot Ulcer:

Ulcers are slow healing wounds on the skin. Diabetic foot ulcers occur on the feet of people with type 1 and type 2 diabetes

#### Source:

#### E056 - Gout:

a disease in which defective metabolism of uric acid causes arthritis, especially in the smaller bones of the feet, deposition of chalkstones, and episodes of acute pain.

#### Source:

#### E057 - Hemodialysis:

Healthy kidneys clean your blood and remove extra fluid in the form of urine. They also make substances that keep your body healthy.

In hemodialysis, a dialysis machine and a special filter called an artificial kidney, or a dialyzer, are used to clean your blood. To get your blood into the dialyzer, the doctor needs to make an access, or entrance, into your blood vessels. This is done with minor surgery, usually to your arm.

#### Source: National Kidney Foundation

#### E058 - Hyperthyroidism:

Hyperthyroidism is a disorder that occurs when the thyroid gland makes more thyroid hormone than the body needs. Hyperthyroidism is sometimes called thyrotoxicosis, the technical term for too much thyroid hormone in the blood. Thyroid hormones circulate throughout the body in the bloodstream and act on virtually every tissue and cell in the body. Hyperthyroidism causes many of the body's functions to speed up.

#### Source: HHS

#### E059 - Hypothyroidism:

Hypothyroidism is a disorder that occurs when the thyroid gland does not make enough thyroid hormone to meet the body's needs. Thyroid hormone regulates metabolism the way the body uses energy and affects nearly every organ in the body. Without enough thyroid hormone, many of the body's functions slow down

#### Source: HHS

#### E063 - Nonalcoholic Fatty Liver Disease (NAFLD):

NAFLD ist he build up of extra fat in liver cells that is not caused by alcohol. It is normal for the liver to contain some fat. However, if more than 5% - 10% percent of the liver's weight is fat, then this is considered NAFLD.

Source: American Liver Foundation



#### C. Event

#### E064 - Sleep Apnea:

A sleep disorder characterized in 2 ways:

-Obstructive sleep apnea(OSA): The blockage of the airway, usually when the soft tissue in the back of the throat collapses during sleep.

-Central sleep apnea: Unlike OSA, the airway is not blocked, but the brain fails to signal the muscles to breathe due to instability in the respiratory control center.

Source:

#### E065 - Syncope:

Syncope is defined as the transient loss of consciousness and postural tone.

Source:

Seq. #: 5136 Name: Event Date

Coding Instructions: Indicate all dates, if documented, of cardiac events that occurred.

Note(s):

All occurrences on current encounter.

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

#### **Technical Specifications**

ShortName: EventDate

Parent Seq #: 5135
Parent Name: Event ID
Parent Value: Not Null

Missing Data: Report

Harvested: Yes (DCR,PINN)

Date (mm/dd/yyyy)

Default Value: NULL

Format:

**Usual Range:** 

Valid Range:

DataSource: User





#### D. Encounter Information

Name: Insurance - Private Health Insurance **Seq. #:** 3020

Coding Instructions: Indicate if the patient's insurance payor(s) included private health insurance.

Note(s):

A health maintenance organization (HMO) is considered private health insurance.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes

Supporting Definitions: Private Health Insurance:

Private health insurance is coverage by a health plan provided through an employer or union or purchased by an individual from a private health insurance company.

Source: U.S. Census Bureau

**Technical Specifications** Seq. #: 3022 Name: Insurance - Medicaid

Coding Instructions: This element has been retired effective PINNACLE v1.3.

Target Value: N/A

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: (none)

Name: Insurance - Military Health Care **Seq.** #: 3023

Coding Instructions: Indicate if the patient's insurance payor(s) included Military Health Care.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0

Yes

Supporting Definitions: Military Health Care:

Military Health care - Military health care includes TRICARE/CHAMPUS (Civilian Health and Medical Program of the Uniformed Services) and CHAMPVA (Civilian Health and Medical Program of the Department of Veterans Affairs), as well as care provided by the

Department of Veterans Affairs (VA).

Source: U.S. Census Bureau

**Technical Specifications** 

InsPrivate

Insurance - None

Yes (DCR,PINN)

Text (Categorical)

3027

Report

User

ShortName:

Parent Seq #:

**Parent Name:** 

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

ShortName: InsMedicaid

3027 Parent Seq #:

**Parent Name:** Insurance - None

Parent Value: No

Missing Data: Report

> Harvested: Yes (DCR,PINN)

> > Text (Categorical)

**Default Value:** 

Format:

**Usual Range:** Valid Range:

DataSource: User

Technical Specifications

InsMilitary ShortName:

3027 Parent Seq #:

Parent Name: Insurance - None

Parent Value:

Missing Data: Report

> Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource: User





#### D. Encounter Information

Seq. #: 3024 Name: Insurance - State Specific Plan (non-Medicaid)

Coding Instructions: Indicate if the patient's insurance payor(s) included State-specific Plan.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: State Specific Plan:

State-specific plan - Some states have their own health insurance programs for low-income uninsured individuals. These health plans may be known by different names in

different states. (Non-Medicaid)

Source: U.S. Census Bureau

Technical Specifications

User

**Technical Specifications** 

InsState

Insurance - None

Yes (DCR,PINN)

Text (Categorical)

3027

Nο

Report

ShortName:

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

ShortName: InsIHS

Parent Seq #: 3027

Parent Name: Insurance - None

Parent Value: No

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User

Seq. #: 3025 Name: Insurance - Indian Health Service

Coding Instructions: Indicate if the patient's insurance payor(s) included Indian Health Service (IHS).

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Indian Health Service:

Indian Health Service (IHS) is a health care program through which the Department of Health and Human Services provides medical assistance to eligible American Indians at IHS facilities. In addition, the IHS helps pay the cost of selected health care services provided at non-IHS facilities.

Source: U.S. Census Bureau

Seq. #: 3026 Name: Insurance - Non-US Insurance

Coding Instructions: Indicate if the patient's insurance payor(s) included Non-US Insurance.

**Target Value:** The value on current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Non-US Insurance:

Non-U.S. Insurance refers to individuals with a payor that does not originate in the

United States.

Source: U.S. Census Bureau

**Technical Specifications** 

ShortName: InsNonUS

Parent Seq #: 3027

Parent Name: Insurance - None

Parent Value: No

Missing Data: Report

Harvested: Yes (DCR,P)

arvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No

**Usual Range:** 

Valid Range:

DataSource: User





#### D. Encounter Information

**Technical Specifications** Name: Insurance - None **Seq. #:** 3027 ShortName: InsNone Coding Instructions: Indicate if the patient has no insurance payor(s). Parent Seq #: **Parent Name:** Target Value: The value on current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Yes (DCR,PINN) Harvested: 0 No Format: Text (Categorical) Yes **Default Value:** Supporting Definitions: None: **Usual Range:** None refers to individuals with no or limited health insurance thus, the individual is the Valid Range: payor regardless of ability to pay. DataSource:

Name: Insurance - Medicare (Fee for service) Seq. #: 3028

Source: NCDR

Coding Instructions: Indicate if the patient's insurance payor(s) included Medicare Fee for Service.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

> No 0 Yes 1

Supporting Definitions: Medicare:

Medicare is the Federal program which helps pay health care costs for people 65 and older and for certain people under 65 with long-term disabilities.

The traditional system of reimbursement under health insurance and Medicare. Health care providers bill patients for services supplied, and costs are shared according to a contractual agreement between the patient and insurance company. A fee-for-service system allows patients maximum flexibility in the choice of providers and services.

Source: U.S. Census Bureau

#### **Technical Specifications** ShortName: InsMedicare\_Feefor Ser

User

Parent Seq #: 3027

**Parent Name:** Insurance - None

Parent Value: Nο Missing Data: Report

> Yes (DCR,PINN) Harvested:

Format: Text (Categorical)

**Default Value: Usual Range:** Valid Range:

DataSource: User





#### D. Encounter Information

Seq. #: 3029 Name: Insurance - Medicare (Managed care)

Coding Instructions: Indicate if the patient is insured by Medicare (managed care/HMO).

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Medicare:

Medicare is the Federal program which helps pay health care costs for people 65 and

older and for certain people under 65 with long-term disabilities.

A type of Medicare Advantage Plan that is available in some areas of the country. In most managed care plans, you can only go to doctors, specialists, or hospitals on the plan's list. Plans must cover all Medicare Part A and Part B health care. Some managed care plans cover extras, like prescription drugs. Your costs may be lower than in Original

Source: U.S. Census Bureau

Technical Specifications

**Technical Specifications** 

3027

Report

InsMedicare\_MngdC

Insurance - None

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #:

**Parent Name:** 

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range: DataSource:

Harvested:

Format:

ShortName: InsMedicaid\_Feefor

User

Ser

Parent Seq #: 3027

Parent Name: Insurance - None

Parent Value: No

Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User

Seq. #: 3030 Name: Insurance - Medicaid (Fee for service)

Coding Instructions: Indicate if the patient's insurance payor(s) included Medicaid Fee for Service.

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Supporting Definitions: Medicaid:

Medicaid is a program administered at the state level, which provides medical assistance to the needy. Families with dependent children, the aged, blind, and disabled who are in financial need are eligible for Medicaid. It may be known by different names in different states.

Source: U.S. Census Bureau

Seq. #: 3031 Name: Insurance - Medicaid (Managed Care)

Coding Instructions: Indicate if the patient's insurance payor(s) included Medicaid (managed care/HMO).

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: Medicaid:

Medicaid is a program administered at the state level, which provides medical assistance to the needy. Families with dependent children, the aged, blind, and disabled who are in financial need are eligible for Medicaid. It may be known by different names in different

states.

Source: U.S. Census Bureau

**Technical Specifications** 

ShortName: InsMedicaid\_MngdC

are

Parent Seq #: 3027

Parent Name: Insurance - None

Parent Value: No
Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: No Usual Range:

Valid Range:

DataSource: User





#### D. Encounter Information

**Technical Specifications** Name: Payer ID Seq. #: 3100 ShortName: PayerID Coding Instructions: Indicate the Payer ID of the patient's primary insurance payer. Payer ID is a national Parent Seq #: numbering system that identifies healthcare payers authorized by CMS for healthcare **Parent Name:** claims processing and other electronic data interchange transactions. Parent Value: Target Value: The value on current encounter Missing Data: Report Selections: (none) Harvested: Yes (DCR,PINN) Format: Text (5) Supporting Definitions: (none) Default Value: NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Seq. #: 6000 Name: Height (in) ShortName: Ht inches Coding Instructions: Indicate the patient's Height in inches (in). Parent Seq #: **Parent Name:** Target Value: The value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Decimal (5,2) **Default Value:** NULL **Usual Range:** Valid Range: 7.87-102.36 DataSource: User **Technical Specifications** Name: Height (cm) **Seq.** #: 6001 ShortName: Ht\_cms Coding Instructions: Indicate the patient's Height in centimeters (cm). Parent Seq #: **Parent Name:** Target Value: The value on current encounter **Parent Value:** Selections: (none) **Missing Data:** Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Decimal (5,2) **Default Value:** NULL **Usual Range:** Valid Range: 20.00-260.00 DataSource: User





O. v. W. COAO. Names Systelia Blood Propoura	Technical S	pecifications
Seq. #: 6010 Name: Systolic Blood Pressure	ShortName:	SystolicBP
Coding Instructions: Indicate the patient's systolic blood pressure in mmHg.	Parent Seq #:	
Target Value: The value on current encounter	Parent Name:	
	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Integer (3)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	1-300
	DataSource:	User
Seq. #: 6011 Name: Diastolic Blood Pressure		pecifications
•	ShortName:	DiastolicBP
Coding Instructions: Indicate the patient's diastolic blood pressure in mmHg.	Parent Seq #: Parent Name:	
Target Value: The value on current encounter		
Selections: (none)	Parent Value:	Danant
	Missing Data: Harvested:	Report Yes (DCR,PINN)
Supporting Definitions: (none)	Format:	Integer (3)
	Default Value:	NULL
	Usual Range:	NOLL
	Valid Range:	1-200
	DataSource:	User
		pecifications
Seq. #: 6015 Name: Heart Rate	ShortName:	
Coding Instructions: Indicate the patient's heart rate in beats per minute.	Parent Seq #:	. 100.111 10.10
	Parent Name:	
Target Value: The value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (PINN)
	Format:	Integer (3)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	1-300
	DataSource:	User



O #- COOO N	Joight (lbs)	Technical S	Specifications
<b>Seq.</b> #: 6020 Name:	veignt (ibs)	ShortName:	Wt_lbs
Coding Instructions:	dicate the patient's weight in pounds (lbs).	Parent Seq #:	6025
Target Value:	he value on current encounter	Parent Name:	Patient unable to be weighed
Selections:	none)	Parent Value:	No
Supporting Definitions:	one)	Missing Data:	Report
Supporting Definitions:	one)	Harvested:	Yes (DCR,PINN)
		Format:	Decimal (6,2)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	22.00-1540.00
		DataSource:	User
Seq. #: 6021 Name:	Veight (kg)	<u>Technical S</u>	Specifications
		ShortName:	Wt_kgs
Coding Instructions:	dicate the patient's weight in kilograms (kg).	Parent Seq #:	6025
Target Value:	he value on current encounter	Parent Name:	Patient unable to be weighed
Selections:	none)	Parent Value:	No
Supporting Definitions	one)	Missing Data:	Report
Supporting Definitions:	one)	Harvested:	Yes (DCR,PINN)
		Format:	Decimal (5,2)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	10.00-700.00
		DataSource:	User
Sea #: 6025 Name:	atient unable to be weighed	<u>Technical S</u>	Specifications
•			CannotWeigh
_	dicate if the patient was unable to be weighed during the	Parent Seq #: Parent Name:	
•	he value on current encounter	Parent Value:	
Selections:	Code Selection Text Definition	Missing Data:	Report
	No	Harvested:	Yes (DCR,PINN)
		Format:	Text (Categorical)
		Default Value:	No
Supporting Definitions:	one)	Usual Range:	
		Valid Range:	
		DataSource:	User





O # 0000 N-	Woist Circ	umforonos (in)		<u>Technical S</u>	Specifications
<b>Seq. #</b> : 6026 <b>Name</b> :	waist Circ	umerence (m)		ShortName:	WaistCir_inches
Coding Instructions:	Indicate the p	patient's waist circumferen	ce in inches (in).	Parent Seq #:	
Target Value	The value on	current encounter		Parent Name:	
_		Tourism officialities		Parent Value:	
Selections:	(none)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (DCR)
				Format:	Decimal (5,2)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 6027 Name:	Waist Circ	umference (cm)			Specifications
Coding Instructions:	Indicate the p	patient's waist circumferen	ce in centimeters (cm).		WaistCir_cm
<b>3</b>			,	Parent Seq #: Parent Name:	
Target Value:	The value on	current encounter		Parent Value:	
Selections:	(none)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (DCR)
oupporting benintions.	()			Format:	Decimal (5,2)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 6030 Name:	Tobacco I	lse		<u>Technical S</u>	Specifications
•				ShortName:	TobaccoUse
Coding Instructions:		patient's use of tobacco pro igars, pipe) and smokeless	oducts. Tobacco products include smoke s (chewing tobacco).	Parent Seq #: Parent Name:	
Target Value:	The value on	current encounter		Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	•
				Harvested:	Yes (DCR,PINN)
	1	Never		Format:	Text (Categorical)
	2	Current		Default Value:	NULL
	3	Quit within past 12 months		Usual Range:	
	4	Quit more than 12 months ago		Valid Range: DataSource:	User
	5	Tobacco Screening not performed for medical reasons			
Supporting Definitions:	(none)				





	Olamani II			Technical S	pecifications
<b>Seq.</b> #: 6035 Name:	Cigarettes	3		ShortName:	Cigarettes
Coding Instructions:	Indicate if the	e patient is a cigarette smo	oker currently or quit within the past 12 months.	Parent Seq #:	6030
Townst Value	The color by	-t 40thit-		Parent Name:	Tobacco Use
Selections:		Selection Text	current encounter and current encounter  Definition	Parent Value:	Current, Quit within past 12 months
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
				Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 6036 Name:	Cigars			Technical S	pecifications
seq. #: 0030 Name:	Olyais			ShortName:	Cigars
Coding Instructions:	Indicate if the	e patient is a cigar smoker	r currently or quit within the past 12 months.	Parent Seq #:	6030
Target Value	The value he	etween 12 months prior to	current encounter and current encounter	Parent Name:	Tobacco Use
Selections:		Selection Text	Definition	Parent Value:	Current, Quit within past 12 months
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
				Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 6037 Name:	Dino			Technical S	pecifications
Seq. #: 003/ Name.	ripe			ShortName:	Pipe
Coding Instructions:	Indicate if the	e patient is a pipe smoker	currently or quit within the past 12 months.	Parent Seq #:	6030
Target Value	The value be	otwoon 12 months prior to	current encounter and current encounter	Parent Name:	Tobacco Use
Selections:		Selection Text	Definition	Parent Value:	Current, Quit within past 12 months
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
				Usual Range:	
				Valid Range:	
				DataSource:	User





Seq. #: 6038 Name:			Technical S	Specifications	
<b>Seq. #:</b> 0038 <b>Name:</b>	SHIOKEIESS	3		ShortName:	Smokeless
Coding Instructions:	Indicate if the	patient uses smokele	ess tobacco currently or quit within the past 12 months.	Parent Seq #:	6030
Target Value	The value he	twoon 12 months aris	r to current encounter and current encounter	Parent Name:	Tobacco Use
Selections:		Selection Text	Definition	Parent Value:	Current, Quit within past 12 months
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
5				Usual Range:	
				Valid Range:	
				DataSource:	User
0 # 0040 Name	Constinu C	Pagastian Caumas	lin n	Technical S	Specifications
<b>Seq. #:</b> 6040 <b>Name:</b>	Smoking C	essation Counse	ling	ShortName:	SmokeCounsel
Coding Instructions:	Coding Instructions: Indicate if the patient received smoking cessation counseling for smoking cessation if they		Parent Seq #:	6030	
	are a current	smoker or quit within	12 months.	Parent Name:	Tobacco Use
	Note(s):			Parent Value:	Current, Quit within
		ective PINNACLE v1.3 this element is specific to counseling only. For armacological therapy code the specific medication prescribed.			past 12 months
	pnarmacolog	ical therapy code the	specific medication prescribed.	Missing Data:	Report
Target Value	Any occurren	oco botwoon start of su	urrent encounter and completion of current encounter	Harvested:	Yes (DCR,PINN)
•	•	ice between start of ct	arrent encounter and completion of current encounter	Format:	Text (Categorical)
Selections:	Code	Selection Text	Definition	Default Value:	NULL
	0	No		Usual Range:	
	0	Yes		Valid Range:	
	1	169		DataSource:	User
Supporting Definitions:	(none)				
	Datiant		evious encounter in the past	Technical S	Specifications

0 " COAE Names	Datient co	kad during any pravi	ious apacintar in the past	Technical S	Specifications
Seq. #: 6045 Name:		s about the use of tol	lous encounter in the past bacco	ShortName:	UseofTobacco_24m onths
Coding Instructions:	Indicate of the about the use		ng any previous encounter in the past 24 months,	Parent Seq #: Parent Name:	
Target Value:	Any occurre	nce between 24 months p	rior to current encounter and completion of current	Parent Value:	
	encounter			Missing Data:	Report
Selections:	Code	Selection Text	Definition	Harvested:	Yes (DCR,PINN)
	0	No		Format:	Text (Categorical)
				Default Value:	NULL
	1	Yes		Usual Range:	
Supporting Definitions:	(none)			Valid Range:	
				DataSource:	User





#### D. Encounter Information

Name: Alcohol History **Seq. #**: 6047

Coding Instructions: Indicate the patient estimate of alcohol consumption.

Target Value: The value on current encounter

Parent Value: Selections: Code Selection Text Definition Missing Data: 1 None 2 One or fewer alcoholic drinks per week **Default Value:** 3 2 to 7 alcoholic drinks **Usual Range:** per week 8 to 14 alcoholic drinks per week 15 or more alcoholic 5

Supporting Definitions: (none)

Name: Advance Care Plan Discussed or Discussion of Advance **Seq.** #: 6050

drinks per week

Care Plan Documented

Coding Instructions: For patients 65 and older, indicate if an advance care plan was documented in the

medical record or the creation of an advance care plan was discussed with the patient or

surrogate decision maker.

Target Value: The value between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition No Selection Retired (v1.4) 0 Yes There was documentation that Advance Care 1 Planning was discussed or there is documentation of a advance care plan or surrogate decision maker in the medical record. No - Not documented There is no documentation as to the reason why 2 advance care was not discussed. Patient reason could include a situation where No - patient reason 3 the patient's cultural and/or spiritual beliefs preclude a discussion of advance care planning, as it would be viewed as harmful to the patient's

> This could also include documentation that Patient did not Wish or was not able to Name a Surrogate Decision Maker or Provide an Advance

beliefs and thus harmful to the physician-patient

Care Plan.

relationship.

Supporting Definitions: (none)

recillical S	pecifications
ShortName:	AdvCarePlanDiscus sed
Parent Seq #: Parent Name:	
Parent Value:	
Missing Data:	Report
Harvested:	Yes (DCR,PINN)

Text (Categorical)

NULL

User

Format:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Tachnical Specifications

**Technical Specifications** 

Alcohol\_Hist

Report

NULL

User

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #: **Parent Name:** 

Harvested:

Valid Range:

DataSource:

Format:





Default Value: Usual Range:

Valid Range:

DataSource:

User

#### D. Encounter Information

**Technical Specifications** Name: Patient screened for evidence of nephropathy **Seq. #:** 6055 ShortName: PatScrEviNephro Coding Instructions: Indicate if the patient was screened or had evidence of nepropathy. Evidence of Parent Seq #: nephropathy can be considered if any of these apply: **Parent Name:** microalbuminuria or macroalbuminuria test result documented and reviewed OR documentation of treatment for nephropathy (e.g. patient receiving dialysis, patient being Parent Value: treated for End Stage Renal Disease, or any visit to a nephrologist in the chart) OR patient receiving ACE or ARB therapy. Missing Data: Report Harvested: Yes (DCR) Target Value: The last value on current encounter Format: Text (Categorical)

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: (none)

**Technical Specifications** Name: Discussion of Lifestyle Modifications Documented **Sea.** #: 6100 ShortName: LifeModify Coding Instructions: Indicate if the patient has a documented lifestyle modifications. Parent Seq #: **Parent Name:** Target Value: Any occurrence between start of current encounter and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (DCR,PINN) No 0 Text (Categorical) Format: Yes **Default Value:** NULL Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Patient enrolled in weight loss program WeightLossPrgm ShortName: Coding Instructions: Indicate if the patient was enrolled in a weight loss program at the time of this current visit. Parent Seq #: **Parent Name:** Target Value: Any occurrence between start of current encounter and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (DCR) No 0 Format: Text (Categorical) Yes **Default Value:** No Supporting Definitions: (none) **Usual Range:** Valid Range:

DataSource:

User





#### D. Encounter Information

**Technical Specifications** Name: Patient Education **Seq. #:** 6110 ShortName: PatientEdu Coding Instructions: Indicate if the patient has received counseling or instruction for diabetes management, Parent Seq #: cardiac symptoms or primary prevention in the past 24 months. **Parent Name:** Parent Value: Target Value: Any occurrence between 24 month prior to current encounter and completion of current encounter Missing Data: Report Selections: Code Selection Text Definition Harvested: Yes (DCR) Format: Text (Categorical) No - Patient Not 2 Counseled or Educated **Default Value:** No Counseling or **Usual Range:** 3 Education - Medical Valid Range: Reason DataSource: User Yes

Supporting Definitions: (none)

" 0400 Name	برطناهما	Diet Counceline		Technical S	Specifications
<b>Seq. #</b> : 6120 <b>Name</b> :	пеанну	Diet Counseling		ShortName:	HealthDietCounse
Coding Instructions:	Counselin	g can include any of the	ealthy diet counseling within 24 months. Healthy Diet e below: • Eating a variety of fruits, vegetables, ducts, fish, legumes, poultry, and lean meats •	Parent Seq #: Parent Name:	
				Parent Value:	
Target Value:	Any occur	rence between start of	current encounter and completion of current encounter	Missing Data:	Report
Selections:	Code	Selection Text	Definition	Harvested:	Yes (DCR)
				Format:	Text (Categorical)
	0	No		Default Value:	No
	1	Yes		Usual Range:	
Supporting Definitions: (none)		Valid Range:			
				DataSource:	User
# 6101 Name	Modicat	ion Instruction		<u>Technical S</u>	Specifications
eq. #: 6121 Name:	Medical	ion instruction		ShortName:	MedInstruct
Coding Instructions:			d patient education on medication instruction within the	Parent Seq #:	
	past 24 m	onins.		Parent Name:	
Target Value:	Any occur	rence between start of	current encounter and completion of current encounter	Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
		Colodian Tox	20mmion	Harvested:	Yes (DCR)
	0	No		Format:	Text (Categorical)
	1	Yes		Default Value:	No
Supporting Definitions:	(none)			Usual Range:	
	` ,			Valid Range:	
				DataSource:	User



**Seq.** #: 6123



## NCDR® Outpatient Registries v1.4 Data Dictionary - Full Specifications

#### D. Encounter Information

Seq. #: 6122 Name: Physical Activity Counseling

Coding Instructions: Indicate if the patient received physical activity counseling within the past 24 months.

Target Value: Any occurrence between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition

0 No

1 Yes

Name: Symptom Management

past 24 months.

Supporting Definitions: Physical Activity Counseling:

Physical activity counseling includes all levels of physical activity, including leisure activities, recreational sports, and competitive professional performance.

Can include moderate-intensity aerobic physical activity or vigorous intensity aerobic physical activity.

Source: American Diabetes Association AHA/ACC 2010 Primary Prevention Performance Measures

Coding Instructions: Indicate if the patient has received patient education on symptom management within the

Selection Text

Selection Text

No

Yes

Target Value: Any occurrence between start of current encounter and completion of current encounter

Definition

Technical Specifications

**Technical Specifications** 

PhyActCounsel

Report

User

Yes (DCR)

Text (Categorical)

ShortName:

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

ShortName: SymptMgmt

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Text (Categorical)

Default Value: No

**Usual Range:** 

Valid Range:

DataSource: User

Seg. #: 6124 Name: Weight Monitoring

Selections: Code

Supporting Definitions: (none)

Selections: Code

0

1

Coding Instructions: Indicate if the patient has received patient education on weight monitoring within the past

24 months.

Target Value: Any occurrence between start of current encounter and completion of current encounter

get value. This occurrence between that of current encounter and completion of current encounter.

Definition

0 No

1 Yes

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: WeightMonitor

Parent Seq #:

Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User





#### D. Encounter Information

Seq. #: 6130 Name: New York Heart Association Functional Classification for

**Heart Failure** 

Coding Instructions: Indicate the patient's New York Heart Association functional classification for Heart

Failure.

Target Value: The value on current encounter

Selections: Code Selection Text Definition Patient has cardiac disease but without resulting 1 limitations of ordinary physical activity. Ordinary physical activity (e.g., walking several blocks or climbing stairs) does not cause undue fatigue, palpitation, dyspnea, or anginal pain. Limiting symptoms may occur with marked exertion. Ш Patient has cardiac disease resulting in slight limitation of ordinary physical activity. Patient is comfortable at rest. Ordinary physical activity such as walking more than two blocks or climbing more than one flight of stairs results in limiting symptoms (e.g., fatigue, palpitation, dyspnea, or anginal pain). 3

III Patient has cardiac disease resulting in marked limitation of physical activity. Patient is comfortable at rest. Less than ordinary physical

activity (e.g., walking one to two level blocks or climbing one flight of stairs) causes fatigue, palpitation, dyspnea, or anginal pain.

IV Patient has dyspnea at rest that increases with any physical activity. Patient has cardiac disease resulting in inability to perform any physical activity without discomfort. Symptoms may be

undertaken, discomfort is increased.

present even at rest. If any physical activity is

Supporting Definitions: (none)

Seg. #: 6135 Name: Kansas City Cardiomyopathy Questionnaire Completed

Coding Instructions: Indicate if the patient has completed the Kansas City Cardiomyopathy Questionnaire

(KCCQ).

Target Value: Any occurrence between start of current encounter and completion of current encounter

**Selections:** Code Selection Text Definition

NoYes

Supporting Definitions: (none)

**Technical Specifications** 

**Technical Specifications** 

Report

NULL

User

Yes (DCR,PINN)

Text (Categorical)

ShortName:

Parent Seq #:

Parent Value:

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:

**ShortName:** KCCQCompleted

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report
Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User



#### **D. Encounter Information**

**Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -**Seq. #:** 6136 ShortName: **KCCQOverallScore** Overall Summary Score Parent Seq #: 6135 Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Name: Kansas City Cardiomyopathy Target Value: N/A Questionnaire Completed Selections: (none) Parent Value: Yes Missing Data: Supporting Definitions: (none) No Action Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -**Seq. #:** 6137 **KCCQClinSummSco** ShortName: Clinical Summary Score Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6135 **Parent Name:** Kansas City Target Value: N/A Cardiomyopathy Questionnaire Selections: (none) Completed Parent Value: Yes Supporting Definitions: (none) **Missing Data:** No Action Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -Seq. #: 6138 ShortName: KCCQPhysLimitScor Physical Limitation Score Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6135 **Parent Name:** Kansas City Target Value: N/A Cardiomyopathy Questionnaire Completed Selections: (none) Parent Value: Yes Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User



#### D. Encounter Information

**Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -**Seq. #:** 6139 KCCQSymStabScor ShortName: Symptom Stability Score Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6135 **Parent Name:** Kansas City Cardiomyopathy Target Value: N/A Questionnaire Selections: (none) Completed Parent Value: Yes Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (PINN) Format: Integer (3) Default Value: **NULL Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -**Seq. #:** 6140 ShortName: KCCQSelfEfficScore Self Efficacy Score Parent Seq #: 6135 Coding Instructions: This element has been retired effective PINNACLE v1.3. **Parent Name:** Kansas City Cardiomyopathy Target Value: N/A Questionnaire Completed Selections: (none) Parent Value: Yes **Missing Data:** Supporting Definitions: (none) No Action Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Kansas City Cardiomyopathy Questionnaire (KCCQ) -Seq. #: 6141 ShortName: **KCCQLifeQltyScore** Quality of Life Score Parent Seq #: 6135 Coding Instructions: This element has been retired effective PINNACLE v1.3. **Parent Name:** Kansas City Cardiomyopathy Target Value: N/A Questionnaire Completed Selections: (none) **Parent Value:** Yes Supporting Definitions: (none) **Missing Data:** No Action Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User





Com # C440 Nome	Kansas City Cardiomyopathy Questionnaire (KCCQ) -	Technical S	Specifications	
<b>Seq.</b> #: 0142 Name:	Social Limitation Score	ShortName:	KCCQSocialLimitSc ore	
Coding Instructions:	This element has been retired effective PINNACLE v1.3.	Parent Seq #:	6135	
		Parent Name:	Kansas City	
Target Value:	N/A		Cardiomyopathy Questionnaire	
Selections:	(none)		Completed	
Supporting Definitions:	(none)	Parent Value:	Yes	
3		Missing Data:	No Action	
		Harvested:	Yes (PINN)	
		Format:	Integer (3)	
		Default Value:	NULL	
		Usual Range:		
		Valid Range:		
		DataSource:	User	
O # C142 Nome	Kanaga City Cardiamyonathy Quastiannaira (KCCQ)	Technical S	pecifications	
Seq. #: 6143 Name:	Kansas City Cardiomyopathy Questionnaire (KCCQ) - Total Symptom Score	ShortName:	KCCQTotalSymScor e	
Coding Instructions:	This element has been retired effective PINNACLE v1.3.	Parent Seq #:	6135	
Tannat Value	N/A	Parent Name:	Kansas City	
Target Value: Selections:			Cardiomyopathy Questionnaire Completed	
		Parent Value:	Yes	
Supporting Definitions:	(none)	Missing Data:	No Action	
		Harvested:	Yes (PINN)	
		Format:	Integer (3)	
		Default Value:	NULL	
		Usual Range:		
		Valid Range:		
		DataSource:	User	
		Technical S	pecifications	
<b>Seq.</b> #: 6145 <b>Name</b> :	Chronic Heart Failure Questionnaire from Guyatt Completed	ShortName:	GuyattCompleted	
Coding Instructions:	Indicate if the patient completed the Chronic Heart Failure Questionnaire from Guyatt.	Parent Seq #: Parent Name:		
Target Value:	Any occurrence between start of current encounter and completion of current encounter	Parent Value:		
_		Missing Data:	Report	
Selections:	Code Selection Text Definition	Harvested:	Yes (PINN)	
	0 No	Format:	Text (Categorical)	
	1 Yes	Default Value:	No	
Supporting Definitions	(none)	Usual Range:		
Supporting Definitions:	(mone)	Valid Range:		
		DataSource:	User	





Com #1 6150 Nome	: Minnesota Living with Heart Failure Questionnaire Completed				Technical Specifications		
<b>Seq.</b> #: 6150 <b>Name</b> :					ShortName:	MLFHQCompleted	
Coding Instructions:	s: Indicate if the patient has completed the Minnesota Living with Heart Failure						
	Parent Name:						
Tanget Value	<b>A</b>	h		l completion of compact concernation	Parent Value:		
_	-	nce between start of cu	irrent encounter and	I completion of current encounter	Missing Data:	Report	
Selections:	Code	Selection Text	Definition		Harvested:	Yes (PINN)	
	0	No			Format:	Text (Categorical)	
	1	Yes			Default Value:	No	
	•	103			Usual Range:		
Supporting Definitions:	(none)				Valid Range:		
					DataSource:	User	
Seq. #: 6155 Name:	Other Too	I/Method used to a	assess Heart Fa	ilure Activity	<u>Technical S</u>	<u>pecifications</u>	
<b>604.</b> <i>III</i> • 100	Completed				ShortName:	OtherHFActvityAssm ntCompleted	
Coding Instructions:	Indicate if and	other tool/method was	used to assess the	patient's heart failure symptoms	Parent Seq #:		
	and activity o	other than the NYHA, K e or Chronic Heart Fail	Parent Name:				
	Quodidinian		are ecore from eay	att.	Parent Value:		
Target Value:	Any occurrer	nce between start of cu	irrent encounter and	I completion of current encounter	Missing Data:	Report	
Selections:	(none)				Harvested:	Yes (PINN)	
	(222)				Format:	Text (Categorical)	
Supporting Definitions:	(none)				Default Value:	No	
					Usual Range:		
					Valid Range:		
					DataSource:	User	
Can # 6200 Name	: Dyspnea F	Procent			Technical S	pecifications	
<b>Seq.</b> #: 6200 Name:	Dyspilea i	resem			ShortName:	Dyspnea	
Coding Instructions:	Indicate if the	e patient has dyspnea.			Parent Seq #:		
Target Value	The value on	n current encounter			Parent Name:		
•					Parent Value:		
Selections:	Code	Selection Text	Definition		Missing Data:	Report	
	0	No			Harvested:	Yes (PINN)	
	1	Yes			Format:	Text (Categorical)	
Cumpantina Definition	•				Default Value: Usual Range:	NULL	
Supporting Definitions:	Supporting Definitions: (none)						
					Valid Range:		
					DataSource:	User	





#### D. Encounter Information **Technical Specifications** Name: Orthopnea Present Seq. #: 6210 ShortName: Orthopnea Coding Instructions: Indicate if the patient has orthopnea. Parent Seq #: **Parent Name:** Target Value: The value on current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (PINN) 0 No Format: Text (Categorical) Yes **Default Value:** Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Rales Present **Seg. #:** 6220 ShortName: Rales Coding Instructions: Indicate if the patient has rales. Parent Seq #: **Parent Name:** Target Value: The value on current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (PINN) No 0 Format: Text (Categorical) Yes 1 **Default Value:** NULL Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Peripheral Edema Present **Seq.** #: 6230 ShortName: PeriEdema Coding Instructions: Indicate if the patient has peripheral edema. Parent Seq #: **Parent Name:** Target Value: The value on current encounter Parent Value: Selections: Code Selection Text Definition **Missing Data:** Report Harvested: Yes (PINN) No 0 Text (Categorical) Format: Yes **Default Value:** NULL Supporting Definitions: (none) **Usual Range:**

Valid Range: DataSource:

User





Target Value: The value on current encounter

### NCDR® Outpatient Registries v1.4 **Data Dictionary - Full Specifications**

**Parent Name:** 

Parent Value:

**Parent Name:** 

DataSource:

Parent Seq #:

DataSource:

User

User

#### D. Encounter Information **Technical Specifications** Name: S3 Gallop Present **Seq. #**: 6240 ShortName: S3Gallop Coding Instructions: Indicate if the patient has an S3 gallop. Parent Seq #:

Selections:	Code	Selection Text	Definition	Missing Data:	Report
•	0	No		Harvested:	Yes (PINN)
	4	Yes		Format:	Text (Categorical)
	1	163		Default Value:	NULL

Supporting Definitions: (none)

**Usual Range:** Valid Range: DataSource: User

**Technical Specifications** Name: Ascites Present **Seq. #**: 6250 ShortName: Ascites Coding Instructions: Indicate if the patient has Ascites. Parent Seq #:

Target Value: The value on current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (PINN)

No 0 Format: Text (Categorical) Yes 1 **Default Value:** NULL

Supporting Definitions: (none) **Usual Range:** Valid Range:

**Technical Specifications** Name: Hepatomegaly Present **Seq.** #: 6260 ShortName: Hepatomegaly Coding Instructions: Indicate if the patient has Hepatomegaly.

**Parent Name:** Target Value: The value on current encounter Parent Value:

Selections: Code Selection Text Definition **Missing Data:** Report Harvested: Yes (PINN) No 0 Text (Categorical) Format: Yes

**Default Value:** NULL Supporting Definitions: (none) **Usual Range:** Valid Range:





	040			Technical Specifications		
Seq. #: 6270 Name:	S4 Gallop	Present		ShortName:	S4Gallop	
Coding Instructions:	Indicate if the	e patient has an S4 gallop		Parent Seq #:		
Target Value	The value or	n current encounter		Parent Name:		
_				Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:	Report	
	0	No		Harvested:	Yes (PINN)	
	1	Yes		Format:	Text (Categorical)	
Owner and the Particular and	•			Default Value:	NULL	
Supporting Definitions:	(none)			Usual Range:		
				Valid Range:		
				DataSource:	User	
Seq. #: 6275 Name:	Jugular V	enous Distention Pre	sent	Technical S	pecifications	
•				ShortName:	JVD	
Coding Instructions:	Indicate if the	e patient has jugular venou	us distention.	Parent Seq #:		
Target Value:	The value or	n current encounter		Parent Name:		
			D # 11	Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:	Report	
	0	No		Harvested:	Yes (PINN)	
	1	Yes		Format:	Text (Categorical)	
Summerting Definitions	(none)			Default Value:	NULL	
Supporting Definitions:	(Horie)			Usual Range:		
				Valid Range:		
				DataSource:	User	
Seq. #: 6278 Name:	HF Educa	ation Completed/Docu	imented	Technical S	<u>pecifications</u>	
•				ShortName:	HFEduCompleted	
Coding Instructions:	This element	t has been retired effective	PINNACLE v1.2.	Parent Seq #:		
Target Value:	N/A			Parent Name:		
_				Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:		
	0	No		Harvested:		
	1	Yes		Format:	Text (Categorical)	
Owner and the D. C. 10				Default Value:	NULL	
Supporting Definitions:	Supporting Definitions: (none)			Usual Range:		
				Valid Range:		
				DataSource:	User	





			D. Encounter Information				
Seq. #: 6280 Name: HF Education - All of the following					Technical Specifications		
<b>3eq. #.</b> 0200 <b>Na</b> ille.	TII Laac	adon Anorthe for	lowing	ShortName:	HFEduAll		
Coding Instructions:	Indicate if the	he patient received all o	of the following education for heart failure.	Parent Seq #:			
Target Value:	Anv occurr	ence between start of c	urrent encounter and completion of current encounter	Parent Name:			
_	-			Parent Value:			
Selections:	Code	Selection Text	Definition	Missing Data:	Report		
	0	No		Harvested:	Yes (PINN)		
	1	Yes		Format:	Text (Categorical)		
Supporting Definitions:	(none)			Default Value:	No		
Supporting Deminions.	(HOHO)			Usual Range:			
				Valid Range:			
				DataSource:	User		
Seq. #: 6281 Name:	HF Educ	ation - Weight Mon	nitoring		Specifications		
		· ·	· ·	ShortName:	HFEduWtMonitoring		
Coding instructions:	indicate if the	ne patient received weig	ght monitoring education for heart failure.	Parent Seq #:	6435		
_	-	ence between start of c	urrent encounter and completion of current encounter	Parent Name:	Seattle Angina Questionnaire (SAQ Completed		
Selections:	Code	Selection Text	Definition	Parent Value:	Yes		
	0	No		Missing Data:	Report		
	1	Yes		Harvested:	Yes (PINN)		
0 4 5 6 4	•			Format:	Text (Categorical)		
Supporting Definitions:	(HOHE)			Default Value:	No		
				Usual Range:			
				Valid Range:			
				DataSource:	User		
Can # 6202 Name	UE Educ	ation - Diet (Sodiur	n Restriction)	Technical S	pecifications		
<b>Seq.</b> #: 6282 Name:	HF Educ	ation - Diet (Sodiul	II Restriction)	ShortName:	HFEduDiet		
Coding Instructions:	Indicate if the	he patient received a so	dium-restricted dietary education for heart failure.	Parent Seq #:	6435		
Target Value:	Any occurr	ence between start of c	urrent encounter and completion of current encounter	Parent Name:	Seattle Angina Questionnaire (SAC Completed		
Selections:	Code	Selection Text	Definition	Parent Value:	Yes		
	0	No		Missing Data:	Report		
	1	Yes		Harvested:	Yes (PINN)		
	•			Format:	Text (Categorical)		
Supporting Definitions:	(none)			Default Value:	No		
				Usual Range:			
				Valid Range:			
				DataSource:	User		





			D. Encounter Informatio				
O #- C202 Nome-	eq. #: 6283 Name: HF Education - Symptom Management				Technical Specifications		
Seq. #: 6283 Name:	HE Education - Symptom Management			ShortNar	ne: HFEduSympMgmt		
Coding Instructions:	Indicate if the	he patient received symp	otom management education for	or heart failure. Parent Sec	<b>#:</b> 6435		
_	•		urrent encounter and completion	n of current encounter	ne: Seattle Angina Questionnaire (SAQ Completed		
Selections:	Code	Selection Text	Definition	Parent Valu	ıe: Yes		
	0	No		Missing Da	a: Report		
	1	Yes		Harveste	ed: Yes (PINN)		
Supporting Definitions:	(none)			Forma	at: Text (Categorical)		
Supporting Deminions.	(HOHC)			Default Valu	e: No		
				Usual Rang	e:		
				Valid Rang	e:		
				DataSource	e: User		
Seq. #: 6284 Name:	HE Educ	ation - Physical Act	ivity	<u>Technic</u>	cal Specifications		
•					ne: HFEduPhyAct		
Coding Instructions:	Indicate if the	he patient received phys	sical activity education for heart	failure. Parent Sec			
_	-	ence between start of cu	urrent encounter and completion	n of current encounter	ne: Seattle Angina Questionnaire (SAQ Completed		
Selections:	Code	Selection Text	Definition	Parent Valu	ie: Yes		
	0	No		Missing Da	ta: Report		
	1	Yes		Harveste	ed: Yes (PINN)		
Owner and to a Deffect them a	(nono)			Form	at: Text (Categorical)		
Supporting Definitions:	(Horie)			Default Valu	e: No		
				Usual Rang	e:		
				Valid Rang	e:		
				DataSourc	e: User		
Seq. #: 6285 Name:	HE Educ	eation - Smoking Co	esation	<u>Technic</u>	al Specifications		
<b>Seq.</b> #: 0200 Name.	III Luuc	ation - Smoking Ce	ssauori	ShortNar	ne: HFEduSmokeCess		
Coding Instructions:	Indicate if the	he patient received smol	king cessation education for he	art failure. Parent Sec	<b>#:</b> 6435		
_	•	ence between start of cu	urrent encounter and completion	n of current encounter	ne: Seattle Angina Questionnaire (SAQ Completed		
Selections:	Code	Selection Text	Definition	Parent Valu	•		
	0	No		Missing Da			
	1	Yes		Harveste			
				Form	at: Text (Categorical)		
Supporting Definitions:	(none)			Default Valu	e: No		
				Usual Rang	e:		
				Valid Rang	ie:		
				Valia italig	17.		





Soa #1 6296 Name:	eq. #: 6286 Name: HF Education - Medication Instruction				Technical S	Technical Specifications		
·						HFEduMedInstr		
Coding Instructions:	Indicate if the	he patient received med	lication instruction	n education for heart failure.	Parent Seq #:	6435		
_	-			and completion of current encounter	Parent Name:	Seattle Angina Questionnaire (SAQ) Completed		
Selections:	Code	Selection Text	Definition		Parent Value:	Yes		
	0	No			Missing Data:	Report		
	1	Yes			Harvested:	Yes (PINN)		
Supporting Definitions	(none)				Format:	Text (Categorical)		
Supporting Definitions:	(HOHE)				Default Value:	No		
					Usual Range:			
					Valid Range:			
					DataSource:	User		
Seq. #: 6287 Name:	HF Educ	ation - Prognosis/E	nd-of-l ife lss	2011	Technical S	<u>specifications</u>		
		-			ShortName:	HFEduPrognosis		
Coding Instructions:	Indicate if the	he patient received prog	nosis/end-of-life	issues education for heart failure.	Parent Seq #:	6435		
_	-			and completion of current encounter	Parent Name:	Seattle Angina Questionnaire (SAQ) Completed		
Selections:	Code	Selection Text	Definition		Parent Value:	Yes		
	0	No			Missing Data:	Report		
	1	Yes			Harvested:	Yes (PINN)		
Summerting Definitions	(none)				Format:	Text (Categorical)		
Supporting Definitions:	(Horie)				Default Value:	No		
					Usual Range:			
					Valid Range:			
					DataSource:	User		
Seq. #: 6288 Name:	HE Educ	ation - Minimizing o	or Avoiding us	e of NSAIDs	Technical S	pecifications		
•					ShortName:	HFEduNSAIDs		
Coding Instructions:	Indicate if the failure.	he patient received mini	mizing or avoidin	g use of NSAIDs education for heart	Parent Seq #:	6435		
					Parent Name:	Seattle Angina Questionnaire (SAQ)		
Target Value:	Any occurre	ence between start of co	urrent encounter	and completion of current encounter		Completed		
Selections:	Code	Selection Text	Definition		Parent Value:	Yes		
					Missing Data:	Report		
	0	No			Harvested:	Yes (PINN)		
	1	Yes			Format:	Text (Categorical)		
Supporting Definitions:	(none)				Default Value:	No		
					Usual Range:			
					Valid Range:			
					DataSource:	User		





#### **D. Encounter Information**

Seq. #: 6289 Name: HF Education - Referral for visiting nurse or specific

education or management programs

Coding Instructions: Indicate if the patient received a referral for visiting nurse or specific education or

management programs education for heart failure.

Target Value: Any occurrence between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: (none)

Seq. #: 6300 Name: ICD Counseling

Coding Instructions: Indicate if patient has been counseled regarding Implantable Cardioverter Defibrillator

Implantation(ICD).

Note(s):

Code 'Yes' for single chamber ICD, dual chamber ICD, cardiac resynchronization

therapy device and defibrillator (CRT-D).

Target Value: Any occurrence between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition

Counseled

1 Yes - Patient Counseled
2 No - Patient Not

3 No Counseling - Medical Reason

Supporting Definitions: (none)

Seq. #: 6310 Name: HF Plan of Care

Coding Instructions: Indicate if the patient has a documented plan of care for management of heart failure

symptoms.

Target Value: Any occurrence between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition

0 No

1 Yes Plan of care was documented

Supporting Definitions: HF Plan of Care:

A documented plan of care may include one or more of the following: reevaluation of medical therapy including up-titration of doses, consideration of electrical device therapy, recommended lifestyle modifications, initiation of palliative care, referral for more advanced therapies (e.g. transplant, ventricular assist device), or referral to disease

management programs.

Source: 2012 ACCF/AHA/AMA-PCPI Heart Failure Performance Measures

Technical Specifications

ShortName: HFEduPgms

Parent Seq #: 6435

Parent Name: Seattle Angina

Questionnaire (SAQ)

Completed

Parent Value: Yes

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

DataSource: User

Technical Specifications

ShortName: Counsel\_ICD

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

**Technical Specifications** 

HF\_PlanCare

Report

NULL

User

Yes (PINN)

Text (Categorical)

Default Value: NULL

Usual Range:

Valid Range:

ShortName:

Parent Seq #:

**Parent Name:** 

Parent Value: Missing Data:

Harvested:

**Default Value:** 

**Usual Range:** 

Valid Range:

DataSource:

Format:

DataSource: User



#### **D. Encounter Information**

Seq. #: 6400 Name: Left Ventricular Ejection Fraction (LVEF) Date

Coding Instructions: Indicate the date of the most recent left ventricular ejection fraction.

Target Value: The last value between birth and completion of current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: LVEF\_Date

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

**Harvested:** Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User

Seq. #: 6410 Name: Left Ventricular Ejection Fraction (LVEF) Percent

Coding Instructions: Indicate the patient's left ventricular quantitative assessment.

Note(s):

The "LVEF percent" element should only be used if a single percentage is documented

in the medical record.

If a LVEF range or a descriptive term (e.g. Moderately reduced) is documented in the medical record, then report the LV function using the "LV Qualitative Assessment"

element.

Target Value: The last value between birth and completion of current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: LVEF

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR,PINN)

NULL

Format: Integer (2)

Default Value: Usual Range:

Valid Range: 1-99

DataSource: User

#### D. Encounter Information

Selection Retired (v1.3)

Seq. #: 6420 Name: Left Ventricular Qualitative Assessment

Coding Instructions: Indicate the patient's LV Qualitative Assessment.

#### Note(s):

4

If a percentage is documented in the medical record, use the "LVEF Percent" element to document the percentage.

If a LVEF percentage range is documented in the medical record, average the percentages, round up and reference the "LV Qualitative Assessment" selections to report.

Target Value: The last value between birth and completion of current encounter

Selections: Code Selection Text Definition

1 Normal: >=50 Selection Retired (v1.3)

2 Mildly reduced: 40 - 49 3 Moderately reduced: 26 - 39

Severely reduced: Selection Retired (v1.3)

5 Hyperdynamic: >706 Normal: 50 - 70

<=25

7 Moderately reduced: 30
- 39
8 Severely reduced:

<=29

Supporting Definitions: (none)

Seq. #: 6430 Name: Canadian Cardiovascular Society (CCS) Class

Coding Instructions: Indicate the patient's Canadian Cardiovascular Society (CCS) classification for angina.

Target Value: The value on current encounter

Selections: Code Selection Text Definition No angina 0 Ordinary physical activity (for example, walking or climbing stairs) does not cause angina; angina occurs with strenuous or rapid or prolonged exertion at work or recreation, Slight limitation of ordinary activity (for example, Ш angina occurs walking or stair climbing after meals, in cold, in wind, under emotional stress, or only during the few hours after awakening; walking more than 2 blocks on the level or climbing more than Ш Marked limitation of ordinary activity (for example, angina occurs with walking 1 or 2 blocks on the

level or climbing 1 flight of stairs in normal conditions and at a normal pace).

4 IV Inability to perform any physical activity without discomfort; angina syndrome may be present at

rest.

Supporting Definitions: (none)

Technical Specifications					
ShortName:	LV_Qlty_Assemnt				
Parent Seq #: Parent Name:					
Parent Value:					
Missing Data:	Report				
Harvested:	Yes (DCR,PINN)				
Format:	Text (Categorical)				
Default Value:	NULL				
Usual Range:					
Valid Range:					
DataSource:	User				

**Technical Specifications** 

Report

NULL

User

Yes (DCR,PINN)

Text (Categorical)

ShortName: CCSClass

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

Default Value:

**Usual Range:** 

Valid Range:

DataSource:

Harvested:

Format:





Seq. #: 6435 Name: Seattle Angina Questionnaire (SAQ) Completed					Technical Specifications	
Seq. #: 6435 Name:	Seattle P	angina Questionnai	ire (SAQ) Completed	ShortName:	SAQCompleted	
Coding Instructions:	Indicate if the	he patient has complete	ed the Seattle Angina Questionnaire (SAQ).	Parent Seq #:		
Target Value:	Any occurr	ence between start of c	current encounter and completion of current encounter	Parent Name:		
· ·	•		·	Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:	Report	
	0	No		Harvested:	Yes (PINN)	
	1	Yes		Format:	Text (Categorical)	
Supporting Definitions:	(none)			Default Value:	No	
Supporting Deminions.	(HOHO)			Usual Range:		
				Valid Range:		
				DataSource:	User	
Com # 6440 Nome	Other To	r Tool/Method used to assess Angina Symptoms and		Technical S	pecifications	
Seq. #: 6440 Name:		Completed	assess Angina Cymptoms and	ShortName:	OtherAnginaToolCo mpleted	
Coding Instructions:		another tool/method was er than the CCS or SAC	s used to assess the patient's angina symptoms and $\ensuremath{\mathtt{Q}}.$	Parent Seq #: Parent Name:		
Target Value:	Any occurr	ence between start of c	surrent encounter and completion of current encounter	Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:	Report	
Coloculono		Selection Text	Delinition	Harvested:	Yes (PINN)	
	0	No		Format:	Text (Categorical)	
	1	Yes		Default Value:	No	
Supporting Definitions: (none)				Usual Range:		
	/			Valid Range:		
				DataSource:	User	

#### D. Encounter Information

Seq. #: 6450 Name: Cardiac Rehabilitation Referral or Plan for Qualifying

Event/Diagnosis

**Coding Instructions:** Indicate if the patient had a cardiac event within the past 12 months requiring cardiac rehabilitation. Cardiac events includes Myocardial Infarction, Valve Replacement, Heart

Transplant, CABG or PCI.

Note(s):

Cardiac rehabilitation is a medically supervised program to help cardiac patients slow and stabilize the progression of cardiovascular disease thus reducing the risk of heart disease, another cardiac event or death. Cardiac rehabilitation programs include patient counseling, an exercise program, nutrition counseling and risk factor education (smoking, obesity, high blood pressure, high cholesterol).

Target Value: Any occurrence between start of current encounter and completion of current encounter

Selections: Code Selection Text Definition

System Reason

1	Yes - Referral/Plan documented	
2	No qualifying event/diagnosis	
3	Patient already participating in rehab	
4	No Referral/Plan - Medical Reason	
5	No Referral/Plan - Patient Reason	Selection Retired (v1.3)
6	No Referral/Plan -	

#### Supporting Definitions: Referral:

A referral is defined as an official communication between the health care provider and the patient to recommend and carry out a referral order to an outpatient CR program. This includes the provision of all necessary information to the patient that will allow the patient to enroll in an outpatient CR program. This also includes a written or electronic communication between the healthcare provider or healthcare system and the cardiac rehabilitation program that includes the patient's enrollment information for the program. A hospital discharge summary or office note may potentially be formatted to include the necessary patient information to communicate to the CR program [the patient's cardiovascular history, testing, and treatments, for instance]. According to standards of practice for cardiac rehabilitation programs, care coordination communications are sent to the referring provider, including any issues regarding treatment changes, adverse treatment responses, or new non-emergency condition (new symptoms, patient care questions, etc.) that need attention by the referring provider. These communications also include a progress report once the patient has completed the program. All communications must maintain an appropriate level of confidentiality as outlined by the 1996 Health Insurance Portability and Accountability Act [HIPAA].)

Source: Thomas RJ, King M,Lui K, et al. "AACVPR/ACC/AHA 2007 Performance Measures on Cardiac Rehabilitation for Referral to and Delivery of Cardiac Rehabilitation/Secondary Prevention Services." Journal of American College of Cardiology. 2007: 50(14), pp 1400-1433

Technical Specifications					
ShortName:	CardRehabReferral				
Parent Seq #: Parent Name:					
Parent Value:					
Missing Data:	Report				
Harvested:	Yes (DCR,PINN)				
Format:	Text (Categorical)				
Default Value:	NULL				
Usual Range:					
Valid Range:					
DataSource:	User				





O #- 0400 N	Poterral for consideration for coronary reveasularization	Technical S	Specifications		
<b>Seq.</b> #: 6460 Name:	Referral for consideration for coronary revascularization	ShortName:	CorRevasReferral		
Coding Instructions:	<b>Coding Instructions:</b> Indicate if the patient has a documented referral for consideration for coronary revascularization.				
	Tevascularization.	Parent Name:			
Target Value:	Any occurrence between start of current encounter and completion of current encounter	Parent Value:			
Selections:	Code Selection Text Definition	Missing Data:	Report		
		Harvested:	Yes (DCR,PINN)		
	0 No	Format:	Text (Categorical)		
	1 Yes	Default Value:	NULL		
Supporting Definitions:	(none)	Usual Range:			
		Valid Range:			
		DataSource:	User		
Seg #: 6470 Name:	Referral for additional evaluation/treatment of anginal	Technical S	Specifications		
00q 0 0	symptoms	ShortName:	EvalTreatReferral		
Coding Instructions:	Indicate if the patient has a documented referral for additional evaluation/treatment of anginal symptoms.	Parent Seq #: Parent Name:			
		Parent Value:			
Target Value:	Any occurrence between start of current encounter and completion of current encounter	Missing Data:	Report		
Selections:	Code Selection Text Definition	Harvested:	Yes (DCR,PINN)		
	N.	Format:	Text (Categorical)		
	0 No	Default Value:	NULL		
	1 Yes	Usual Range:			
Supporting Definitions:	(none)	Valid Range:			
		DataSource:	User		
Seq. #: 6481 Name:	Seattle Angina Questionnaire (SAQ) - Physical Function	<u>Technical S</u>	Specifications		
•	Score	ShortName:	SAQAnginaPhyFunc Score		
Coding Instructions:	This element has been retired effective PINNACLE v1.3.	Parent Seq #:	6105		
Target Value:	N/A	Parent Name:	Patient enrolled in weight loss program		
Selections:	(none)	Parent Value:	Yes		
Supporting Definitions:	(none)	Missing Data:	No Action		
Supporting Deminions:	(none)	Harvested:	Yes (PINN)		
		Format:	Integer (3)		
		Default Value:	NULL		
		Usual Range:			
		Valid Range:	0-100		
		DataSource:	User		



#### D. Encounter Information

**Technical Specifications** Name: Seattle Angina Questionnaire (SAQ) - Angina Stability **Seq. #:** 6482 SAQAnginaStability ShortName: Score Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6105 **Parent Name:** Patient enrolled in Target Value: N/A weight loss program Parent Value: Yes Selections: (none) Missing Data: No Action Supporting Definitions: (none) Harvested: Yes (PINN) Format: Integer (3) Default Value: NULL **Usual Range:** Valid Range: 0-100 DataSource: User **Technical Specifications** Name: Seattle Angina Questionnaire (SAQ) - Angina Frequency **Seq. #:** 6483 ShortName: SAQAnginaFreqSco Score Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6105 **Parent Name:** Patient enrolled in Target Value: N/A weight loss program Parent Value: Selections: (none) Missing Data: No Action Supporting Definitions: (none) Yes (PINN) Harvested: Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: 0-100 DataSource: User **Technical Specifications** Name: Seattle Angina Questionnaire (SAQ) - Treatment **Seq. #:** 6484 SAQAnginaTreatme ShortName: Satisfaction Score ntsatiScore Coding Instructions: This element has been retired effective PINNACLE v1.3. Parent Seq #: 6105 **Parent Name:** Patient enrolled in Target Value: N/A weight loss program Parent Value: Yes Selections: (none) Missing Data: No Action Supporting Definitions: (none) Harvested: Yes (PINN) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: 0-100 DataSource: User





Con #1 GAGE Name:	Spattle Ar	eattle Angina Questionnaire (SAQ) - Quality of Life			Technical Specifications		
Seq. #: 0405 Name:	Score	ngina Questionnair	e (3AQ) - Q	uality of Life		ShortName:	SAQAnginaQuallifeS core
Coding Instructions:	This element	t has been retired effect	tive PINNACLE	v1.3.		Parent Seq #:	6105
Target Value:	N/A					Parent Name:	Patient enrolled in weight loss program
Selections:	: (none)					Parent Value:	Yes
	, ,					Missing Data:	No Action
Supporting Definitions:	(none)					Harvested:	Yes (PINN)
						Format:	Integer (3)
						Default Value:	NULL
						Usual Range:	
						Valid Range:	0-100
						DataSource:	User
Seq. #: 6490 Name:	Hypertens	sion Plan of Care Γ	Occumented			Technical S	pecifications
<b>3eq.</b> #: 0490 Name.	riyperterio	Sion i lan oi care L	ocumented			ShortName:	HTPlanofCare
-		t has been retired effect	tive PINNACLE	E v1.3.		Parent Seq #: Parent Name:	
Target Value:	N/A					Parent Value:	
Selections:	Code	Selection Text	Definition			Missing Data:	Report
		No				Harvested:	Yes (PINN)
	0	Yes				Format:	Text (Categorical)
	1	165				Default Value:	NULL
Supporting Definitions:	(none)					Usual Range:	
						Valid Range:	
						DataSource:	User
Com #1 6500 Namo	ΛΕίΚ/ΕΙμ <del>ι</del>	ter Duration				Technical S	pecifications
<b>Seq.</b> #: 6500 Name:	, Al Ib/I lutt	lei Duiation				ShortName:	Afib_Dur
Coding Instructions:	Indicate the	duration of the patient's	AFib/Flutter.			Parent Seq #:	
Target Value:	The value or	n current encounter				Parent Name:	
Selections:	Code	Selection Text	Definition			Parent Value:	Danad
		Coloculor Toxi	Bonnaon			Missing Data:	Report
	1	First diagnosed				Harvested:	Yes (PINN)
	2	Paroxysmal				Format:	Text (Categorical)
	3	Persistent				Default Value: Usual Range:	NULL
	4	Long-standing Persistent				Valid Range:	
	5	Permanent				DataSource:	User
Supporting Definitions:					'		





Atrial Fibrillation

# NCDR® Outpatient Registries v1.4 Data Dictionary - Full Specifications

### D. Encounter Information

0 " 0540 Name	Λ Cib/Clutto	A File /Flootheau Toure		pecifications					
Seq. #: 6510 Name:	eq. #: 6510 Name: AFib/Flutter Type								
Coding Instructions:	Indicate the if	the patient has valvular of non-valvular AFib/Flutter	Parent Seq #:						
Target Value:	The value on	current encounter	Parent Name:						
<b>U</b>		current encounter	Parent Value:						
Selections:	Code	Selection Text Definition	Missing Data:	Report					
	1	Non - valvular	Harvested:	Yes (PINN)					
	2	Valvular	Format:	Text (Categorical)					
	_		Default Value:	NULL					
Supporting Definitions:	AFib/Flutter	Type:	Usual Range:						
	Valid Range:								
	vaive, Of filst	ory of mitral valve repair.	DataSource:	User					

Source: 2014 AHA/ACC/HRS Guideline for the Management of Patients With

Seq. #: 6520 Name: Etiology - Transient/reversible Cause				Technical S	Specifications
•			due to a transient and/or reversible cause.	ShortName:	Afib_Etiology_rev_c ause
		n current encounter		Parent Seq #: Parent Name:	
Selections:	Code	Selection Text	Definition	Parent Value:	
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
				Usual Range:	
				Valid Range:	
				DataSource:	User
Can # CEO1 Name	Etiology	Cardiac Surgery with	nin nast 2 manths	Technical S	specifications
	-	nt has been retired effective	·	ShortName:	Afib_Etiology_Card_ Srg
Target Value:	N/A			Parent Seq #: Parent Name:	
Selections:	Code	Selection Text	Definition	Parent Value:	
				Missing Data:	Report
	0	No		Harvested:	Yes (PINN)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:	(none)			Default Value:	No
				Usual Range:	
				Valid Range:	





Seq. #: 6522 Name:	-tiology - Pregnancy	Technical S	pecifications
•	This element has been retired effective PINNACLE v1.3.	ShortName:	Afib_Etiology_Pregancy
Target Value:		Parent Seq #: Parent Name:	
Selections:	Code Selection Text Definition	Parent Value:	
		Missing Data:	Report
	0 No	Harvested:	Yes (PINN)
	1 Yes	Format:	Text (Categorical)
Supporting Definitions:	none)	Default Value:	No
		Usual Range:	
		Valid Range:	
		DataSource:	User
Seg. #: 6530 Name:	nternational Normalized Ratio (INR) Value		Specifications
•		ShortName:	INR_Value
Coding Instructions:	ndicate all values of the patient's International Normalized Ratio (INR).  Note(s):	Parent Seq #: Parent Name:	
	All occurrences between birth and completion of current encounter	Parent Value:	
Torret Value	·	Missing Data:	Report
Target Value:		Harvested:	Yes (PINN)
Selections:	(none)	Format:	Decimal (3,1)
Supporting Definitions:	none)	Default Value:	NULL
		Usual Range:	
		Valid Range:	0.1-99.0
		DataSource:	User
Con #. 6522 Names	nternational Normalized Ratio (INR) Date	Technical S	pecifications
Seq. #: 0002 Name.	The mational Normalized Natio (INN) Date	ShortName:	INR_Dt
Coding Instructions:	ndicate all dates the patient's International Normalized Ratio (INR) was assessed.	Parent Seq #:	6530
	Note(s):	Parent Name:	International Normalized Ratio
	All occurrences between birth and completion of current encounter		(INR) Value
Target Value:	N/A	Parent Value:	Not Null
Selections:		Missing Data:	No Action
Jeiecuolis.	(none)	Harvested:	Yes (PINN)
Supporting Definitions:	none)	Format:	Date (mm/dd/yyyy)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User





#### D. Encounter Information

Seq. #: 6540 Name: Electrophysiology Study

Coding Instructions: Indicate if the patient received an electrophysiology study (EP study).

Note(s):

An EP study consists of one or more catheters capable of recording and pacing which are placed in one or more of the cardiac chambers. The catheters may be used to measure conduction of the impulse from the sinus node to the ventricle; induce a tachycardia; and/or localize (map) the location where the tachycardia originates.

Target Value: Any occurrence between birth and completion of current encounter

Target Value: Any occurrence between birth and completion of current encounter

Selection Text

No

Yes

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: (none)

Technical Specifications

ShortName: EPStudy

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: No

Usual Range:

Valid Range:

**Parent Value:** 

Missing Data:

**Default Value:** 

**Usual Range:** 

Valid Range: DataSource:

Harvested:

Format:

Report

User

Yes (PINN)

Text (Categorical)

DataSource: User

Con # 6E40 Name	: Electrophysiology Study Date	Technical S	pecifications
<b>Seq.</b> #: 6542 <b>Name</b> :	Electrophysiology Study Date	ShortName:	EPStudy_Date
Coding Instructions:	Indicate all dates the patient received an electrophysiology study.	Parent Seq #:	6540
	Note(s):	Parent Name:	Electrophysiology Study
	All occurrences between birth and completion of current encounter	Parent Value:	Yes
Target Value:	N/A	Missing Data:	No Action
Selections:	(none)	Harvested:	Yes (PINN)
• 4 <b>•</b> • • •	(1000)	Format:	Date (mm/dd/yyyy)
Supporting Definitions:	(none)	Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User
Con # 6550 Namo	Atrial Ablation	Technical S	pecifications
<b>Seq.</b> #: 6550 <b>Name</b> :	Attial Abiation	ShortName:	AtrialAblation
Coding Instructions:	Indicate if an atrial ablation was performed. Ablation is the application of an energy source delivered through a catheter to eliminate or modify a focus or re-entry circuit that causes an arrhythmia.	Parent Seq #: Parent Name:	

Definition

Selections: Code

Supporting Definitions: (none)

0





O # 0550 Names	Atrial Ablation Data	Technical Specifications		
Seq. #: 6552 Name:	Allial Abiation Date	ShortName:	AtrialAblation_Date	
Coding Instructions:	Indicate all dates the patient received an atrial ablation.	Parent Seq #:	6550	
	Note(s):	Parent Name:	Atrial Ablation	
	All occurrences between birth and completion of current encounter	Parent Value:	Yes	
Target Value:	N/A	Missing Data:	No Action	
_		Harvested:	Yes (PINN)	
Selections:	(none)	Format:	Date (mm/dd/yyyy)	
Supporting Definitions:	(none)	Default Value:	NULL	
		Usual Range:		
		Valid Range:		
		DataSource:	User	
Seq. #: 6560 Name:	Atrial Fibrillation Recurrence		Specifications	
		ShortName:	AFRecurrence	
Coding Instructions:	Indicate if the patient had a documented case of atrial fibrillation of any type after the performance of an atrial fibrillation ablation.	Parent Seq #: Parent Name:		
Target Value:	Any occurrence between birth and current encounter	Parent Value:		
Selections:	Code Selection Text Definition	Missing Data:	Report	
	Sold Sold Fox	Harvested:	Yes (PINN)	
	0 No	Format:	Text (Categorical)	
	1 Yes	Default Value:	No	
Supporting Definitions:	(none)	Usual Range:		
		Valid Range:		
		DataSource:	User	
Sog #: 6562 Name:	Atrial Fibrillation Recurrence Date	Technical S	<u>Specifications</u>	
•		ShortName:	AFRecurrence_Dat	
Coding Instructions:	Indicate all dates the patient had an atrial fibrillation recurrence.	Parent Seq #:	6560	
	Note(s):	Parent Name:	Atrial Fibrillation Recurrence	
	All occurrences between birth and completion of current encounter	Parent Value:		
Target Value:	N/A	Missing Data:	No Action	
Selections:	(none)	Harvested:	Yes (PINN)	
Gelections.	(none)	Format:	Date (mm/dd/yyyy)	
Supporting Definitions:	(none)	Default Value:	NULL	
		Usual Range:		
			Hear	
		Valid Range: DataSource:	User	





#### D. Encounter Information

0 // 0570 Name	Atrial Fib	rillation Cumptom Fron	NUODO!	Technical S	pecifications
<b>Seq. #:</b> 6570 <b>Name</b> :				ShortName:	AFSymptom_Freque
Coding Instructions:	Indicate the of atrial fibri		e interval, in days, between symptomatic episodes	Parent Seg #:	ncy
Toward Wolco	<b>^</b>	and the form of the first of the second of t		Parent Name:	
_	-	ence between birth and curr	ent encounter	Parent Value:	
Selections:	(none)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (PINN)
				Format:	Integer (5)
				Default Value:	No
				Usual Range:	
				Valid Range:	1-99999
				DataSource:	User
Seq. #: 6580 Name:	Atrial Fibr	rillation Symptom Dura	ation	Technical S	pecifications
4		• •		ShortName:	AFSymptom_Duratio
Coding Instructions:	Indicate the fibrillation.	patient estimate of duration	n of usual symptomatic episodes for atrial	D	11
				Parent Seq #: Parent Name:	
Target Value:	Any occurre	ence between birth and curr	rent encounter	Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
				-	Yes (PINN)
	1	< 48 hours		Format:	Text (Categorical)
	2	>= 48 hours to 7 days		Default Value:	NULL
	3	> 7 days to 3 months		Usual Range:	
	4	> 3 months		Valid Range:	
Supporting Definitions:	(none)			DataSource:	User
				Technical S	Specifications

Seq. #: 6590 Name: Rate Control (Therapy)

**Coding Instructions:** Indicate if the patient is currently on rate control therapy.

Rate control is the attempted control of ventricular rate with no commitment to restore or maintain sinus rhythm. (Strict rate control is generally defined as <80 bpm while lenient rate control is generally defined as <110 bpm.) Rate control may consist of:

- Pharmacological

- Non pharmacological
- Hybrid

Target Value: The value on current encounter

Selections: Code Selection Text Definition

0 No 1 Yes

Supporting Definitions: (none)

Technical Specifications

ShortName: RateControl

Parent Seq #:

Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: N

Usual Range:

Valid Range:

DataSource: User



#### **D. Encounter Information**

Seq. #: 6595 Name: Rhythm Control (Therapy)

**Coding Instructions:** Indicate if the patient is currently on rhythm control therapy.

Rhythm control is the attempted restoration and/or maintenance of sinus rhythm. Also

requires attention to rate control. Rhythm control may consist of:

Target Value: Any occurrence between start of current encounter and completion of current encounter

Definition

Selection Retired (v1.3)

Selection Retired (v1.2)

- Pharmacological

- Non pharmacological

- Hybrid

Target Value: The value on current encounter

Selections: Code Selection Text Definition

Name: Thromboembolic Risk Factors Assessed

Selection Text

assessed)

Yes (All risk factors

No - Medical Reason

No - Patient Reason

No - System Reason

0 No

Yes

Coding Instructions: This element has been retired effective v1.4

Supporting Definitions: (none)

**Seq.** #: 6596

Technical Specifications

User

**Technical Specifications** 

RhvthmControl

Report

Yes (PINN)

Text (Categorical)

ShortName:

Parent Seq #: Parent Name:

Parent Value:

Missing Data:

**Default Value:** 

Usual Range: Valid Range:

DataSource:

Harvested:

Format:

ShortName: ThrombRskFact

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Text (Categorical)

Default Value: Usual Range:

Valid Range:

DataSource: User

Seg. #: 6600 Name: CHA2DS2 Score

Selections: Code

1

2

3

4

Coding Instructions: Indicate the value of the patient's CHA2DS2 Score.

Target Value: The value between birth and current encounter

Selections: (none)

Supporting Definitions: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: CHA2DS2Score

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (PINN)

Format: Integer (1)

Default Value: NULL

**Usual Range:** 

Valid Range: 0-6

DataSource: User





Co. # 0040 No	CHVDea	JADC2 VACa Caara		Technical Specifications	
<b>Seq.</b> #: 6610 <b>Name</b> :	CHADS2-	VASC Score		ShortName:	CHA2DS2VScore
Coding Instructions:	Indicate the	value of the patient's CHAD	OS2-VASc Score.	Parent Seq #:	
Target Value	The value he	etween birth and current en	counter	Parent Name:	
		or our our and our one on	ocurre.	Parent Value:	
Selections:	(none)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (PINN)
				Format:	Integer (1)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	0-9
				DataSource:	User
Seq. #: 6620 Name:	HAS-BLE	D Score		Technical S	specifications
•				ShortName:	HASBLEDScore
Coding Instructions:	Indicate the	value of the patient's HAS-l	BLED Score.	Parent Seq #:	
Target Value:	The value be	etween birth and current en	counter	Parent Name:	
Selections:				Parent Value:	
Selections.	(Horie)			Missing Data:	Report
Supporting Definitions:	(none)			Harvested:	Yes (PINN)
				Format:	Integer (1)
				Default Value:	NULL
				Usual Range:	
				Valid Range:	0-9
				DataSource:	User
Seq. #: 6630 Name:	Foot Exan	n (Within the Past 12	Months)	<u>Technical S</u>	specifications
•				ShortName:	FootExam
_			within the past 12 months.	Parent Seq #: Parent Name:	
larget value:	encounter	nce between 12 month prio	or to current encounter and completion of current	Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
				Harvested:	Yes (DCR)
	document through vi monofilan 1 Yes A foot exa	No documentation of a foot exam or the	Format:	Text (Categorical)	
			documentation does not include examination through visual inspection, sensory exam with	Default Value:	Null
		monofilament, and pulse exam.	Usual Range:		
		A foot exam should include these 3 elements: visual inspection, sensory exam with	Valid Range:		
			monofilament AND pulse exam.	DataSource:	User
Supporting Definitions:	(none)				





Con # 6600 Nome	eq. #: 6632 Name: Foot Exam Date				
Seq. #: 6632 Name:	FUUL EXAI	II Date		ShortName:	FootExam_Date
Coding Instructions:	Indicate the	date the patient received a	a foot exam.	Parent Seq #:	6630
Target Value:	N/A			Parent Name:	Foot Exam (Within the Past 12 Months)
Selections:	(none)			Parent Value:	Yes
	()			Missing Data:	No Action
Supporting Definitions:	(none)			Harvested:	Yes (DCR)
				Format:	Date (mm/dd/yyyy)
				Default Value:	NULL
				Valid Range:	
				DataSource:	User
Seq. #: 6640 Name:	Monofilan	nent Exam		Technical S	Specifications
•				ShortName:	MonofilExam
Coding Instructions:	Indicate if th	e patient received a mono	filiment exam within the past 12 months.	Parent Seq #:	
Target Value:	N/A			Parent Name:	
_			D # W	Parent Value:	
Selections:	Code	Selection Text	Definition	Missing Data:	Report
	0	No		Harvested:	Yes (DCR)
	1	Yes		Format:	Text (Categorical)
Cunnarting Definitions	(none)			Default Value:	Null
Supporting Definitions:	(Horie)			Usual Range:	
				Valid Range:	
				DataSource:	User
Seq. #: 6650 Name:	Pulse Exa	am		Technical Specifications	
•				ShortName:	PulseExam
Coding Instructions:	Indicate if th	e patient received a pulse	exam within the past 12 months.	Parent Seq #:	
Target Value:	N/A			Parent Name:	
Selections:		Oalestian Tour	Definition	Parent Value:	
Selections.	Code	Selection Text	Definition	Missing Data:	Report
	0	No		Harvested:	Yes (DCR)
	1	Yes		Format:	Text (Categorical)
Supporting Definitions:				Default Value:	Null
Supporting Deminions:	(HOHO)			Usual Range:	
				Valid Range:	
				DataSource:	User





On the CCCO Names Apkle Prophiel Index Test				Technical	Specifications	
Seq. #: 6660 Name: Ankle Brachial Index Test					ABI_Performed	
Coding Instructions:	Indicate if the	e patient received an a	nkle brachial index test within the past 12 months.	Parent Seq #: Parent Name:		
Target Value:	N/A			Parent Value:		
Selections:	Code	Selection Text	Definition	Missing Data:	Report	
	0	No		Harvested:	Yes (DCR)	
	1	Yes		Format:	Text (Categorical)	
		100		Default Value:	Null	
Supporting Definitions:	(none)			Usual Range:		
				Valid Range:		
				DataSource:	User	
<b>Seq.</b> #: 6670 <b>Name</b> :	Negative (	dilated or retinal ey	/e eyam	<u>Technical</u>	Technical Specifications	
Seq. #. 0070 Name.	ricgative	anated of retirial ey	re exam	ShortName:	NegRetDiaExam	
Coding Instructions:			pative retinal or dilated eye exam (negative for ional (optometrist or ophthalmologist) within the pa	Parent Seq #:		
	24 months.					
<b>-</b>	•			Parent Value:		
larget Value:	Any occurrer encounter	nce between 24 months	s prior to current encounter and completion of curr	Missing Data:	Report	
Selections:	Code	Selection Text	Definition	Harvested:	Yes (DCR)	
				Format:	Text (Categorical)	
	0	No - Not documented		Default Value:	Null	
	1	Yes		Usual Range:		
Supporting Definitions:	(none)			Valid Range:		
7. 0				DataSource:	User	





#### D. Encounter Information

**Technical Specifications** Name: Retinal or Dialated Eye Exam Seq. #: 6680 ShortName: RetDiaExam Coding Instructions: Indicate if the patient has had an eye exam with an eye care provider within the past 12 Parent Seq #: **Parent Name:** Parent Value: Target Value: Any occurrence between 12 month prior to current encounter and completion of current encounter Missing Data: Report Selections: Code Selection Text Definition Harvested: Yes (DCR) Text (Categorical) Format: No - Not documented 0 Indicate if there was no documentation of a retinal or dilated eye exam by an Eye Care Professional **Default Value:** or the documentation did not include any of the **Usual Range:** 1) Retinal or dilated eye exam interpretation by an Valid Range: ophthalmologist or optometrist was documented and reviewed. DataSource: User 2) Seven standard field stereoscopic photos with interpretation by an ophthalmologist or optometrist documented and reviewed. 3) Eye imaging validated to match diagnosis from seven standard field stereoscopic photos results documented and reviewed.

> Indicate if the Retinal or Dilated Eye Exam was Performed by an Eye Care Professional. This

1) Retinal or dilated eye exam interpretation by an ophthalmologist or optometrist was documented

2) Seven standard field stereoscopic photos with interpretation by an ophthalmologist or optometrist

3) Eye imaging validated to match diagnosis from seven standard field stereoscopic photos results

must include one of the following:

documented and reviewed.

documented and reviewed.

and reviewed.

Supporting Definitions: (none)

Seq. #: 6682 Name: Eye Exam Date

Coding Instructions: Indicate the date the patient received an eye exam.

Yes

Target Value: N/A

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: EyeExam\_Date

**Parent Seq #:** 6680

Parent Name: Retinal or Dialated

Eye Exam

Parent Value: Yes

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User





Seq. #: 6700 Name: Insulin Pump				Technical Specifications	
<b>Seq.</b> #: 6700 Name:	ShortName:	InsulinPmp			
Coding Instructions:	Indicate if a	patient has been prescribed to start or continue to use an insulin pump.	Parent Seq #:		
Target Value:	N/A		Parent Name:		
<b>G</b>			Parent Value:		
Selections:	Code	Selection Text Definition	Missing Data:	Report	
	0	No	Harvested:	Yes (DCR)	
	1	Yes	Format:	Text (Categorical)	
	1		Default Value:	No	
Supporting Definitions:	Insulin Pun	np:	Usual Range:		
The insulin pump is not an artificial pancreas (because you still have to monitor your blood glucose level).			Valid Range:	Harr	
	Source: AD		DataSource:	User	

	Source: A	ADA			DataSource:	User
					Technical S	pecifications
Seq. #: 6702 N	ame: Insulin I	Pump Date			ShortName:	InsulinPmp_Date
Coding Instruc	tions: Indicate the	ne date the patient was p	prescribed to start or continu	ue use of an insulin pump.	Parent Seq #:	6700
Target \	/alue: N/A				Parent Name:	Insulin Pump
_					Parent Value:	Yes
Selec	tions: (none)				Missing Data:	Report
Supporting Defini	itions: (none)				Harvested:	Yes (DCR)
					Format:	Date (mm/dd/yyyy
					Default Value:	NULL
					Usual Range:	
					Valid Range:	
					DataSource:	User
	emer Continu	ious Clusoss Monit	toring		Technical S	pecifications
<b>Seq. #</b> : 6710 <b>N</b>	ame: Continu	ious Glucose Monit	oning		ShortName:	ContGluMonitor
Coding Instruc	tions: Indicate if monitoring	Indicate if the patient has been prescribed to start or continue continuous glucose			Parent Seq #:	
	monitoring	oring.			Parent Name:	
Target \	/alue: N/A				Parent Value:	
Selec	tions: Code	Selection Text	Definition		Missing Data:	Report
00.00		Selection rext			Harvested:	Yes (DCR)
	0	No			Format:	Text (Categorical)
	1	Yes			Default Value:	No
Supporting Defini	itions: (none)				Usual Range:	
ppo:g 2011111					Valid Range:	
					DataSource:	User



#### **D. Encounter Information**

**Technical Specifications** Name: Continuous Clucose Monitoring Date **Seq. #:** 6712 ShortName: ContGluMonitor\_Dat Coding Instructions: Indicate the date the patient was prescribed to start or continue continuous glucose monitoring. Parent Seq #: 6710 **Parent Name:** Continuous Glucose Target Value: N/A Monitoring Parent Value: Yes Selections: (none) Missing Data: No Action Supporting Definitions: (none) Harvested: Yes (DCR) Format: Date (mm/dd/yyyy) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Device ID **Seq. #:** 6720 ShortName: DeviceID Coding Instructions: Reserved for Future Use. Parent Seq #: **Parent Name:** Target Value: N/A Parent Value: Selections: (none) No Action Missing Data: Harvested: Yes (DCR) Supporting Definitions: (none) Format: Integer (5) **Default Value:** NULL **Usual Range:** Valid Range: 1-99999 DataSource: User **Technical Specifications Seq.** #: 6730 Name: Device Manufacturer ShortName: DevMfr Coding Instructions: Reservied for Future Use. Parent Seq #: **Parent Name:** Target Value: N/A Parent Value: Selections: (none) Missing Data: No Action Harvested: Yes (DCR) Supporting Definitions: (none) Format: Text (100) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User





	Technical S	Specifications
Seq. #: 6740 Name: Device Model	ShortName:	
Coding Instructions: Reserved for Future Use.	Parent Seq #:	Bormodol
	Parent Name:	
Target Value: N/A	Parent Value:	
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Text (100)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 6900 Name: Body Mass Index Screening	Technical S	<u>specifications</u>
·	ShortName:	BMIScreening
Coding Instructions: Indicate if the patient had a Body Mass Index screening was performed.	Parent Seq #:	
Target Value: Any occurrence between start of current encounter and completion of current encounter	Parent Name:	
Selections: Code Selection Text Definition	Parent Value:	
Selections. Code Selection Text Delimition	Missing Data:	Report
0 No	Harvested:	Yes (PINN)
1 Yes	Format:	Text (Categorical)
Supporting Definitions: (none)	Default Value:	No
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 6902 Name: Body Mass Index Screening Date		Specifications
Coding Instructions: Indicate the most recent documented date a Body Mass Index screening was performed.	ShortName:	BMIScreening_Date
County made make and mode reconstruction and a body made made estecting was performed.	Parent Seq #: Parent Name:	6900 Body Mass Index
Target Value: The last value on current encounter	T di oni i tanio.	Screening
Selections: (none)	Parent Value:	Yes
Ourse and the Definition of (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User



Supporting Definitions: (none)



# NCDR® Outpatient Registries v1.4 Data Dictionary - Full Specifications

DataSource:

User

#### D. Encounter Information

**Technical Specifications** Name: Body Mass Index Management Plan ShortName: BMIManagement\_PI Coding Instructions: Indicate if the patient has a documented BMI management plan. Parent Seq #: Note(s): **Parent Name:** A BMI management plan may include the following: documentation of future Parent Value: appointment, education, referral, prescription/administration of medication/dietary supplements, weight loss surgery. Missing Data: Report Harvested: Yes (PINN) Target Value: The value on current encounter Format: Text (Categorical) Selections: Code Selection Text Definition **Default Value:** No 0 **Usual Range:** Yes 1 Valid Range:

**Technical Specifications** Name: Prescription given for any Medication **Seq. #:** 8000 ShortName: RxEncounter Coding Instructions: This element has been retired effective v1.4. Parent Seq #: **Parent Name:** Target Value: The value between start of current encounter and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Yes (PINN) Harvested: No 0 Text (Categorical) Format: Yes 1 **Default Value:** Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Prescription generated and transmitted using an e-**Seg.** #: 8005 ShortName: prescribing system Parent Seq #: Coding Instructions: This element has been retired effective v1.4. **Parent Name:** Prescription given for any Medication Target Value: The value between start of current encounter and completion of current encounter Parent Value: Selections: Code Selection Text Definition Missing Data: Report Harvested: Yes (PINN) Nο 0 Format: Text (Categorical) Yes Default Value: NULL Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User





## E. Laboratory Results

O # 7000 News Linid David Obtained Data	Technical S	pecifications
Seq. #: 7000 Name: Lipid Panel Obtained Date	ShortName:	LipidPanelDate
Coding Instructions: Indicate the date blood was drawn for the most recent lipid panel.	Parent Seq #:	
Target Value. The leat value between birth and completion of current encounter	Parent Name:	
Target Value: The last value between birth and completion of current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7005 Name: Lipid Panel Fasting	Technical S	pecifications
Seq. #: 7005 Name. Lipid Faller Fasting	ShortName:	LipidPanelFasting
Coding Instructions: This element has been retired effective v1.4	Parent Seq #:	7000
Target Value: The last value between birth and completion of current encounter	Parent Name:	Lipid Panel Obtained Date
	Parent Value:	Not Null
Selections: Code Selection Text Definition	Missing Data:	No Action
0 No	Harvested:	Yes (PINN)
1 Yes	Format:	Text (Categorical)
Supporting Definitions: (none)	Default Value:	No.
Supporting Definitions. (11010)	Usual Range:	110
	Valid Range:	
	DataSource:	User
		Specifications
Seq. #: 7010 Name: Total Cholesterol		TotalCholesterol
Coding Instructions: Indicate the patient's most recent cholesterol in milligrams per deciliter (mg/dL) for the	Parent Seg #:	
most recent lipid panel.	Parent Name:	Lipid Panel
Target Value: The last value between birth and completion of current encounter		Obtained Date
	Parent Value:	Not Null
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Integer (4)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	1-1000
	DataSource:	User



#### E. Laboratory Results

**Technical Specifications** Name: High Density Lipoprotein (HDL) **Seq. #:** 7020 ShortName: Coding Instructions: Indicate the patient's most recent high density lipoproteins (HDL) in milligrams per Parent Seq #: 7000 deciliter (mg/dL) for the most recent lipid panel. Parent Name: Lipid Panel **Obtained Date** Target Value: The last value between birth and completion of current encounter Parent Value: Not Null Selections: (none) Missing Data: Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Integer (3) Default Value: **NULL Usual Range:** Valid Range: 1-300 DataSource: User **Technical Specifications Seq.** #: 7030 Name: Low Density Lipoprotein (LDL) ShortName: LDL Coding Instructions: Indicate the patient's most recent low density lipoproteins (LDL) in milligrams per deciliter Parent Seq #: 7000 (mg/dL) for the most recent lipid panel. **Parent Name:** Lipid Panel **Obtained Date** Target Value: The last value between birth and completion of current encounter Parent Value: Not Null Selections: (none) **Missing Data:** Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Integer (3) **Default Value:** NULL **Usual Range:** Valid Range: 1-800 DataSource: User **Technical Specifications Seq. #:** 7040 Name: Direct Low Density Lipoprotein (DLDL) ShortName: DLDL Coding Instructions: Indicate the patient's most recent direct low density lipoproteins (LDL) in milligrams per Parent Seq #: 7000 deciliter (mg/dL) for the most recent lipid panel. **Parent Name:** Lipid Panel **Obtained Date** Target Value: The last value on current encounter Parent Value: Not Null Selections: (none) Missing Data: Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Decimal (5,2) Default Value: NULL **Usual Range:** Valid Range: 1-7000 DataSource: User





## E. Laboratory Results

	<b>-</b>				Technical Specifications		
<b>Seq. #:</b> 7050 <b>Name:</b>	Triglycerio	des		ShortNan	ne: Triglycerides		
Coding Instructions:			glycerides in milligrams per de				
	most recent			Parent Nam			
Target Value:	The last value	ue between birth and co	mpletion of current encounter	Parent Valu	e: Not Null		
Selections:	(none)			Missing Dat	a: Report		
Supporting Definitions:	(none)			Harveste	d: Yes (DCR,PINN)		
	,			Forma	t: Integer (4)		
				Default Valu	e: NULL		
				Usual Rang	e:		
				Valid Rang	<b>e</b> : 1-7000		
				DataSourc	e: User		
Seq. #: 7052 Name:	Lipid Pan	el Ordered		<u>Technic</u>	al Specifications		
•				ShortNan	ne: LipidPanelOrdered		
_		t has been retired effect		Parent Seq Parent Nam			
_	-	ence between start of cul	rrent encounter and completior	n of current encounter Parent Valu	e:		
Selections:	Code	Selection Text	Definition	Missing Dat	a: Report		
	0	No		Harveste	d: Yes (PINN)		
	1	Yes		Forma	t: Text (Categorical)		
	-	100		Default Valu	e: No		
Supporting Definitions:	(none)			Usual Rang	e:		
				Valid Rang	e:		
				DataSourc	e: User		
Seq. #: 7054 Name:	Serum Gl	lucose Ordered		<u>Technic</u>	al Specifications		
•				ShortNan	e: GlucoseOrdered		
_		t has been retired effect		Parent Seq Parent Nam			
_		ue between birth and co	mpletion of current encounter	Parent Valu	e:		
Selections:	Code	Selection Text	Definition	Missing Dat	a: Report		
	0	No		Harveste	d: Yes (PINN)		
	1	Yes		Forma	t: Text (Categorical)		
	•	. 55		Default Valu	e: No		
Supporting Definitions:	(none)			Usual Rang	e:		
				Valid Rang	e:		
				DataSourc	e: User		





## E. Laboratory Results

O. v. # 7050 Names Chicago Data	Technical S	pecifications
Seq. #: 7056 Name: Glucose Date	ShortName:	SerumGlucoseDate
Coding Instructions: This element has been retired effective v1.4.	Parent Seq #:	
Target Value: The last value between birth and completion of current encounter	Parent Name:	
Target Value: The last value between birth and completion of current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7070 Name: Plasma Glucose Results	Technical S	pecifications
	ShortName:	PlasGluRes
<b>Coding Instructions:</b> Indicate the patient's plasma glucose level in milligrams per deciliter (mg/dL) for the most recent plasma glucose test.	Parent Seq #: Parent Name:	
Target Value: The last value between birth and completion of current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Control (no.15)	Harvested:	Yes (DCR,PINN)
Supporting Definitions: (none)	Format:	Decimal (5,2)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	1-1500
	DataSource:	User
Seq. #: 7072 Name: Plasma Glucose Results Date	Technical Sp	pecifications
·	ShortName:	PlasGluRes_Date
Coding Instructions: Indicate the date blood was drawn for the most recent plasma glucose test.	Parent Seq #:	7070
Target Value: The last value between birth and completion of current encounter	Parent Name:	Plasma Glucose Results
Selections: (none)	Parent Value:	Not Null
Supporting Definitions: (none)	Missing Data:	Report
Supporting Definitions. (1010)	Harvested:	Yes (DCR,PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User



#### E. Laboratory Results

**Technical Specifications** Name: HbA1c Percentage **Seq. #:** 7080 ShortName: HbA1c Coding Instructions: Indicate the patient's Hemoglobin A1c (HbA1c) percentage for the most recent Parent Seq #: Hemoglobin A1c (HbA1c) test. **Parent Name:** Parent Value: Target Value: The last value between birth and completion of current encounter Missing Data: Report Selections: (none) Yes (DCR,PINN) Harvested: Supporting Definitions: (none) Format: Decimal (4,1) Default Value: NULL **Usual Range:** Valid Range: 0.1-100.0 DataSource: User **Technical Specifications** Seq. #: 7082 Name: HbA1c Date ShortName: HbA1cDate Coding Instructions: Indicate the date blood was drawn for the most recent Hemoglobin A1c (HbA1c) test. Parent Seq #: 7080 **Parent Name:** HbA1c Percentage Target Value: The last value between birth and completion of current encounter Parent Value: Not Null Selections: (none) Missing Data: Report Yes (DCR,PINN) Harvested: Supporting Definitions: (none) Format: Date (mm/dd/yyyy) Default Value: NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: 2 Hour Plasma Glucose During Oral Glucose Tolerance **Seq.** #: 7090 ShortName: PlasGluOralTest Parent Seq #: Coding Instructions: Indicate the patient's most recent 2 hour plasma glucose during oral glucose tolerance **Parent Name:** test in mg/dL. Parent Value: Target Value: The last value on current encounter Missing Data: Report Harvested: Yes (DCR) Selections: (none) Format: Decimal (5,2) Supporting Definitions: PAD: Default Value: NULL **Usual Range:** 

PAD excludes renal, coronary, cerebral, and mesenteric vessels and aneurysm. Major symptoms can include

- ? Asymptomatic (confirmed by noninvasive diagnostic test)
- ? Claudication relieved by rest
- ? Ischemic rest pain
- ? Tissue loss (including ischemic ulcer and/or gangrene)
- ? Amputation for critical limb ischemia
- ? Vascular reconstruction, bypass surgery, or percutaneous revascularization in the arteries of the lower extremities
- ? Positive noninvasive test (e.g., ABI 0.90, ultrasound, MR or CT imaging demonstrating 50% diameter stenosis in any peripheral artery, i.e., aorta, iliac, femoral, popliteal, tibial,

Source: 2012 ACCF/AHA/ACR/SCAI/SIR/STS/SVM/SVN/SVS Key Data Elements and Definitions for Peripheral Atherosclerotic Vascular Disease

Valid Range:

DataSource:

1-1500

User





#### E. Laboratory Results

**Technical Specifications** Name: 2 Hour Plasma Glucose During Oral Glucose Tolerance **Seq. #:** 7092 PlasGluOralTest\_Da ShortName: **Test Date** Coding Instructions: Indicate the date of the patient's most recent 2 hour plasma glucose during oral glucose Parent Seq #: 7090 tolerance test. **Parent Name:** 2 Hour Plasma Glucose During Oral Target Value: The last value on current encounter Glucose Tolerance Test Selections: (none) Parent Value: Not Null Supporting Definitions: (none) Missing Data: Report Harvested: Yes (DCR) Format: Date (mm/dd/yyyy) Default Value: **NULL Usual Range:** Valid Range: DataSource: User **Technical Specifications Seq. #:** 7100 Name: Initial Labs ordered for newly diagnosed Heart Failure or ShortName: InitialLabsforHF patient new to the practice Parent Seq #: Coding Instructions: Indicate if the physician ordered Initial Labs for newly diagnosed Heart Failure. Newly **Parent Name:** diagnosed Heart Failure is defined as HF diagnosed within the past 12 months. Parent Value: Target Value: The value between 12 months prior to current encounter and completion of current Missing Data: Report encounter Yes (PINN) Harvested: Selections: Code Selection Text Definition Format: Text (Categorical) No **Default Value:** 0 **Usual Range:** Yes Valid Range: Supporting Definitions: (none) DataSource: User **Technical Specifications** Name: Estimated Glomerular Filtration Rate Electronic Medical **Seq. #:** 7105 ShortName: eGFR\_Emr Record Parent Seq #: Coding Instructions: This element has been retired effective v1.4. **Parent Name:** Parent Value: Target Value: N/A **Missing Data:** Report Selections: (none) Harvested: Yes (PINN) Supporting Definitions: (none) Format: Decimal (5,2) Default Value: NULL **Usual Range:** Valid Range: 0.01-999.99

DataSource:

User





#### E. Laboratory Results

Name: Estimated Glomerular Filtration Rate (eGFR) **Seq. #:** 7200

Coding Instructions: Indicate the most recent estimated glomerular filtration rate in ml/min.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName:

Parent Seq #: **Parent Name:** 

Parent Value:

Missing Data: Report

> Yes (DCR,PINN) Harvested:

Format: Decimal (5,2)

Default Value: No **Usual Range:** 60-120 1-999 Valid Range: DataSource:

Seq. #: 7202 Name: Estimated Glomular Filtration Rate (eGFR) Date

Coding Instructions: Indicate the date of the patient's most recent eGFR.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

User

eGFR\_Date ShortName:

Parent Seq #: 7200

**Parent Name:** Estimated

Glomerular Filtration Rate (eGFR)

Parent Value: Not Null Missing Data: Report

> Harvested: Yes (DCR,PINN)

Date (mm/dd/yyyy) Format:

**Default Value:** 

**Usual Range:** Valid Range:

DataSource: User

Seq. #: 7212 Name: Evidence of nephropathy Date

Coding Instructions: Indicate the date of the most recent screening for evidence of nephropathy.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

EviNephro\_date ShortName:

Parent Seq #: **Parent Name:** Parent Value:

Missing Data: No Action

Harvested: Yes (DCR)

> Format: Date (mm/dd/yyyy)

Default Value: NULL

**Usual Range:** Valid Range:

DataSource: User





#### E. Laboratory Results

**Technical Specifications** Name: Estimated Glomerular Filtration Rate Imputed **Seq. #:** 7215 ShortName: eGFR Imputed Coding Instructions: This element has been retired effective v1.4. Parent Seq #: **Parent Name:** Target Value: N/A Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (PINN) Supporting Definitions: (none) Format: Decimal (5,2) Default Value: NULL **Usual Range:** Valid Range: 0.01-999.99 DataSource: User **Technical Specifications** Name: Creatinine Clearance Seg. #: 7220 ShortName: CreatinineClearance Coding Instructions: Indicate the most recent document creatinine clearance in mL/min. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR,PINN) Supporting Definitions: (none) Format: Decimal (5,2) **Default Value:** NULL **Usual Range:** Valid Range: 0.01-999.99 DataSource: User **Technical Specifications** Name: Creatinine Clearance Date Seq. #: 7222 CreatinineClearance ShortName: Coding Instructions: Indicate the most recent documented date where creatinine clearance rate was recorded. Parent Seg #: Target Value: The last value on current encounter Parent Name: Creatinine Clearance Selections: (none) Parent Value: Not Null Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (DCR,PINN) Format: Date (mm/dd/yyyy) Default Value: NULL **Usual Range:** Valid Range: DataSource: User



#### E. Laboratory Results

**Seq. #:** 7225 **Name:** Creatinine Clearance Units

Coding Instructions: This element has been retired effective v1.4.

Target Value: N/A

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: CreatinineClearance

\_Units

Parent Seq #: 7220

Parent Name: Creatinine

Clearance

Decimal (5,2)

SerumCreatinine

Parent Value: Not Null

Missing Data: Report

Format:

Harvested: Yes (DCR,PINN)

Default Value: NULL

**Usual Range:** 

Valid Range: 0.01-999.99

DataSource: User

Seq. #: 7230 Name: Serum Creatinine

Coding Instructions: Indicate the most recent serum creatinine in mg/dL.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

Parent Seq #: Parent Name:

ShortName:

Parent Value:

Missing Data: Report

3

Harvested: Yes (DCR,PINN)

Decimal (5,2)

Default Value: NULL

**Usual Range:** 

Format:

Valid Range: 0.01-999.99

DataSource: User

Seq. #: 7232 Name: Serum Creatinine Date

Coding Instructions: Indicate the most recent documented date where serum creatinine rate was recorded.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: SerumCreatinine\_D

ate

Parent Seq #: 7230

Parent Name: Serum Creatinine

Parent Value: Not Null

Harvested: Yes (DCR,PINN)

Format: Date (mm/dd/yyyy)

No Action

Default Value: NULL

Usual Range: Valid Range:

Missing Data:

DataSource: User



#### E. Laboratory Results

**Technical Specifications** Name: Liver Function Tests - ALT **Seq. #:** 7300 ShortName: LiverFuncTestALT Coding Instructions: Indicate the patient's most recent ALT (alanine transaminase) in U/L. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Integer (4) **Default Value:** NULL **Usual Range:** 7-56 Valid Range: 1-2000 DataSource: User **Technical Specifications** Seq. #: 7302 Name: Liver Function Tests - ALT Date ShortName: LiverFuncTestALT\_ Coding Instructions: Indicate the most recent documented date of the ALT test. Parent Seq #: 7300 Target Value: The last value on current encounter **Parent Name: Liver Function Tests** - ALT Selections: (none) Parent Value: Not Null Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (DCR) Date (mm/dd/yyyy) Format: **Default Value: Usual Range:** Valid Range: DataSource: User **Technical Specifications** Seq. #: 7310 Name: Amylase ShortName: Amylase Coding Instructions: Indicate the patient's Amylase levels in U/L. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Integer (3) **Default Value:** NULL **Usual Range:** 23-140 Valid Range: 1-999 DataSource: User





## E. Laboratory Results

e « 7040 N. Amulaca Data	Technical S	pecifications
Seq. #: 7312 Name: Amylase Date	ShortName:	Amylase_Date
Coding Instructions: Indicate the date of the patient's most recent amylase result	Parent Seq #:	7310
Target Value: The lest value on current encounter	Parent Name:	Amylase
Target Value: The last value on current encounter	Parent Value:	Not Null
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7320 Name: Liver Function Tests - AST	Technical S	pecifications
·	ShortName:	LiverFuncTestAST
Coding Instructions: Indicate the patient's most recent AST (aspartate transaminase) in U/L.	Parent Seq #: Parent Name:	
Target Value: The last value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Integer (4)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7322 Name: Liver Function Tests - AST Date	Technical S	pecifications
Coding Instructions: Indicate the most recent documented date of the AST test.	ShortName:	LiverFuncTestAST_ Date
	Parent Seq #:	7320
Target Value: The last value on current encounter	Parent Name:	Liver Function Tests - AST
Selections: (none)	Parent Value:	
Supporting Definitions: (none)	Missing Data:	No Action
	Harvested:	Yes (DCR)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	





#### E. Laboratory Results

**Technical Specifications** Name: Liver Function Tests - Direct Bilirubin **Seq. #:** 7340 ShortName: LiverFuncTestDB Coding Instructions: Indicate the patient's most recent Direct Bilirubin in mg/dL. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Decimal (5,2) Default Value: NULL Usual Range: 0-0.3 Valid Range: 0-5 DataSource: User **Technical Specifications** Seq. #: 7342 Name: Liver Function Tests - Direct Bilirubin Date ShortName: LiverFuncTestDB\_D Coding Instructions: Indicate the most recent documented date of the direct Bilirubin test Parent Seq #: 7340 Target Value: The last value on current encounter **Parent Name: Liver Function Tests** - Direct Bilirubin Selections: (none) Parent Value: Not Null Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (DCR) Date (mm/dd/yyyy) Format: **Default Value: Usual Range:** Valid Range: DataSource: User **Technical Specifications Seq.** #: 7350 Name: Liver Function Tests - Total Bilirubin ShortName: LiverFuncTestTB Coding Instructions: Indicate the patient's most recent Total Bilirubin in mg/dL. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value:

Selections: (none)

Supporting Definitions: (none)

Missing Data: Report
Harvested: Yes (DCR)

Format: Decimal (5,2)

Default Value: NULL
Usual Range: 0-2
Valid Range: 0-10
DataSource: User



#### E. Laboratory Results

Seq. #: 7352 Name: Liver Function Tests - Total Bilirubin Date

Coding Instructions: Indicate the most recent documented date of the total Bilirubin test

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

**ShortName:** LiverFuncTestTB\_D

ate

Parent Seq #: 7350

Parent Name: Liver Function Tests

Total Bilirubin

Parent Value: Not Null

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User

Seq. #: 7360 Name: Blood Urea Nitrogen (BUN)

Coding Instructions: Indicate the most recent documented blood urea nitrogen (BUN) level. Blood urea

nitrogen (BUN) is a waste product in the blood from the breakdown of protein. The kidneys filter blood to remove urea. As kidney function decreases, the BUN levels

increase.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: BUN:

Blood urea nitrogen (BUN) is a waste product in the blood from the breakdown of protein.

The kidneys filter blood to remove urea. As kidney function decreases, the BUN levels

increase.

Source: ADA

Technical Specifications

ShortName: BUN

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Decimal (5,2)

Default Value: NULL

Usual Range: 6-24

Valid Range: 0-100

DataSource: User

Seq. #: 7362 Name: Blood Urea Nitrogen (BUN) Date

Coding Instructions: Indicate the most recent documented date where blood urea nitrogen (BUN) was

recorded.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: BUN\_Date

**Parent Seq #:** 7360

Parent Name: Blood Urea Nitrogen

(BUN)

Parent Value: Not Null

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: User



### PINNACLE Registry

## NCDR® Outpatient Registries v1.4 Data Dictionary - Full Specifications

#### E. Laboratory Results

**Technical Specifications** Seq. #: 7370 Name: Cystatin-C (Cystatin) ShortName: Cystatin Coding Instructions: Indicate the patient's most recent cystatin-C (cystatin). Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Decimal (5,2) Default Value: NULL 0.57-1.52 **Usual Range:** Valid Range: 0.01-9.99 DataSource: User **Technical Specifications** Seq. #: 7372 Name: Cystatin-C (Cystatin) Date ShortName: Cystatin\_Date Coding Instructions: Indicate the date of the patient's most recent cystatin results. Parent Seq #: Parent Name: Cystatin-C (Cystatin) Target Value: The last value on current encounter Parent Value: Not Null Selections: (none) Missing Data: No Action Harvested: Yes (DCR) Supporting Definitions: (none) Format: Date (mm/dd/yyyy) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: High-Sensitivity C-Reactive Protein (hs-CRP) **Seq. #:** 7380 ShortName: hsCRP Coding Instructions: Indicate the patient's most recent high-sensitivity C-reactive protein in mg/dL. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) **Missing Data:** Report Harvested: Yes (DCR) Supporting Definitions: (none) Decimal (5,2) Format: Default Value: NULL **Usual Range:** 0.1-10 0.01-20 Valid Range: DataSource: User



#### E. Laboratory Results

**Technical Specifications** Name: High-Sensitivity C-Reactive Protein (hs-CRP) Date **Seq. #:** 7382 ShortName: hsCRP\_Date Coding Instructions: Indicate the most recent documented date of the hs-CRP test. Parent Seq #: 7380 Parent Name: High-Sensitivity C-Target Value: The last value on current encounter Reactive Protein (hs -CRP) Selections: (none) Parent Value: Not Null Supporting Definitions: (none) Missing Data: No Action Harvested: Yes (DCR) Format: Date (mm/dd/yyyy) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Lipase **Seq. #:** 7390 ShortName: Lipase Coding Instructions: Indicate the patient's Lipase levels in U/L. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) **Missing Data:** Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Integer (4) **Default Value:** NULL **Usual Range:** 10-180 Valid Range: 1-3000 DataSource: User **Technical Specifications** Seq. #: 7392 Name: Lipase Date ShortName: Lipase\_Date Coding Instructions: Indicate the date of the patient's most recent lipase result Parent Seq #: 7390 Parent Name: Lipase Target Value: The last value on current encounter Parent Value: Not Null Selections: (none) Missing Data: No Action Harvested: Yes (DCR) Supporting Definitions: (none) Format: Date (mm/dd/yyyy) Default Value: NULL **Usual Range:** Valid Range: DataSource: User



### E. Laboratory Results

	Technical 9	pecifications
Seq. #: 7400 Name: Thyroid-Stimulating Hormone (TSH)	ShortName:	
Coding Instructions: Indicate the patient's most recent thyroid-stimulating hormone test in mg/dL.	Parent Seq #:	1011
	Parent Name:	
Target Value: The last value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR)
Cupporting Definitions. (Note)	Format:	Decimal (5,2)
	Default Value:	NULL
	Usual Range:	0.4-2.5
	Valid Range:	0.1-9.9
	DataSource:	User
Seq. #: 7402 Name: Thyroid-Stimulating Hormone (TSH) Date	Technical S	pecifications
Seq. #: 7402 Name. Thyroid-offindiating Floriflone (1011) Date	ShortName:	TSH_Date
Coding Instructions: Indicate the most recent documented date of the TSH test.	Parent Seq #:	7400
Target Value: The last value on current encounter	Parent Name:	Thyroid-Stimulating Hormone (TSH)
Selections: (none)	Parent Value:	Not Null
Own setting Definitions (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7410 Name: Uric Acid	<u>Technical S</u>	pecifications
	ShortName:	UricAcid
Coding Instructions: Indicate the most recent documented uric acid.	Parent Seq #: Parent Name:	
Target Value: The last value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Decimal (5,2)
	Default Value:	NULL
	Usual Range:	2.4-7.2
	Valid Range:	0.1-999.9
	DataSource:	User





### E. Laboratory Results

O # 7440 Name Urio Acid Data	Technical Specifications	
Seq. #: 7412 Name: Uric Acid Date	ShortName:	UricAcid_Date
Coding Instructions: Indicate the most recent documented date of the Uric Acid test	Parent Seq #:	7410
Target Value: The last value on current encounter	Parent Name:	Uric Acid
	Parent Value:	Not Null
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User
Seq. #: 7420 Name: 24 Hour Urine Protein	Technical S	Specifications
•	ShortName:	UrineProtein
Coding Instructions: Indicate the most recent documented 24 hour urine protein.	Parent Seq #: Parent Name:	
Target Value: The last value on current encounter	Parent Value:	
Selections: (none)	Missing Data:	Report
Supporting Definitions: (none)	Harvested:	Yes (DCR)
	Format:	Decimal (5,2)
	Default Value:	NULL
	Usual Range:	1-10
	Valid Range:	0.1-999.9
	DataSource:	User
Seq. #: 7422 Name: 24 Hour Urine Protein Date	Technical S	Specifications
•	ShortName:	UrineProtein_Date
Coding Instructions: Indicate the date of the patient's most recent urine protein test.	Parent Seq #:	7420
Target Value: The last value on current encounter	Parent Name:	24 Hour Urine Protein
Selections: (none)	Parent Value:	
Supporting Definitions: (none)	Missing Data:	No Action
capporting communio. (******/	Harvested:	Yes (DCR)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	User



#### E. Laboratory Results

Name: Urine albumin: Creatine ratio (UACR) **Seq. #:** 7430

Coding Instructions: Inidcate the most recent urine albumin:creatine ratio (UACR).

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: Urine albumin:

Creatine ratio is a test for levels of albumin and creatine in the blood as an indicator of

nephropathy

Albuminuria is a condition in which the urine has more than normal amounts of a protein

called albumin.

Albuminuria may be a sign of nephropathy (kidney disease)

Source: ADA Creatinine:

Creatinine is a waste product from protein in the diet and from the muscles of the body. Creatinine is removed from the body by the kidneys; as kidney disease progresses, the

level of creatinine in the blood increases.

Source: ADA

**Technical Specifications** 

ShortName: **UACR** 

Parent Seq #: **Parent Name:** 

Parent Value:

Missing Data: Report Yes (DCR) Harvested:

> Format: Date (mm/dd/yyyy)

> > User

Default Value: NULL **Usual Range:** 1-30 Valid Range: 1-999

DataSource:

Name: Urine albumin:creatine ratio (UACR) Date **Seq. #**: 7432

Coding Instructions: Indicate the date of the patient's most recent urine albumin test.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

UACR\_Date ShortName:

Parent Seq #: 7430

**Parent Name:** Urine albumin:

Creatine ratio (UACR)

**Parent Value:** Not Null Missing Data: No Action

> Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

NULL

**Default Value: Usual Range:** 

Valid Range:

DataSource: User

Name: Complete Blood Count - White Blood Cells (WBC)

Coding Instructions: Indicate the patient's white blood cell (WBC) count.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: WBC

Parent Seg #: **Parent Name:** 

**Parent Value:** 

Missing Data: Report Harvested: Yes (DCR)

> Format: Integer (5) NULL

Default Value: **Usual Range:** 3500-10500

Valid Range: 1-99999 DataSource:



#### E. Laboratory Results

Seq. #: 7502 Name: Complete Blood Count - White Blood Cells (WBC) Date

Coding Instructions: Indicate the most recent documented date of the white blood cell count.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

Technical Specifications

ShortName: WBC\_Date

Parent Seq #: 7500

Parent Name: Complete Blood

Count - White Blood

Cells (WBC)

Parent Value: Not Null

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User

Seq. #: 7510 Name: Complete Blood Count - Hemoglobin (HgB)

Coding Instructions: Indicate the patient's Hemoglobin (HgB) count.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: HgB

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Report

Harvested: Yes (DCR)

Format: Decimal (3,1)

Default Value: NULL
Usual Range: 12-17.5
Valid Range: 0.1-99.9

DataSource: User

Seq. #: 7512 Name: Complete Blood Count - Hemoglobin (HgB) Date

Coding Instructions: Indicate the most recent documented date of the hemoglobin count

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

**ShortName:** HgB\_Date

Parent Seq #: 7510

Parent Name: Complete Blood

Count - Hemoglobin

(HgB)

Parent Value: Not Null

Missing Data: No Action

Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

Usual Range: Valid Range:

DataSource: User





#### E. Laboratory Results

**Technical Specifications** Name: Complete Blood Count - Hematocrit **Seq. #:** 7520 ShortName: Hematocrit Coding Instructions: Indicate the patient's Hematocrit count. Parent Seq #: **Parent Name:** Target Value: The last value on current encounter Parent Value: Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Decimal (4,1) Default Value: NULL **Usual Range:** 34.9-50 Valid Range: 0.1-100 DataSource: User **Technical Specifications** Seq. #: 7522 Name: Complete Blood Count - Hematocrit Date ShortName: Hematocrit\_Date Coding Instructions: Indicate the most recent documented date of the hematocrit count. Parent Seq #: 7520 **Parent Name:** Complete Blood Target Value: The last value on current encounter Count - Hematocrit **Parent Value:** Not Null Selections: (none) **Missing Data:** No Action Supporting Definitions: (none) Harvested: Yes (DCR) Format: Date (mm/dd/yyyy) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Name: Complete Blood Count - Platelet **Sea.** #: 7530 ShortName: Platelet Coding Instructions: Indicate the patient's platelet count. Parent Seq #: Parent Name: Target Value: The last value on current encounter **Parent Value:** Selections: (none) Missing Data: Report Harvested: Yes (DCR) Supporting Definitions: (none) Format: Decimal (4,1) Default Value: NULL **Usual Range:** 34.9-50 Valid Range: 0.1-100 DataSource:



#### **E. Laboratory Results**

Seq. #: 7532 Name: Complete Blood Count - Platelet Date

Coding Instructions: Indicate the most recent documented date of the hematocrit count.

Target Value: The last value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

**ShortName:** Platelet\_Date

**Parent Seq #:** 7530

Parent Name: Complete Blood

Count - Platelet

Parent Value: Not Null

Missing Data: No Action
Harvested: Yes (DCR)

Format: Date (mm/dd/yyyy)

Default Value: NULL

**Usual Range:** 

Valid Range:

DataSource: User





### F. Medications

Seq. #: 9300 Name: M	edication ID		Technical S	specifications
•			ShortName:	MedID
Coding Instructions: Inc	dicate the NCDR-assigned IDs for	or the medications the patient was prescribed.	Parent Seq #:	
Target Value: The	ne value between start of current	t encounter and completion of current encounter	Parent Name:	
_		· oncounter and completion of canonic encounter	Parent Value:	
Selections: (r	ione)		Missing Data:	Report
Supporting Definitions: (n	one)		Harvested:	Yes (DCR,PINN)
			Format:	Integer (3)
			Default Value:	NULL
			Usual Range:	
			Valid Range:	1-999
			DataSource:	User
<b>Seq. #</b> : 9301 <b>Name</b> : D	ose Strength		Technical S	pecifications
•			ShortName:	DoseStrength
Coding Instructions: Inc	dicate the dosing strength for each	ch medication that is prescibed/continued.	Parent Seq #:	9300
Target Value: Th	ne last value on current encounte	er	Parent Name:	Medication ID
_			Parent Value:	Not Null
Selections: (r	ione)		Missing Data:	Report
Supporting Definitions: (n	one)		Harvested:	Yes (DCR,PINN)
			Format:	Decimal (6,2)
			Default Value:	NULL
			Usual Range:	
			Valid Range:	0.01-9999.99
			DataSource:	User
<b>Seq. #</b> : 9302 <b>Name</b> : D	osing Measure		Technical S	pecifications
•			ShortName:	DosMeasure
Coding Instructions: Inc	dicate the dosage measurment for	or each medication prescribed/continued (eg. g, mg).	Parent Seq #:	9300
Target Value: Th	ne last value on current encounte	er	Parent Name:	Medication ID
_			Parent Value:	Not Null
Selections: (	Code Selection Text	Definition	Missing Data:	Report
1	mg		Harvested:	Yes (DCR,PINN)
2	-		Format:	Text (Categorical)
3			Default Value:	NULL
4			Usual Range:	
			Valid Range:	
Supporting Definitions: (n	one)		DataSource:	User





#### F. Medications

Seq. #: 9303 Name: Dose Frequency

Coding Instructions: Indicate the frequency for which the patient should take the prescribed medication

dosage.

Target Value: The last value on current encounter

_			
Selections:	Code	Selection Text	Definition
	1	once daily	
	2	twice daily	
	3	three times daily	
	4	four times daily	
	5	five times daily	
	6	with meals	
	7	once every other day	
	8	once weekly	
	9	twice weekly	
	10	three times weekly	

Technical Specifications

ShortName: DoseFrqncy

Parent Seq #: 9300

Parent Name: Medication ID

Parent Value: Not Null
Missing Data: Report

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: NULL Usual Range:

Valid Range:

DataSource: User

Supporting Definitions: (none)

**Selections** 

Seq. #: 9305 Name: Medication Administered

Coding Instructions: Indicate if the medication was prescribed/continued or was not prescribed for either a

medical, system, or patient reason.

Target Value: The value between start of current encounter and completion of current encounter

S:	Code	Selection Text	Definition
	1	Yes	Medications was administered or prescribed.
	4	No - Patient Reason	Unable to administer/prescribe due to a patient reason such as patient refusal of medication. Patient reason may include religious
	5	No - Medical Reason	Unable to administer/prescribe due to a medical reason such as an allergies, contraindications side effects, intolerances, medical interactions, and safety concerns.
	6	No - System Reason	Unable to administer/prescribe due to system

Supporting Definitions: (none)

	•
ShortName:	MedAdmin
Parent Seq #:	9300
Parent Name:	Medication ID
Parent Value:	Not Null
Missing Data:	Report
Harvested:	Yes (DCR,PINN)
Format:	Text (Categorical)

**Technical Specifications** 

Default Value: NULL
Usual Range:
Valid Range:
DataSource: User





### G. Hospitalizations

o.z.o. N	Hereitel A Julia in Date	Technical S	pecifications
<b>Seq. #:</b> 9500 <b>Name</b> :	: Hospital Admission Date	ShortName:	HospitalAdmit_Date
Coding Instructions:	Indicate the most recent date of admission to a hospital or other acute healthcare facility for the patient.	Parent Seq #: Parent Name:	
Target Value:	The last value between birth and current encounter	Parent Value:	
Selections	: (none)	Missing Data:	Report
		Harvested:	Yes (DCR,PINN)
Supporting Definitions	: (none)	Format:	Date (mm/dd/yyyy)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User
Seg #: 9502 Name	: Hospital Discharge Date	Technical S	pecifications
•		ShortName:	HospitalDCDate
Coding Instructions:	Indicate the date the patient was discharged from the most recent hospitalization admission.	Parent Seq #:	
		Parent Name:	
Target Value:	The last value between birth and current encounter	Parent Value:	
Selections	: (none)	Missing Data:	Report
	(nana)	Harvested:	Yes (DCR,PINN)
Supporting Definitions	: (none)	Format:	Date (mm/dd/yyyy)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	User
Seg #: 9505 Name:	Primary Reason for Admission		pecifications
•	Indicate the primary diagnosis of the event that prompted the most recent hospitalization	ShortName:	Admission_Reason_ Code
-	admission, as determined by the judgment of the investigator. Utilize latest ICD code (e.g., ICD-9 or ICD-10). May be the same as principal discharge diagnosis.	Parent Seq #: Parent Name:	9500 Hospital Admission Date
Target Value:	The last value between birth and current encounter	Parent Value:	Not Null
Selections	: (none)	Missing Data:	Report
Supporting Definitions	· (none)	Harvested:	Yes (DCR,PINN)
Supporting Deminions		Format:	Text (20)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	





#### G. Hospitalizations

**Technical Specifications** Name: Secondary Diagnosis **Seq. #:** 9507 ShortName: SecondDiag Coding Instructions: Indicate the secondary diagnosis of the even that prompted the most recent Parent Seq #: hospitalization admission, as determined by the judgement of the investigator if a **Parent Name:** secondary diagnosis is made. Utilize latest ICD code (e.g., ICD-9 or ICD-10). May be the same as principal discharge diagnosis. Parent Value: Missing Data: Report Target Value: The last value between birth and current encounter Yes (DCR,PINN) Harvested: Selections: (none) Format: Text (20) Supporting Definitions: (none) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: User **Technical Specifications** Seg. #: 9510 Name: Coding Standard ShortName: Coding\_Standard Coding Instructions: Indicate the coding standard used in recording admission reason. Parent Seq #: 9505 **Parent Name:** Primary Reason for Target Value: The last value between birth and current encounter Admission **Parent Value:** Not Null Selections: Code Selection Text Definition **Missing Data:** Report ICD-9 1 Yes (DCR,PINN) Harvested: 2 ICD-10 Format: Text (Categorical) **Default Value:** NULL Supporting Definitions: (none) **Usual Range:** Valid Range: DataSource: User



### Z. Administration

On the 4000 Marrian Data File Name	Technical S	pecifications
Seq. #: 1000 Name: Data File Name	ShortName:	DataFile_Name
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Parent Seq #:	
Target Value: N/A	Parent Name:	
	Parent Value:	
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (100)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	Automatic
Seq. #: 1005 Name: Data File Creation Date Time	Technical S	pecifications
·	ShortName:	DataFile_CreationDt Time
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Doront Son #.	Time
Target Value: N/A	Parent Seq #: Parent Name:	
Selections: (none)	Parent Value:	
	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Date (mm/dd/yyyy)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	Automatic
Seq. #: 1010 Name: Data File Total Visits	Technical S	pecifications
Seq. #. 1010 Name. Data File Total Visits	ShortName:	Datafile_TotalVisits
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Parent Seq #: Parent Name:	
Target Value: N/A	Parent Value:	
Selections: (none)	Missing Data:	No Action
Supposition Policitions (none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions: (none)	Format:	Integer (9)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	Automatic





### Z. Administration

Seq. #: 1015 Name: Data File Source Identification Number	Technical S	pecifications
Seq. #: 1015 Name: Data File Source Identification Number	ShortName:	Datafile_SourceID
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Parent Seq #:	
Target Value: N/A	Parent Name:	
	Parent Value:	
Selections: (none)	Missing Data:	No Action
Supporting Definitions: (none)	Harvested:	Yes (DCR,PINN)
	Format:	Text (20)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	DataSource:	Automatic
Seq. #: 1020 Name: Practice Total Visits		Specifications
		Practice_TotalVisits
Coding Instructions: This element has been retired effective PINNACLE v1.3.	Parent Seq #:	
Target Value: N/A	Parent Name:	
Selections: (none)	Parent Value:	Nia Aattaa
	Missing Data:	No Action
Supporting Definitions: (none)	Harvested: Format:	Yes (DCR,PINN)
	Default Value:	Integer (9) NULL
	Usual Range:	NULL
	Valid Range: DataSource:	Automatic
		Specifications
Seq. #: 1021 Name: Timeframe of Data Submission	ShortName:	-
Coding Instructions: Indicate the time frame of data included in the data submission. Format: YYYYQQ.		Timename
e.g.,2013Q4	Parent Seq #: Parent Name:	
Target Value: N/A	Parent Value:	
	Missing Data:	Illegal
Selections: (none)		Yes (DCR,PINN)
Supporting Definitions: (none)	Format:	Text (6)
	Default Value:	NULL
	Usual Range:	
	Valid Range:	
	1	





### Z. Administration

O 4- 4005 N	Location Total Visite	Technical S	pecifications
<b>Seq. #:</b> 1025 <b>Name:</b>	Location Total visits	ShortName:	Location_TotalVisits
Coding Instructions:	This element has been retired effective PINNACLE v1.3.	Parent Seq #:	
Target Value:	N/Δ	Parent Name:	
_		Parent Value:	
Selections	: (none)	Missing Data:	No Action
Supporting Definitions:	: (none)	Harvested:	Yes (DCR,PINN)
		Format:	Integer (9)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	Automatic
Seg #: 1030 Name:	Encounter Unique Key	Technical S	<u>specifications</u>
•		ShortName:	EncounterKey
Coding Instructions:	Indicate the unique key associated with each patient encounter as assigned by the EMR/EHR or your software application.	Parent Seq #: Parent Name:	
Target Value:	N/A	Parent Value:	
Selections	(none)	Missing Data:	Illegal
		Harvested:	Yes (DCR,PINN)
Supporting Definitions:	(none)	Format:	Text (50)
		Default Value:	NULL
		Usual Range:	
		Valid Range:	
		DataSource:	Automatic
Seg #: 1040 Name:	Transmission Number	Technical S	pecifications
•		ShortName:	Xmsnld
Coding Instructions:	This is a unique number created, and automatically inserted by the software into export file. It identifies the number of times the software has created a data submission file. The transmission number should be incremented by one every time the data submi	Parent Seq #: Parent Name:	
		Parent Value:	
Target Value:	N/A	Missing Data:	Illegal
Selections	(none)	Harvested:	Yes (DCR,PINN)
Supporting Definitions:	(none)	Format:	Integer (9)
Supporting Demindons.		Default Value:	NULL
		Usual Range:	
		Valid Range:	1-99999999
		DataSource:	Automatic



#### Z. Administration

**Technical Specifications** Name: Vendor Identifier **Seq. #**: 1050 ShortName: Vendorld Coding Instructions: Vendor identification (agreed upon by mutual selection between the vendor and the Parent Seq #: NCDR) to identify software vendor. This is entered into the schema automatically by **Parent Name:** vendor software. Vendors must use consistent name identification across sites. Changes to vendor name identification must be approved by the NCDR. Parent Value: Missing Data: Illegal Target Value: N/A Yes (DCR,PINN) Harvested: Selections: (none) Format: Text (15) Supporting Definitions: (none) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: Automatic **Technical Specifications** Name: Vendor Software Version **Seq. #:** 1060 ShortName: VendorVer Coding Instructions: Vendor's software product name and version number identifying the software which Parent Seq #: created this record (assigned by vendor). Vendor controls the value in this field. This is **Parent Name:** entered into the schema automatically by vendor software. Parent Value: Target Value: N/A Missing Data: Illegal Selections: (none) Harvested: Yes (DCR,PINN) Format: Text (20) Supporting Definitions: (none) **Default Value:** NULL **Usual Range:** Valid Range: DataSource: Automatic **Technical Specifications** Seq. #: 1070 Name: Registry Identifier ShortName: RegistryId Coding Instructions: The NCDR registry identifier describes the data registry to which these records apply. It is Parent Seq #: implemented in the software at the time the data is collected and records are created. **Parent Name:** This is entered into the schema automatically by software. **Parent Value:** Target Value: N/A Missing Data: Illegal Selections: (none) Harvested: Yes (DCR,PINN) Format: Text (20) Supporting Definitions: (none) **Default Value:** ACC-NCDR-PINN **Usual Range:** Valid Range: DataSource: Automatic





#### Z. Administration

Seq. #: 1080 Name: Registry Version

Coding Instructions: Registry version describes the version number of the Data Specifications/Dictionary, to

which each record conforms. It identifies which fields should have data, and what are the valid data for each field. It is the version implemented in the software at the time the data is collected and the records are created. This is entered into the schema automatically by

software.

Target Value: N/A

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: RegistryVer

Parent Seq #: Parent Name:

Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Text (10)

Default Value: 1.4

Usual Range:

Valid Range:

DataSource: Automatic

Seq. #: 1095 Name: Submission Type

Coding Instructions: Indicate if the data contained in the harvest/data file contains PINNACLE registry records,

diabetic records, or all patient encounter records.

Target Value: N/A

Selections: Code Selection Text Definition

1 All Encounter Records Contains all patients and all encounter records with eligible visits to the physician office with an

Encounter Date.

2 PINNACLE Encounter Records Only Patients and all encounter records with eligible visits to the physician office with an Encounter Date.

3 Diabetes Encounter
Records Only
Contains all completed PINN-Diabetes
Collaborative Registry (DCR) dataset for all
patients and all encounter records with eligible
visits to the physician office with an Encounter

Date.

Supporting Definitions: (none)

Technical Specifications

ShortName: SubmissionType

Parent Seq #: Parent Name: Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Text (Categorical)

Default Value: Usual Range:

Valid Range:

DataSource: User

Seq. #: 1520 Name: Practice ID

Coding Instructions: Indicate the Practice Identification number assigned to the Practice by the ACC-NCDR.

Note(s):

The Practice ID will display in the General Information Section of the data collection form however the coding instructions will move to Administration Section in the data

dictionary.

Target Value: The value on current encounter

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: PracticeID

Parent Seq #: Parent Name: Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Integer (6)

Default Value: NULL

Usual Range: Valid Range:

DataSource: Automatic



#### Z. Administration

Seq. #: 1521 Name: Practice Name

Coding Instructions: Indicate the full name of the practice.

Target Value: N/A

Selections: (none)

Supporting Definitions: (none)

**Technical Specifications** 

ShortName: PracName

Parent Seq #:
Parent Name:
Parent Value:

Missing Data: Illegal

Harvested: Yes (DCR,PINN)

Format: Text (50)

Default Value: NULL

Usual Range:

Valid Range:

DataSource: Automatic