





Section: Demographics		Parent: Root	
Element: 2000	Last Name		Technical Specification

Coding Instruction: Indicate the patient's last name. Hyphenated names should be recorded with a hyphen.

Target Value: The value on arrival at this facility

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

Code: 1000142463

Code System: ACC NCDR Short Name: LastName

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Yes
Data Type: LN
Precision: 50
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Element: 2010 First Name Technical Specification

Coding Instruction: Indicate the patient's first name.

Target Value: The value on arrival at this facility

Code: 1000142463
Code System: ACC NCDR
Short Name: FirstName
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Yes
Data Type: FN
Precision: 50
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Element: 2020 Middle Name Technical Specification Code: 1000142463 Code: 1000142463 Code: System: ACC NCDP

Note(s):

It is acceptable to specify the middle initial.

If there is no middle name given, leave field blank.

If there are multiple middle names, enter all of the middle names sequentially.

If the name exceeds 50 characters, enter the first 50 letters only.

Target Value: The value on arrival at this facility

Code: 1000142463
Code System: ACC NCDR
Short Name: MidName
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: MN
Precision: 50
Selection Type: Single
Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Demographics		Parent: Root
Element: 2050	Birth Date	Technical Specification
Coding Instruction:	Indicate the patient's date of birth.	Code: 1000142447 Code System: ACC NCDR
Target Value:	The value on arrival at this facility	Short Name: DOB Missing Data: Report
		Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
		Is Identifier: No
		Is Base Element: Yes
		Is Followup Yes
		Data Type: DT
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:

Technical Specification Element: 2030 SSN Code: 2.16.840.1.113883.4.1 Coding Instruction: Indicate the patient's United States Social Security Number (SSN). Code System: United States Social Security Number (SSN) Short Name: SSN If the patient does not have a US Social Security Number (SSN), leave blank and check 'SSN Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, Target Value: The value on arrival at this facility TMVrpr, TTVP) Is Identifier: No Vendor Instruction: Patient's SSN must be 9 numeric characters long Is Base Element: Yes Is Followup Yes Data Type: ST Precision: 9 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Element: 2031 SSN N/A
Operator: Equal

Data Source: User

Value: No (or Not Answered)

Coding Instruction: Indicate if the patient does not have a United States Social Security Number (SSN).

Target Value: The value on arrival at this facility

SSN N/A

Technical Specification
Code: 2.16.840.1.113883.4.1
Code System: United States Social Security

Parent/Child Validation

Number (SSN)

Short Name: SSNNA Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Element: 2031







Valid Range: 1 - 999,999,999 Data Source: Automatic

Data Source: User

Data Source: User

Technical Specification

Section: Demographics	Parent: Root		
Element: 2040	Patient ID	Technic	al Specification
Coding Instruction:	Indicate the number created and automatically inserted by the software that uniquely identifies this patient.	Code System: Short Name:	NCDRPatientID
	Note(s): Once assigned to a patient at the participating facility, this number will never be changed or reassigned to a different patient. If the patient returns to the same participating facility or for		Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
Target Value:	follow up, they will receive this same unique patient identifier. The value on arrival at this facility	Is Identifier: Is Base Element: Is Followup Element:	Yes
		Data Type: Precision:	
		Selection Type: Unit of Measure: Default Value: Usual Range:	·

Technical Specification Element: 2045 Other ID Code: 2.16.840.1.113883.3.3478.4.843 Coding Instruction: Indicate an optional patient identifier, such as medical record number, that can be associated Code System: ACC NCDR with the patient. Short Name: OtherID Target Value: N/A Missing Data: No Action Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: Yes Data Type: ST Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

On the selection of the	Indicate the materials and a bidle	Code:	1000142448
Coding Instruction:	Indicate the patient's sex at birth.	Code System:	ACC NCDR
Target Value:	The value on arrival at this facility	Short Name:	Sex
		Missing Data:	Report
		Harvested:	Yes (BDS, TAVR, TMVR,
			TMVrpr, TTVP)
		Is Identifier:	No
		Is Base Element:	
		Is Followup	Vas
		Element:	103
		Data Type:	CD
		Precision:	
		Selection Type:	Single
		Unit of Measure:	
		Default Value:	Null
		Usual Range:	
		Valid Range:	

Person Sex - 1.3.6.1.4.1.19376.1.4.1.6.5.19

Sex

Element: 2060

Selection	Definition	Source	Code	Code System
Male			M HL7 Ac	dministrative Gender
Female			F HL7 Ac	dministrative Gender







Section: Demographics	Parent: Root	
Element: 2065	Patient Zip Code	Technical Specification
Cadina Instruction	Indicate the national claim of Chatan Double Commission with and of their arithmen, uncidence	Code: 1000142449
Coding instruction:	Indicate the patient's United States Postal Service zip code of their primary residence.	Code System: ACC NCDR
	Note(s):	Short Name: ZipCode
	If the patient does not have a U.S. residence, or is homeless, leave blank and check 'Zip Code	Missing Data: Report
	NA'.	Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Vendor Instruction:	Patient's zip code must be 5 numeric characters long.	Is Base Element: Yes
	Tallotto 2p ood matro o tallotto o	Is Followup Yes
		Element:
		Data Type: ST
		Precision: 5
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation
		Element: 2066 Zip Code N/A
		Operator: Equal
		Value: No (or Not Answered)

Element: 2066	Zip Code N/A	Technic	al Specification
0 - 11 1 1 1 1	Indicate if the patient does not have a United States Postal Service zip code.	Code:	1000142449
Coding instruction:		Code System:	ACC NCDR
	Note(s):	Short Name:	ZipCodeNA
	This includes patients who do not have a U.S. residence or are homeless.	Missing Data:	Report
Target Value:	The value on arrival at this facility	Harvested:	Yes (TAVR, TMVR, TMVrpr, TTVP)
		Is Identifier:	No
		Is Base Element:	
		Is Followup Element:	Yes
		Data Type:	BL
		Precision:	
		Selection Type:	Single
		Unit of Measure:	
		Default Value:	Null
		Usual Range:	
		Valid Range:	
		Data Source:	User

Element: 2070	Race - White	Technical Specification
O a dia a la atau ati a a	Ladicate With a national is Mileton and determine allow the mathematical formation	Code: 2106-3
Coding Instruction:	Indicate if the patient is White as determined by the patient/family.	Code System: HL7 Race
	Note(s):	Short Name: RaceWhite
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (BDS, TAVR, TMVR,
Tannat Value	The value on emissi at this facility.	TMVrpr, TTVP)
rarget value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	White (race)	Is Base Element: Yes
	Having origins in any of the original peoples of Europe, the Middle East, or North Africa.	Is Followup No
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Element:
		Data Type: BL
	Limbity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User







Data Source: User

Section: Demographics	Parent: Root		
Element: 2071	Race - Black/African American	Technic	al Specification
Coding Instruction:	Indicate if the patient is Black or African American as determined by the patient/family.	Code: Code System:	2054-5 HL7 Race
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: Missing Data: Harvested:	Report Yes (BDS, TAVR, TMVR,
Target Value:	The value on arrival at this facility	Is Identifier:	TMVrpr, TTVP) No
Supporting Definition:	Black/African American (race) Having origins in any of the black racial groups of Africa. Terms such as "Haitian" or "Negro"	Is Base Element:	
	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	Element: Data Type: Precision:	BL
	Ethnicity	Selection Type: Unit of Measure:	Single
		Default Value: Usual Range: Valid Range:	Null

Element: 2073	Race - American Indian/Alaskan Native	Technical Specification
Coding Instruction	: Indicate if the patient is American Indian or Alaskan Native as determined by the patient/family.	Code: 1002-5
County instruction.		Code System: HL7 Race
	Note(s):	Short Name: RaceAmIndian
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (BDS, TAVR, TMVR,
Towns Value	The value on emissi at this facility.	TMVrpr, TTVP)
rarget value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	American Indian or Alaskan Native (race)	Is Base Element: Yes
	Having origins in any of the original peoples of North and South America (including Central	Is Followup No
	America), and who maintains tribal affiliation or community attachment.	Element: 110
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Data Type: BL
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User

Element: 2072	Race - Asian	Technical Specification
Cadina Instruction	Indicate if the neticut is Asian as determined by the neticut/family	Code: 2028-9
Coding instruction:	: Indicate if the patient is Asian as determined by the patient/family.	Code System: HL7 Race
	Note(s):	Short Name: RaceAsian
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (BDS, TAVR, TMVR,
Tananat Walana	The code of a second of the facility	TMVrpr, TTVP)
rarget value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Asian (race)	Is Base Element: Yes
	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	Is Followup No
		Element: 110
		Data Type: BL
		Precision:
	Ethnicity	Selection Type: Single
	•	Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User







Section: Demographics	Parent: Root	
Element: 2080	Race - Asian Indian	Technical Specification
On the sales at several and	Indicate William attention Aster Indian and attentional by the matter (Many Sci	Code: 2029-7
Coding instruction:	Indicate if the patient is Asian Indian as determined by the patient/family.	Code System: HL7 Race
	Note(s):	Short Name: RaceAsianIndian
	If the patient has multiple race origins, specify them using the other race selections in addi	
	to this one.	Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Asian Indian	Is Base Element: Yes
	Having origins in any of the original peoples of India.	Is Followup Element: No
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race	and Data Type: BL
	Ethnicity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation
		Element: 2072 Race - Asian
		Operator: Equal
		Value: Yes

Element: 2081	Race - Chinese	Technical Specification
		Code: 2034-7
Coding Instruction:	Indicate if the patient is Chinese as determined by the patient/family.	Code System: HL7 Race
	Note(s):	Short Name: RaceChinese
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Asian - Chinese	Is Base Element: Yes
•	Having origins in any of the original peoples of China.	Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and	Data Type: BL
	Ethnicity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation

Element: 2072 Race - Asian

Operator: Equal Value: Yes







Section: Demographics	Parent: Root	
Element: 2082	Race - Filipino	Technical Specification
Coding Instruction:	Indicate if the patient is Filipino as determined by the patient/family.	Code: 2036-2 Code System: HL7 Race
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: RaceFilipino Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Asian - Filipino	Is Base Element: Yes
	Having origins in any of the original peoples of the Philippines.	Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Data Type: BL Precision:
		Selection Type: Single Unit of Measure:
		Default Value: Null Usual Range:
		Valid Range: Data Source: User
		Parent/Child Validation Element: 2072 Race - Asian Operator: Equal

Technical Specification Element: 2083 Race - Japanese Code: 2039-6 Coding Instruction: Indicate if the patient is Japanese as determined by the patient/family. Code System: HL7 Race Short Name: RaceJapanese Missing Data: Report If the patient has multiple race origins, specify them using the other race selections in addition Harvested: Yes (TAVR, TMVR, TMVrpr, to this one. Target Value: The value on arrival at this facility Is Identifier: No Is Base Element: Yes Supporting Definition: Asian - Japanese Is Followup Element: No 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 Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 2072 Race - Asian

Operator: Equal Value: Yes

Value: Yes







Section: Demographics	Parent: Root	
Element: 2084	Race - Korean	Technical Specification
Coding Instruction:	Indicate if the patient is Korean as determined by the patient/family.	Code: 2040-4 Code System: HL7 Race
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: RaceKorean Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr,
-	The value on arrival at this facility	TTVP) Is Identifier: No
Supporting Definition:	Asian - Korean Having origins in any of the original peoples of Korea.	Is Base Element: Yes Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	
		Selection Type: Single Unit of Measure:
		Default Value: Null Usual Range:
		Valid Range: Data Source: User
		Parent/Child Validation Element: 2072 Race - Asian Operator: Equal Value: Yes

Element: 2085	Race - Vietnamese	Technical Specification
	Indicate if the patient is Vietnamese as determined by the patient/family.	Code: 2047-9
Coding Instruction:		Code System: HL7 Race
	Note(s):	Short Name: RaceVietnamese
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Asian - Vietnamese	Is Base Element: Yes
	Having origins in any of the original peoples of Viet Nam.	Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and	Data Type: BL
	Ethnicity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation
		Floment: 2072 Page Asian

Element: 2072 Race - Asian

Operator: Equal Value: Yes







Section: Demographics	Parent: Root	
Element: 2086	Race - Other Asian	Technical Specification
Coding Instruction:	Indicate if the patient is of Other Asian descent as determined by the patient/family.	Code: 100001130 Code System: ACC NCDR
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: RaceAsianOther Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	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	Is Base Element: Yes Is Followup Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range:
		Valid Range: Data Source: User Parent/Child Validation Element: 2072 Race - Asian Operator: Equal Value: Yes

Element: 2074	Race - Native Hawaiian/Pacific Islander	Technical Specification
On the standard and		Code: 2076-8
Coding Instruction:	Indicate if the patient is Native Hawaiian or Pacific Islander as determined by the patient/family.	Code System: HL7 Race
	Note(s):	Short Name: RaceNatHaw
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Race - Native Hawaiian/Pacific Islander - Native Hawaiian	Is Base Element: Yes
	Having origins in any of the original peoples of Hawaii, Guam, Samoa, or other Pacific Islands.	Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and	Data Type: BL
	Ethnicity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User







Section: Demographics	Parent: Root	
Element: 2090	Race - Native Hawaiian	Technical Specification
Coding Instruction:	Indicate if the patient is Native Hawaiian as determined by the patient/family.	Code: 2079-2 Code System: HL7 Race
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: RaceNativeHawaii Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Native Hawaiian	Is Base Element: Yes
	Having origins in any of the original peoples of the islands of Hawaii.	Is Followup No Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Data Type: BL Precision:
		Selection Type: Single Unit of Measure:
		Default Value: Null Usual Range:
		Valid Range: Data Source: User
		Parent/Child Validation
		Element: 2074 Race - Native Hawaiian/Pacific Islander Operator: Equal Value: Yes

Element: 2091	Race - Guamanian or Chamorro	Technical Specification
On the selection of	Indicate if the patient is Guamanian or Chamorro as determined by the patient/family.	Code: 2086-7
Coding Instruction:		Code System: HL7 Race
	Note(s):	Short Name: RaceGuamChamorro
	If the patient has multiple race origins, specify them using the other race selections in addition	Missing Data: Report
	to this one.	Harvested: Yes (TAVR, TMVR, TMVrpr,
Target Value	The value on arrival at this facility	TTVP)
rarget value.	The value on anival at this facility	Is Identifier: No
Supporting Definition:	Native Hawaiian/Pacific Islander - Guamanian or Chamorro	Is Base Element: Yes
	Having origins in any of the original peoples of the Mariana Islands or the island of Guam.	Is Followup No
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and	Element:
	Ethnicity	Data Type: BL
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation

Parent/Child Validation

Element: 2074 Race - Native Hawaiian/Pacific

Islander

Operator: Equal Value: Yes







Section: Demographics	Parent: Root	
Element: 2092	Race - Samoan	Technical Specification
Coding Instruction:	Indicate if the patient is Samoan as determined by the patient/family.	Code: 2080-0 Code System: HL7 Race
	Note(s): If the patient has multiple race origins, specify them using the other race selections in addition to this one.	Short Name: RaceSamoan Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Native Hawaiian/Pacific Islander - Samoan	Is Base Element: Yes
	Having origins in any of the original peoples of the island of the Samoa.	Is Followup Element: No
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Data Type: BL Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range: Data Source: User
		Parent/Child Validation
		Element: 2074 Race - Native Hawaiian/Pacific
		Operator: Equal

Technical Specification Element: 2093 Race - Other Pacific Islander Code: 2500-7 Coding Instruction: Indicate if the patient is Other Pacific Islander as determined by the patient/family. Code System: HL7 Race Short Name: RacePacificIslandOther Missing Data: Report If the patient has multiple race origins, specify them using the other race selections in addition Harvested: Yes (TAVR, TMVR, TMVrpr, to this one. TTVP) Target Value: The value on arrival at this facility Is Identifier: No Is Base Element: Yes Supporting Definition: Native Hawaiian/Pacific Islander - Other Pacific Island Is Followup Element: No 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 Data Type: BL Ethnicity Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 2074 Race - Native Hawaiian/Pacific Islander

Operator: Equal
Value: Yes

Value: Yes







Section: Demographics	Parent: Root	
Element: 2076	Hispanic or Latino Ethnicity	Technical Specification
Coding Instruction:	Indicate if the patient is of Hispanic or Latino ethnicity as determined by the patient/family.	Code: 2135-2 Code System: HL7 Ethnicity
	Note(s): If the patient has multiple hispanic or latin ethnicity, specify them using the other ethnicity selections in addition to this one.	Short Name: HispOrig Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	Hispanic or Latino Ethnicity A person of 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."	Is Base Element: Yes Is Followup Element: Data Type: BL
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Precision: Selection Type: Single Unit of Measure:
		Default Value: Null Usual Range: Valid Range: Data Source: User

Element: 2100	Hispanic Ethnicity Type - Mexican, Mexican-American, Chicano	Technical Specification
Coding Instruction:	Indicate if the patient is Mexican, Mexican - American, or Chicano as determined by the patient/family.	Code: 2148-5 Code System: HL7 Ethnicity Short Name: HispEthnicityMexican
	Note(s): If the patient has multiple hispanic or latin ethnicity, specify them using the other ethnicity selections in addition to this one.	Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No Is Base Element: Yes
Supporting Definition:	Hispanic Ethnicity - Mexican/Mexican American/Chicano Having origins in any of the original peoples of Mexico.	Is Followup Element: No
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and Ethnicity	Data Type: BL Precision: Selection Type: Single
		Unit of Measure: Default Value: Null
		Usual Range: Valid Range:

Parent/Child Validation

Element: 2076 Hispanic or Latino Ethnicity

Data Source: User

Operator: Equal Value: Yes







Section: Demographics	Parent: Root	
Element: 2101	Hispanic Ethnicity Type - Puerto Rican	Technical Specification
Coding Instruction:	Indicate if the patient is Puerto Rican as determined by the patient/family.	Code: 2180-8 Code System: HL7 Ethnicity
	Note(s): If the patient has multiple hispanic or latin ethnicity, specify them using the other ethnicity selections in addition to this one.	Short Name: HispEthnicityPuertoRico Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
_	The value on arrival at this facility	Is Identifier: No
Supporting Definition:	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	Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range:
		Valid Range: Data Source: User Parent/Child Validation Element: 2076 Hispanic or Latino Ethnicity Operator: Equal Value: Yes

Element: 2102	Hispanic Ethnicity Type - Cuban	Technical Specification
On the standard on		Code: 2182-4
Coding instruction:	Indicate if the patient is Cuban as determined by the patient/family.	Code System: HL7 Ethnicity
	Note(s):	Short Name: HispEthnicityCuban
	If the patient has multiple hispanic or latin ethnicity, specify them using the other ethnicity	Missing Data: Report
	selections in addition to this one.	Harvested: Yes (TAVR, TMVR, TMVrpr,
Tarnet Value	The value on arrival at this facility	TTVP)
raiget value.	The value of arrival at this facility	Is Identifier: No
Supporting Definition:	Hispanic Ethnicity - Cuban	Is Base Element: Yes
	Having origins in any of the original peoples of Cuba.	Is Followup Element:
	Source: U.S. Office of Management and Budget. Classification of Federal Data on Race and	Data Type: BL
	Ethnicity	Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User
		Parent/Child Validation
		Flement: 2076 Hispanic or Latino Ethnicity

Element: 2076 Hispanic or Latino Ethnicity

Operator: Equal Value: Yes







Value: Yes

Section: Demographics	Parent: Root	
Element: 2103	Hispanic Ethnicity Type - Other Hispanic, Latino or Spanish Origin	Technical Specification
Coding Instruction:	Indicate if the patient is another Hispanic, Latino, or Spanish origin as determined by the patient/family.	Code: 100001131 Code System: ACC NCDR Short Name: HispEthnicityOtherOrigin
	Note(s): If the patient has multiple hispanic or latin ethnicity, specify them using the other ethnicity selections in addition to this one.	Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP)
Target Value:	The value on arrival at this facility	Is Identifier: No Is Base Element: Yes
Supporting Definition:	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	Is Followup Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User
		Parent/Child Validation
		Element: 2076 Hispanic or Latino Ethnicity Operator: Equal

Element: 14780	Original Patient ID	Technical Specification
Cadina Instruction		Code: 112000002061
Coding instruction:	This is the ID generated when the patient was first submitted to the STS/ACC TVT Registry. This field will be provided to vendors as part of the participant vendor migration process for all	Code System: ACC NCDR
	patients currently in the Registry. For patients submitted to the STS/ACC TVT Registry the first	Short Name: OrigPtID
	time by a vendor, it should be populated with the NCDR Patient ID assigned by the vendor.	Missing Data: Illegal
Target Value:		Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
		Is Identifier: No
		Is Base Element: Yes
		Is Followup Yes Element:
		Data Type: NUM
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value:
		Usual Range:
		Valid Range:

		Data Source: Automatic
Element: 14781	Original NCDR Vendor	Technical Specification
Coding Instruction:	This is the vendor identifier of the vendor who first submitted the patient to the STS/ACC TVT Registry. This field will be provided to vendors as part of the vendor migration process for all patients currently in the registry. For patients submitted to the STS/ACC TVT Registry for the first time by a vendor, it should be populated with the Vendor Identifier of the submitting vendor.	Code: 112000002062 Code System: ACC NCDR Short Name: OrigNCDRVen Missing Data: Illegal Harvested: Yes (BDS, TAVR, TMVR,
Target Value:	N/A	TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element:
		Data Type: ST Precision: 15 Selection Type: Single Unit of Measure:
		Default Value: Usual Range: Valid Range: Data Source: Automatic



Element: 2999



Full Specifications Data Dictionary v3.0



Section: Episode Information	Parent: Episode of Care

Episode Unique Key Coding Instruction: Indicate the unique key associated with each patient episode record as assigned by the

EMR/EHR or your software application.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.855

Code System: ACC NCDR Short Name: EpisodeKey Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: Yes Is Base Element: Yes Is Followup No Element: Data Type: ST Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: Automatic

Element: 3001 Arrival Date and Time

Coding Instruction: Indicate the date and time the patient arrived at your facility.

Indicate the time (hours:minutes) using the military 24-hour clock, beginning at midnight (00:00

hours).

Target Value: N/A

Vendor Instruction: Arrival Date and Time (3001) must be Less than or Equal to Procedure Start Date and Time

Arrival Date and Time (3001) must be Less than or Equal to Discharge Date (10100)

Technical Specification

Code: 1000142450 Code System: ACC NCDR Short Name: ArrivalDateTime

Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: TS Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

> Valid Range: Data Source: User

Health Insurance Element: 3005

Coding Instruction: Indicate if the patient has health insurance.

Target Value: The value on arrival at this facility

Technical Specification

Code: 63513-6 Code System: LOINC Short Name: HealthIns Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup Element: No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

> Valid Range: Data Source: User







Section: Episode Information Parent: Episode of Care

Element: 3010 Health Insurance Payment Source

Coding Instruction: Indicate the patient's health insurance payment type.

Note(s):

If the patient has multiple insurance payors, select all payors.

If there is uncertainty regarding how to identify a specific health insurance plan, please discuss with your billing department to understand how it should be identified in the registry.

Target Value: The value on arrival at this facility

Technical Specification

Code: 100001072
Code System: ACC NCDR
Short Name: HIPS
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Multiple

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 3005 Health Insurance

Operator: Equal Value: Yes

Payor Category - 1.3.6.1.4.1.19376.1.4.1.6.5.5

Selection	Definition	Source	Code	Code System
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. A health maintenance organization (HMO) is considered private health insurance.	;	5	PHDSC
Medicare Fee-For-Service	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.		1	PHDSC
Medicare Advantage			112000002025	ACC NCDR
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.		2	PHDSC
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 Departmen of Veterans Affairs), as well as care provided by the Department of Veterans Affairs (VA).	t	31	PHDSC
State-Specific Plan (non- Medicaid)	State Specific Plans - 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.		36	PHDSC
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-HIS facilities.		33	PHDSC
Non-US Insurance	Non-US insurance refers to individuals with a payor that does not originate in the United States.		100000812	ACC NCDR







Section: Episode Information Parent: Episode of Care

Technical Specification Element: 12846 Medicare Beneficiary Identifier Code: 2.16.840.1.113883.4.927 Coding Instruction: Indicate the patient's Medicare Beneficiary Identifier (MBI). Code System: Center for medicare and medicaid services, MBI Short Name: MBI Enter the Medicare Beneficiary Identifier (MBI) for those patients insured by Medicare. Patients Missing Data: Report without Medicare will not have a MBI. Harvested: Yes (BDS, TAVR, TMVR, Target Value: The value on arrival at this facility TMVrpr, TTVP) Is Identifier: No Supporting Definition: Medicare Beneficiary Identifier Is Base Element: Yes The Medicare Access and CHIP Reauthorization Act (MACRA) of 2015, requires us to remove Is Followup No Social Security Numbers (SSNs) from all Medicare cards by April 2019. A new Medicare Element: Beneficiary Identifier (MBI) will replace the SSN-based Health Insurance Claim Number (HICN) Data Type: ST on the new Medicare cards for Medicare transactions like billing, eligibility status, and claim Precision: 11 status. Selection Type: Single Source: https://www.cms.gov/Medicare/New-Medicare-Card/index.html Unit of Measure: Default Value: Null **Usual Range:**

Element: 13803 Residence

Coding Instruction: Indicate the primary residence of the patient prior to arrival. If the primary residence is not

available, code not documented.

Target Value: The value on arrival at this facility

Technical Specification

Code: 112000001506
Code System: ACC NCDR
Short Name: Residence
Missing Data: Report

Valid Range: Data Source: User

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation
Element: 13804 Residence Not Documented

Operator: Equal

Value: No (or Not Answered)

Residence - 1.3.6.1.4.1.19376.1.4.1.6.5.562

Selection	Definition	Source	Code	Code System
Home with No Health Aid	The patient lives at home with no health-aid (this includes living in senior living facilities with no assistance).		112000001507	ACC NCDR
Home with Health Aid	The patient lives at home with health-aid (this include living in senior living facilities with assistance).	5	112000001508	ACC NCDR
Long Term Care	The patient lives in a long-term care facility that provides the person's health or personal care needs during a short or long period of time.	National Institute of Aging at the National Institutes of Health	42665001	SNOMED CT
Other			100000351	ACC NCDR







Section: Episode Information Parent: Episode of Care

Element: 13804 Residence Not Documented

Coding Instruction: Indicate if the primary residence of the patient prior to arrival was not documented.

Target Value: N/A

Technical Specification
Code: 112000001506
Code System: ACC NCDR
Short Name: ResidenceND

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Usual Range: Valid Range: Data Source: User

Default Value: Null

Element: 3020 Patient Enrolled in Research Study

Coding Instruction: Indicate if the patient is enrolled in an ongoing ACC - NCDR research study related to this

registry.

Target Value: Any occurrence between arrival at this facility and discharge

Supporting Definition: Patient Enrolled in Research Study

A clinical or research study is one in which participants are assigned to receive one or more interventions (or no intervention) so that researchers can evaluate the effects of the interventions on biomedical or health-related outcomes. The assignments are determined by the study protocol. Participants may receive diagnostic, therapeutic, or other types of

interventions.

Source: Clinicaltrials.gov Glossary of Common Site Terms retrieved from http://clinicaltrials.gov/ct2/about-studies/glossary#interventional-study **Technical Specification**

Code: 100001095
Code System: ACC NCDR
Short Name: EnrolledStudy
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Default Value: Null Usual Range: Valid Range: Data Source: User

Element: 3035 Patient Restriction

Coding Instruction: Indicate if the patient requested for their information not to be used for any research or studies

for the associated episode of care.

Note(s):

Documentation must be found in the patient record to support the request of removal of their

information.

Target Value: The value on arrival at this facility

Technical Specification

Code: 100000922
Code System: ACC NCDR
Short Name: PtRestriction
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User



Element: 13171



Full Specifications Data Dictionary v3.0



Section: Episode Information Parent: Episode of Care

TVT Pathway Coding Instruction: Indicate all TVT Registry procedures performed during this episode of care.

Target Value: The value between arrival at this facility and discharge

Code: 112000001167 Code System: ACC NCDR

Technical Specification

Short Name: TVTPathway Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup Element: No Data Type: CD Precision:

Is Identifier: No

Selection Type: Multiple Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Transcatheter Valve Therapy Pathway - 1.3.6.1.4.1.19376.1.4.1.6.5.450

Selection	Definition	Source	Code	Code System
TAVR	A TVT pathway where the patient underwent a transcatheter aortic valve replacement during the current episode of care.		112000001168	ACC NCDR
TMVr	A TVT Pathway where the patient underwent a transcatheter mitral valve repair during the current episode of care.		112000001169	ACC NCDR
TMVR	A TVT Pathway where the patient underwent a transcatheter mitral valve replacement during the current episode of care.		112000001170	ACC NCDR
Tricuspid Valve Procedure	A TVT Pathway where the patient underwent a transcatheter tricuspid valve repair or replacement procedure during the current episode of care.		112000001171	ACC NCDR



Element: 3050



Full Specifications Data Dictionary v3.0



Section: Admitting Providers

Parent: Episode Information

Coding Instruction: Indicate the last name of the admitting provider.

Admitting Provider's Last Name

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: The value on arrival at this facility

Technical Specification

Code: 1000142451 Code System: ACC NCDR Short Name: AdmLName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: LN Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

> Valid Range: Data Source: User

Element: 3051 Admitting Provider's First Name

Coding Instruction: Indicate the first name of the admitting provider.

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record

Target Value: The value on arrival at this facility

Technical Specification

Code: 1000142451 Code System: ACC NCDR Short Name: AdmFName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: FN Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Element: 3052 Admitting Provider's Middle Name

Coding Instruction: Indicate the middle name of the admitting provider.

Note(s):

It is acceptable to specify the middle initial.

If there is no middle name given, leave field blank.

If there are multiple middle names, enter all of the middle names sequentially.

If the name exceeds 50 characters, enter the first 50 letters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: The value on arrival at this facility

Technical Specification

Code: 1000142451 Code System: ACC NCDR Short Name: AdmMName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes Is Followup No Element: Data Type: MN Precision: 50 Selection Type: Single

Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User







Section: Admitting Providers

Parent: Episode Information

Element: 3053 Admitting Provider's NPI

Coding Instruction: Indicate the National Provider Identifier (NPI) of the provider that admitted the patient. NPI's, assigned by the Centers for Medicare and Medicaid Services (CMS), are used to uniquely

identify physicians for Medicare billing purposes.

Note(s)

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: The value on arrival at this facility

Technical Specification

Code: 1000142451
Code System: ACC NCDR
Short Name: AdmNPI
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: NUM
Precision: 10
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User



Element: 3055



Full Specifications Data Dictionary v3.0



Section: Attending Providers

Parent: Episode Information

Attending Provider's Last Name Coding Instruction: Indicate the last name of the attending provider.

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: All values between arrival at this facility and discharge

Technical Specification

Code: 1000142452 Code System: ACC NCDR Short Name: AttLName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: LN Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User

Element: 3056 Attending Provider's First Name

Coding Instruction: Indicate the first name of the attending provider.

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record

Target Value: All values between arrival at this facility and discharge

Vendor Instruction: An Attending Provider - combination First Name (3056), Last Name (3055) and NPI (3058) - may

only be entered/selected once

Technical Specification

Code: 1000142452 Code System: ACC NCDR Short Name: AttFName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: FN Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null

Usual Range: Valid Range: Data Source: User

Element: 3057 Attending Provider's Middle Name

Coding Instruction: Indicate the middle name of the attending provider.

Note(s):

It is acceptable to specify the middle initial.

If there is no middle name given, leave field blank.

If there are multiple middle names, enter all of the middle names sequentially.

If the name exceeds 50 characters, enter the first 50 letters only.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: All values between arrival at this facility and discharge

Technical Specification

Code: 1000142452 Code System: ACC NCDR Short Name: AttMName Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes Is Followup No Element: Data Type: MN Precision: 50

Selection Type: Single Unit of Measure: Default Value: Null

> Usual Range: Valid Range: Data Source: User







Section: Attending Providers

Parent: Episode Information

Element: 3058

Attending Provider's NPI

Coding Instruction: Indicate the National Provider Identifier (NPI) of the provider that will be listed as the physician of record during the hospitalization. NPI's, assigned by the Centers for Medicare and Medicaid Services (CMS), are used to uniquely identify physicians for Medicare billing purposes.

The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical

record.

Target Value: All values between arrival at this facility and discharge

Technical Specification

Code: 1000142452 Code System: ACC NCDR Short Name: AttNPI Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: NUM Precision: 10 Selection Type: Single Unit of Measure: Default Value: Null

Usual Range: Valid Range: Data Source: User



Element: 3025



Full Specifications **Data Dictionary v3.0**



Section: Research Study Parent: Episode of Care

Coding Instruction: Indicate the research study name as provided by the research study protocol.

The research study from the research study frame as provided by the research study protector

Note(s):

Research Study Name

If the patient is in more than one research study, list each separately.

Target Value: N/A

Vendor Instruction: Research Study Name (3025) must be a valid study name for TVT 3.0

A Research Study Name (3025) may only be entered/selected once

Technical Specification

Code: 100001096 Code System: ACC NCDR Short Name: StudyName Missing Data: Report

Is Identifier: No

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: ST
Precision: 50
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation
Element: 3020 Patient Enrolled in Research Study

Operator: Equal Value: Yes

Element: 3030 Research Study Patient ID

Coding Instruction: Indicate the research study patient identification number as assigned by the research protocol.

Note(s):

If the patient is in more than one research study, list each separately.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.852

Code System: ACC NCDR Short Name: StudyPtID Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: ST
Precision: 50
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 3025 Research Study Name

Operator:

Value: Any Value







Section: History and Risk Factors

Parent: Root

Element: 6000 Height Technical Specification

Code: 8302-2
Coding Instruction: Indicate the patient's height in centimeters.

Code: 8302-2
Code System: LOINC

Short Name: Height

Target Value: The last value prior to the start of the first procedure

Short Name: Height
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Default Value: Null
Usual Range: 100.00 - 225.00 cm
Valid Range: 20.00 - 260.00 cm

Data Source: User

Element: 6005 Weight Technical Specification

Coding Instruction: Indicate the patient's weight in kilograms.

Target Value: The last value prior to the start of the first procedure

Code: 3141-9
Code System: LOINC
Short Name: Weight
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 5,2
Selection Type: Single
Unit of Measure: kg
Default Value: Null

Usual Range: 40.00 - 200.00 kg **Valid Range:** 10.00 - 700.00 kg

Data Source: User

Element: 13697 Number of Prior Open Heart Cardiac Surgeries

Coding Instruction: Indicate the number of open heart cardiac surgeries the patient has had prior to this

procedure. This includes open heart coronary artery bypass, or valve replacement/repairs.

Note: If the patient had more than 4 open heart procedures and the total number is not known,

code 4 prior open heart surgeries.

Target Value: Any occurrence between birth and start of the current procedure

Technical Specification

Code: 112000001411
Code System: ACC NCDR
Short Name: NumPrevCardSurg

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Unit of Measure:







Section: History and Risk Factors

Parent: Root

Element: 13707 Heart Failure Hospitalization Within Past Year

Coding Instruction: Indicate if the patient has been admitted to the hospital for an inpatient admission with a

diagnosis of heart failure within the past year.

Target Value: Any occurrence between 1 year prior to arrival at this facility and arrival at this facility

Heart Failure Hospitalization within Past Year Not Documented

Technical Specification

Code: 112000001855
Code System: ACC NCDR
Short Name: PriorHFAdmit1Year

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Is Identifier: No

Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Technical Specification

Element: 14253 Heart Failure Hospitalization

within Past Year Not Documented

Operator: Equal

Value: No (or Not Answered)

Valid Range: Data Source: User

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Element: 14253

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Licinciii. 14200	ricart randic ricophanzation within ract real rect becamented		
On dia a la stavetica	The Proof of Management of Antonio and Advantage of the Antonio Advanta	Code:	112000001855
Coding instruction:	Indicate if an inpatient admission with a diagnosis of heart failure within the past year was not documented.	Code System:	ACC NCDR
	documented.	Short Name:	PriorHFAdmit1YearND
Target Value:	N/A	Missing Data:	Report
		Harvested:	Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
		Is Identifier:	No
		Is Base Element:	
		Is Followup Element:	No
		Data Type:	BL
		Precision:	
		Selection Type:	Single
		Unit of Measure:	
		Default Value:	Null
		Usual Range:	







Section: History and Risk Factors

Parent: Root

Element: 13172 Anticipated Life Expectancy of Less than 1 Year

Coding Instruction: Indicate if there is physician documentation of the patient's anticipated life expectancy being

less than one year, based on comorbidities and other factors not related to the aortic stenosis

(factors that would not be expected to be favorably altered by valve replacement).

Target Value: The value on start of current procedure

Technical Specification

Code: 112000001172 Code System: ACC NCDR Short Name: LifeLessThan1yr

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User

> > Parent/Child Validation

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

----- AND -----Element: 14454 Anticipated Life Expectancy of

Less than 1 Year Not Documented

Operator: Equal

Value: No (or Not Answered)

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Element: 14454 Anticipated Life Expectancy of Less than 1 Year Not Documented

Coding Instruction: Indicate if there is no physician documentation of the patient's anticipated life expectancy being

less than one year.

Target Value: N/A

Technical Specification

Code: 112000001172 Code System: ACC NCDR Short Name: LifeLessThan1yrND

Missing Data: Report Harvested: Yes (TAVR) Is Identifier: No Is Base Element: Yes

Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Parent/Child Validation

Element: 13171 TVT Pathway Operator: Equal Value: TAVR







Section: History and Risk Factors Parent: Root

Element: 13881 Oxygen at Home

Coding Instruction: Indicate whether patient uses supplemental oxygen at home.

Target Value: The value on arrival at this facility

Technical Specification

Code: 268512000
Code System: SNOMED CT
Short Name: HmO2
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null

Usual Range: Valid Range: Data Source: User

Element: 13882 Immunocompromise Present

Coding Instruction: Indicate whether immunocompromise is present due to immunosuppressive medication therapy

or an existing medical condition. This includes, but is not limited to systemic steroid therapy, anti-rejection medications and chemotherapy. This does not include topical steroid applications, one time systemic therapy, inhaled steroid therapy or preprocedure protocol.

Target Value: The last value on start of the first procedure

Technical Specification

Code: 370388006
Code System: SNOMED CT
Short Name: ImmSupp
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Element: 13880 Currently on Dialysis

Coding Instruction: Indicate if the patient is currently undergoing either hemodialysis or peritoneal dialysis on an

ongoing basis as a result of renal failure.

Note(s):

If a patient is receiving continuous veno-venous hemofiltration (CVVH) as a result of renal

failure (and not as treatment to remove fluid for heart failure), code 'Yes'.

Target Value: The last value on start of the first procedure

Technical Specification

Code: 108241001
Code System: SNOMED CT
Short Name: CurrentlyonDialysis

Missing Data: Report

 $\textbf{Harvested:} \ \ \text{Yes (BDS, TAVR, TMVR,} \\$

TMVrpr, TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User



Element: 4625



Full Specifications Data Dictionary v3.0



Section: History and Risk Factors

Parent: Root

Coding Instruction: Indicate the frequency that the patient uses tobacco.

Note(s): Consider use of any tobacco product as equivalent to a cigarette for referenced

definitions.

Target Value: The value on arrival at this facility

Tobacco Use

Technical Specification Code: 110483000 Code System: SNOMED CT Short Name: TobaccoUse

Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Is Identifier: No

Valid Range: Data Source: User

Tobacco Use - 1.3.6.1.4.1.19376.1.4.1.6.5.427

Selection	Definition	Source	Code	Code System
Never	An individual who has not smoked 100 or more cigarettes during his/her lifetime.	The Office of the National Coordinator for Health Information Technology 2014 Edition Test Procedure for §170.314.a.11.Smoking status	266919005	SNOMED CT
Former	An individual who has smoked at least 100 cigarettes during his/her lifetime but does not currently smoke.		8517006	SNOMED CT
Current - Every Day	An individual who has smoked at least 100 cigarettes during his/her lifetime and still regularly smokes every day.		449868002	SNOMED CT
Current - Some Days	An individual who has smoked at least 100 cigarettes during his/her lifetime and still regularly smokes periodically (not every day), yet consistently.		428041000124106	SNOMED CT
Smoker - Current Status Unknown	An individual known to have smoked at least 100 cigarettes in the past, but whether they currently still smoke is unknown.		77176002	SNOMED CT
Unknown if ever smoked	An individual whose current and prior smoking status is not known.		266927001	SNOMED CT







Section: History and Risk Factors

Parent: Root

Element: 4626 Tobacco Type

Coding Instruction: Indicate all the tobacco type(s) reported by the patient.

Target Value: The value on arrival at this facility

Technical Specification

Code: 266918002
Code System: SNOMED CT
Short Name: TobaccoType
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

 Is Base Element:
 Yes

 Is Followup Element:
 No

 Data Type:
 CD

 Precision:
 Multiple

Is Identifier: No

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 4625 Tobacco Use

Operator: Equal

Value: Current - Every Day
Element: 4625 Tobacco Use

Operator: Equal

Value: Current - Some Days Element: 4625 Tobacco Use

Operator: Equal

Value: Smoker - Current Status Unknown

Tobacco Type

. obaooo . ypc				
Selection	Definition	Source	Code	Code System
Cigarettes			65568007	SNOMED CT
Cigars			59978006	SNOMED CT
Pipe			82302008	SNOMED CT
Smokeless			713914004	SNOMED CT







Section: History and Risk Factors

Parent: Root

Element: 4627 Smoking Amount

Coding Instruction: Indicate the amount of cigarette smoking reported by the patient.

Target Value: The value on arrival at this facility

Technical Specification

Code: 100001256
Code System: ACC NCDR
Short Name: SmokeAmount
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 4625 Tobacco Use

Operator: Equal

Value: Current - Every Day

----- AND -----Element: 4626 Tobacco Type

Operator: Equal Value: Cigarettes

Tobacco Amount - 1.3.6.1.4.1.19376.1.4.1.6.5.457

Selection	Definition	Source	Code	Code System
Light tobacco use (<10/day)	The patient smokes less than 10 cigarettes daily.		428061000124105	SNOMED CT
Heavy tobacco use (>= 10/day)	The patient smokes 10 or more cigarettes daily.		428071000124103	SNOMED CT







Section: Home Medications

Parent: History and Risk Factors

Element: 12297 Home Medication Code

Coding Instruction: Indicate the medication the patient has been taking routinely at home prior to this hospitalization.

Target Value: N/A

Vendor Instruction: When a Home Medication Code (12297) is selected then Home Medication Prescribed (13903)

must not be Null

Technical Specification

Code: 100013057 Code System: ACC NCDR Short Name: HomeMeds Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP)

| Is Identifier: No | Is Base Element: Yes | Is Followup | Element: No | Data Type: CD | Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13171 TVT Pathway

Operator: Equal Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure

Home Medications - 2.16.840.1.113883.3.3478.6.5.302

Selection	Definition	Source	Code	Code System
Angiotensin Convertin Enzyme Inhibitor	g		41549009	SNOMED CT
Aldosterone Antagoni	st		372603003	SNOMED CT
Angiotensin Receptor- Neprilysin Inhibitor	-		112000001832	ACC NCDR
Anticoagulant			112000001416	ACC NCDR
Aspirin			1191	RxNorm
Angiotensin II Recepto	or Blocker		372913009	SNOMED CT
Beta Blocker			33252009	SNOMED CT
Diuretics Not Otherwis Specified	se		112000001417	ACC NCDR
Loop Diuretics			29051009	SNOMED CT
Thiazides			372747003	SNOMED CT
P2Y12 Antagonist			112000001003	ACC NCDR
Selective Sinus Node Channel Inhibitor	l/f		112000001831	ACC NCDR







Section: Home Medications

Parent: History and Risk Factors

Element: 13903 Home Medication Prescribed

Coding Instruction: Indicate whether the patient received the medication at home prior to this hospitalization.

Target Value: The value on arrival at this facility

Technical Specification Code: 33633005

Code System: SNOMED CT
Short Name: PriorMedAdmin_Hom

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 12297 Home Medication Code

Operator:

Value: Any Value

Value: Loop Diuretics

Operator: Equal Value: Yes

Element: 13903 Home Medication Prescribed

Home Medication Prescribed - 1.3.6.1.4.1.19376.1.4.1.6.5.710

Selection	Definition	Source	Code	Code System
Yes			100001247	ACC NCDR
Not Prescribed - No R	Reason		100001048	ACC NCDR

Element: 14575	Loop Diuretic Dose	Technical Specification
Coding Instruction:	Specify the total daily dose of the loop diuretic the patient was taking routinely at home prior to this hospitalization.	Code: 112000001975 Code System: ACC NCDR Short Name: HomeMed_LoopDiureticDose
Target Value:	The value on arrival at this facility	Missing Data: Report Harvested: Yes (TMVR, TMVrpr, TTVP)
		Is Identifier: No Is Base Element: Yes Is Followup Element:
		Data Type: PQ Precision: 3,0
		Selection Type: Single Unit of Measure: mg Default Value: Null
		Usual Range: 1 - 40 mg Valid Range: 1 - 300 mg Data Source: User
		Parent/Child Validation Element: 12297 Home Medication Code
		Operator: Equal



Element: 12903



Full Specifications Data Dictionary v3.0



Section: Condition History Parent: History and Risk Factors

Coding Instruction: The list of medical conditions from which the patient's history is to be determined.

Condition History Name

Target Value: N/A

Technical Specification Code: 312850006 Code System: SNOMED CT

Short Name: ConditionHx Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null

Is Identifier: No

Usual Range: Valid Range: Data Source: User

Condition History Name - 1 3 6 1 4 1 19376 1 4 1 6 5 340

Selection	Definition	Source	Code	Code System
Atrial Fibrillation	AF is a supraventricular tachyarrhythmia with uncoordinated atrial activation and consequently ineffective atrial contraction. Characteristics on an electrocardiogram (ECG) include: 1) irregular R-R intervals (when atrioventricular [AV] conduction is present),	January CT, Wann LS, et al. 2014 AHA/ACC/HRS Guideline for the Management of Patients With Atrial Fibrillation: A Report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines and the Heart Rhythm Society. JACC Vol 64, #21, 2014.	49436004	SNOMED CT
	 absence of distinct repeating P waves, and 3) irregular atrial activity. 			
Atrial Flutter	and don'ny.		5370000	SNOMED CT
Cardiomyopathy			85898001	SNOMED CT
Carotid Artery Stenosis	When one or both carotid arteries was determined from any diagnostic test to have >= 50% stenosis.	Society for Thoracic Surgeons (STS)	64586002	SNOMED CT
Cerebrovascular Accident	An acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction, where the neurological dysfunction lasts for greater than 24 hours.	Society for Thoracic Surgeons (STS)	230690007	SNOMED CT
Cerebrovascular Disease	Cerebrovascular disease includes any of the following: A. Stroke: Stroke is an acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction, where the neurological dysfunction lasts for greater than 24 hours. B. TIA: is defined as a transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours. C. Noninvasive or invasive arterial imaging test demonstrating >=50% stenosis of any of the major extracranial or intracranial vessels to the brain. D. Vertebral artery and internal carotid and intercranial consistent with atherosclerotic disease with document presence as CVD. External carotid disease is excluded. E. Previous cervical or cerebral artery revascularization surgery or percutaneous intervention. F. Brain/cerebral aneurysm. G. Occlusion of veterbral artery, internal carotid artery, and intercranial due to dissection. Note: This does not include chronic (nonvascular) neurological diseases or other acute neurological insults such as metabolic and anoxic ischemic encephalopathy. Subdural hematoma or AVM is not	Society for Thoracic Surgeons (STS)	62914000	SNOMED CT
Chronic Lung Disease	cerebral vascular disease. Chronic lung disease can include patients with	ACC/AHA Key Data Elements and Definitions for Measuring the	413839001	SNOMED CT

chronic obstructive pulmonary disease, chronic

Clinical Management and Outcomes of Patients With Chronic







Section: Condition	History	Parent: History and Risk Factors		
	bronchitis, or emphysema. It can also include a patient who is currently being chronically treated with inhaled or oral pharmacological therapy (e.g., beta-adrenergic agonist, anti-inflammatory agent, leukotriene receptor antagonist, or steroid). Patients with asthma or seasonal allergies are not considered to have chronic lung disease.	Heart Failure Circulation. 2005;112:1888-1916		
Conduction Defect	Conduction disorder as evidenced by a right or left bundle branch block, sick sinus syndrome, or first, second or third degree heart block on ECG.		44808001	SNOMED C
COVID-19 Positive	The patient had a diagnosis of COVID-19, confirmed with a laboratory performed polymerase chain reaction (PCR) test.		112000001982	ACC NCDF
	Notes: It is acceptable to code the diagnosis of COVID-19 based on testing that was NOT performed at your facility if it is accompanied by appropriate documentation in the medical record by the provider.			
	Code no if documentation ONLY included antibody testing (IgG).			
Dementia - Moderate to Severe	Patients with moderate dementia (also termed moderate or severe cognitive decline) are typically oriented to person but not place and time. They are patients who need assistance with activities of daily living.	/	112000001493	ACC NCDF
Diabetes Mellitus		American Diabetes Association Care. 2017;40 Suppl 1:S13.	73211009	SNOMED CT
	1. FPG >=126 mg/dL (7.0 mmol/L). Fasting is defined as no caloric intake for at least 8 h. OR 2. 2-h PG >=200 mg/dL (11.1 mmol/L) during an OGTT. The test should be performed as described by the WHO, using a glucose load containing the equivalent of 75 g anhydrous glucose dissolved in water. OR 3. A1C >=6.5% (48 mmol/mol). The test should be performed in a laboratory using a method that is NGSP certified and standardized to the DCCT assay OR 4. In a patient with classic symptoms of hyperglycemia or hyperglycemic crisis, a random plasma glucose >=200 mg/dL (11.1 mmol/L). Endocarditis must meet the current CDC definition:		56819008	SNOMED C
Endocarditis	Endocarditis must meet at least 1 of the following criteria: 1. Patient has organisms cultured from valve or vegetation. 2. Patient has 2 or more of the following signs or symptoms: fever (>38°C), new or changing murmur*, embolic phenomena*, skin manifestations* (i.e., petechiae, splinter hemorrhages, painful subcutaneous nodules), congestive heart failure*, or cardiac conduction abnormality*		333.333	G. G.II.
	* With no other recognized cause and at least 1 of the following: 1) Organisms cultured from 2 or more blood cultures 2) Organisms seen on Gram's stain of valve when culture is negative or not done 3) Valvular vegetation seen during an invasive procedure or autopsy 4) Positive laboratory test on blood or urine (e.g., antigen tests for H influenzae, S pneumoniae, N meningitidis, or Group B Streptococcus) 5) Evidence of new vegetation seen on echocardiogram and if diagnosis is made antemortem, physician institutes appropriate antimicrobial therapy.			
	Notes: 1. Choose "Yes" for patients with pre-operative endocarditis who begin antibiotics post-op. 2. Code "Yes" for patients who are diagnosed			







Section: Condition I	History	Parent: History and Risk Factors		
	intraoperatively. 3. Marantic Endocarditis (Nonbacterial Thrombotic Endocarditis) (Lupus) should not be coded as infectious endocarditis.			
Heart Failure	Heart failure is a complex clinical syndrome that results from any structural or functional impairment of ventricular filling or ejection of blood. The cardinal manifestations of HF are dyspnea and fatigue, which may limit exercise tolerance, and fluid retention, which may lead to pulmonary and/or splanchnic congestion and/or peripheral edema. Some patients have exercise intolerance but little evidence of fluid retention, whereas others complain primarily of edema, dyspnea, or fatigue. Because some patients present without signs or symptoms of volume overload, the term "heart failure" is preferred over "congestive heart failure." There is no single diagnostic test for HF because it is largely a clinical diagnosis based on a careful history and physical examination.		84114007	SNOMED CT
Hostile Chest	A medical condition that precludes an open chest procedure and that is documented in the medical record. This can include any of the following or other reasons that make redo operation through sternotomy or right anterior thoracotomy prohibitively hazardous: 1. Evidence of abnormal chest wall anatomy due to severe kyphoscoliosis or other skeletal abnormalities (including thoracoplasty, Potts' disease, sternal bone destruction, evidence of indetectable plane between posterior sternal table and important mediastinal structures) 2. Complications from prior surgery 3. Prior radiation involving the mediastinum/thoracic, or evidence of severe radiation damage (e.g., skin burns, bone destruction, muscle loss, lung fibrosis or esophageal stricture) 4. History of multiple recurrent pleural effusions causing internal adhesions. 5. Chronic, ongoing open skin defects or extremely severe soft tissue atrophy. 6. Complete absence of reconstructive options based on plastic surgeon consult.		112000001489	ACC NCDF
Hypertension	Hypertension is defined by any one of the following 1. Documentation of hypertension as a medical problem OR	Derived from: 2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. J Am Coll Cardiol 2018;71:e127-e248.	38341003	SNOMED CT
Liver Disease	A history of hepatitis B, hepatitis C, drug induced hepatitis, autoimmune hepatitis, cirrhosis, portal hypertension, esophageal varices, liver transplant, or congestive hepatopathy. Exclude NASH in the absence of cirrhosis.	Society for Thoracic Surgeons (STS)	235856003	SNOMED CT
Myocardial Infarction	Prior myocardial infarction is defined by any of the following: 1. Documentation of myocardial infarction (MI) as a medical problem. OR 2. Any one of the following criteria meets the diagnosis for prior (sometimes called silent/unrecognized) MI: a. Abnormal Q waves with or without symptoms in the absence of nonischemic causes. b. Imaging evidence of loss of viable myocardium in a pattern consistent with ischemic etiology. c. Patho-anatomical findings of a prior MI.	Thygesen, K, Alpert, J.S., et al Fourth Universal Definition of Myocardial Infarction (2018), J Am Coll Cardiol. 2018 Oct 30;72 (18):2231-2264	22298006	SNOMED C1
Peripheral Arterial Disease		ACCF/AHA 2011 Key Data Elements and Definitions of a Base Cardiovascular Vocabulary for Electronic Health Records (JACC, 2011;58;202-222)	399957001	SNOMED CT







Section: Condition History		Parent: History and Risk Fa	ctors	
	percutaneous revascularization in the arteries of th extremities * Positive noninvasive test (e.g., ankle brachial inde <= 0.9, ultrasound, MR or CT imaging of >50% diameter stenosis in any peripheral artery (i.e., subclavian, femoral, iliac) or angiographic imaging)			
Porcelain Aorta	A porcelain aorta is defined as "severe atherosclerosis of the aorta, calcification may be severe and diffuse, causing an eggshell appearance seen on chest x-ray or CT".	ACCF/AHA/AATS/ACR/ASA/SCA/SCAI/SIR/STS for the Diagnosis and Management of Patients W Aortic Disease (JACC, 2010; 55:27-129)		ACC NCDR
Transient Ischemic Attack (TIA)	A transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.	Society for Thoracic Surgeons (STS)	266257000	SNOMED CT
Element: 14264	Condition History Occurrence		Technical Specific	ation
_	tion: Indicate if the patient does or does not have a	•	Code: 312850006 Code System: SNOMED CT Short Name: ConditionHxC)ccurence

Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User Parent/Child Validation Element: 12903 Condition History Name

Operator:







Section: Condition History

Parent: History and Risk Factors

Element: 14251 Condition History Date

Coding Instruction: Indicate the most recent occurrence date for the condition.

Note(s):

If the month or day of the diagnosis is unknown, please code 01/01/YYYY. If the specific year is unknown in the current record, the year may be estimated based on timeframes found in prior medical record documentation (Example: If the patient had "most recent diagnosis" documented in a record from 2011, then the year 2011 can be utilized and coded as

01/01/2011).

Target Value: The last value between birth and the first procedure in this admission

Vendor Instruction: Condition History Date (14251) must be Less than or Equal to Procedure Start Date and Time

(7000)

Technical Specification

Code: 312850006
Code System: SNOMED CT
Short Name: CondHistDate
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: DT
Precision:
Selection Type: Single

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Cerebrovascular Accident Element: 12903 Condition History Name

Operator: Equal

Value: COVID-19 Positive

Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes







Section: Atrial Fibrillation Parent: Condition History Details

Element: 13179 Atrial Fibrillation Classification

Coding Instruction: Indicate the classification of atrial fibrillation.

Target Value: The last value within 30 days prior to the first procedure in this admission

Technical Specification

Code: 100000935
Code System: ACC NCDR
Short Name: AFibClassification

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Atrial Fibrillation

Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Atrial Fibrillation Classification - 1.3.6.1.4.1.19376.1.4.1.6.5.17

Selection	Definition	Source	Code	Code System
Paroxysmal	AF that terminates spontaneously or with interventior within 7 days of onset. Episodes may recur with variable frequency.		26593000	SNOMED CT
Persistent	Continuous AF that is sustained >7 days or with electrical or pharmacological termination.		62459000	SNOMED CT
Long-standing Persistent	Continuous AF of >12 months duration.		100001029	ACC NCDR
Permanent	The term "permanent AF" is used when the patient ar clinician make a joint decision to stop further attempts restore and/or maintain sinus rhythm.		6934004	SNOMED CT
	 Acceptance of AF represents a therapeutic attitude on the part of the patient and clinician rather than an inherent pathophysiological attribute of the AF. 			
	 Acceptance of AF may change as symptoms, the efficacy of therapeutic interventions, and patient and clinician preferences evolve. 			
None			100001231	ACC NCDR







Section: Atrial Fibrillation

Parent: Condition History Details

Element: 14244 Recent Atrial Fibrillation

Coding Instruction: Indicate if the patient has had atrial fibrillation within the past 30 days.

Target Value: Any occurrence between 30 days prior to the procedure and the procedure

Technical Specification

Code: 112000001790
Code System: ACC NCDR
Short Name: AFib30days
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13179 Atrial Fibrillation Classification

Operator: Equal

Value: Paroxysmal

Element: 13179 Atrial Fibrillation Classification

Operator: Equal

Value: Persistent



Element: 14245



Full Specifications **Data Dictionary v3.0**



Section: Atrial Flutter Parent: Condition History Details

Coding Instruction: Indicate if the patient has had atrial flutter within the past 30 days.

Recent Atrial Flutter

Target Value: Any occurrence between 30 days prior to the procedure and the procedure

Code: 112000001791
Code System: ACC NCDR
Short Name: AFlutter30days
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

Technical Specification

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Atrial Flutter

Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes







Section: Carotid Artery Stenosis

Parent: Condition History Details

Element: 14265 **Current Carotid Artery Stenosis**

Coding Instruction: Indicate if the patient has carotid artery stenosis.

Target Value: The value on arrival at this facility

Supporting Definition: Carotid Artery Stenosis

A narrowing of the carotid artery lumen. It is usually caused by the formation of an atherosclerotic plaque. Symptoms are usually present when there is severe narrowing or obstruction of the arterial lumen and manifest as ischemic cerebrovascular accidents.

Source: NCImetathesaurus NCIm Version: 201706 Version 2.8

CUI C0007282

Technical Specification

Code: 64586002 Code System: SNOMED CT Short Name: CurrendCAS Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Carotid Artery Stenosis

----- AND -----Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Element: 14230 Carotid Artery Stenosis Location

Coding Instruction: Indicate which carotid artery was determined from any diagnostic test to be greater or equal to

Target Value: The last value prior to the start of the first procedure

Supporting Definition: Carotid Artery Stenosis

A narrowing of the carotid artery lumen. It is usually caused by the formation of an atherosclerotic plaque. Symptoms are usually present when there is severe narrowing or obstruction of the arterial lumen and manifest as ischemic cerebrovascular accidents.

Source: NCImetathesaurus NCIm Version: 201706 Version 2.8

CUI C0007282

Technical Specification

Code: 112000002012 Code System: ACC NCDR Short Name: CVDCarsten Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP) Is Identifier: No

Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 14265 Current Carotid Artery Stenosis

Operator: Equal Value: Yes

AND

Element: 14329 Carotid Artery Stenosis Location

Not Documented

Operator: Equal Value: No (or Not Answered)

Carotid Artery Stenosis Location - 1.3.6.1.4.1.19376.1.4.1.6.5.684

Selection	Definition	Source	Code	Code System
Right Carotid Artery Stenosis	There is >=50% stenosis in the right carotid artery.		285201000119100	SNOMED CT
Left Carotid Artery Stenosis	There is >=50% stenosis in the left carotid artery.		285191000119103	SNOMED CT
Bilateral Carotid Artery	There is >=50% stenosis in both the right carotid and		293821000119107	SNOMED CT
Stenosis	left carotid arteries.			







Section: Carotid Artery Stenosis

Parent: Condition History Details

Element: 14329 Carotid Artery Stenosis Location Not Documented

Coding Instruction: Indicate if the severity of carotid artery stenosis was not documented.

Target Value: N/A

Supporting Definition: Carotid Artery Stenosis

A narrowing of the carotid artery lumen. It is usually caused by the formation of an atherosclerotic plaque. Symptoms are usually present when there is severe narrowing or obstruction of the arterial lumen and manifest as ischemic cerebrovascular accidents.

Source: NCImetathesaurus NCIm Version: 201706 Version 2.8

CUI C0007282

Technical Specification

Code: 112000002012
Code System: ACC NCDR
Short Name: CVDCarSteLocND

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14265 Current Carotid Artery Stenosis

Operator: Equal Value: Yes



Element: 4570



Full Specifications Data Dictionary v3.0



Section: Cardiomyopathy **Parent: Condition History Details**

Cardiomyopathy Type Coding Instruction: Indicate the type of cardiomyopathy experienced by the patient.

If the patient has had multiple cardiomyopathies, select all applicable types.

Target Value: Any occurrence between birth and the procedure

Technical Specification

Code: 100000953 Code System: ACC NCDR Short Name: PriorCMType Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Multiple

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Cardiomyopathy

----- AND -----Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

AND

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 13171 TVT Pathway

Operator: Equal Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure

Cardiomyopathy Type - 1.3.6.1.4.1.19376.1.4.1.6.5.193

Selection	Definition	Source Code	Code System
Ischemic cardiomyopathy	The patient has a history of ischemic cardiomyopathy documented by heart failure and reduced systolic function (ejection fraction <40%) and history of any one of the following: 1. History of myocardial infarction (MI) 2. History of Percutaneous Coronary Intervention; 3. History of Coronary Artery Bypass Graft Surgery; 4. Conventional coronary angiography demonstrates >=70% stenosis in at least one major coronary artery. 5. Stress testing (with or without imaging) diagnostic coronary artery disease.		SNOMED CT
Non-ischemic cardiomyopathy	Includes cardiomyopathies resulting from volume or pressure overload, such as hypertension or valvular heart disease.	111000119104	SNOMED CT
Other Cardiomyopathy Type	Cardiomyopathy not otherwise specified.	100001065	ACC NCDR



Element: 13904



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Section: Chronic Lung Disease

Parent: Condition History Details

Coding Instruction: Indicate the severity of chronic lung disease.

Target Value: The last value between birth and the first procedure in this admission

Chronic Lung Disease Severity

Supporting Definition: Chronic Lung Disease

Chronic lung disease can include patients with chronic obstructive pulmonary disease, chronic bronchitis, or emphysema. It can also include a patient who is currently being chronically treated with inhaled or oral pharmacological therapy (e.g., beta-adrenergic agonist, antiinflammatory agent, leukotriene receptor antagonist, or steroid). Patients with asthma or seasonal allergies are not considered to have chronic lung disease.

Source: ACC/AHA Key Data Elements and Definitions for Measuring the Clinical Management and Outcomes of Patients With Chronic Heart Failure Circulation. 2005;112:1888-1916

Technical Specification

Code: 413839001 Code System: SNOMED CT

Short Name: ChronLungDisSeverity

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD

Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14459 Chronic Lung Disease Severity

Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----Element: 12903 Condition History Name

Operator: Equal

Value: Chronic Lung Disease

----- AND -----Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Chronic Lung Disease Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.585

Selection	Definition	Source	Code	Code System
Mild Lung Disease	FEV1 60% to 75% of predicted, and/or on chronic inhaled or oral bronchodilator therapy.	Society of Thoracic Surgeons (STS)	112000001593	ACC NCDR
Moderate Lung Disease	FEV1 50% to 59% of predicted, and/or on chronic steroid therapy aimed at lung disease.	Society of Thoracic Surgeons (STS)	112000001594	ACC NCDR
Severe Lung Disease	FEV1 <50% predicted, and/or Room Air pO2 < 60 or Room Air pCO2 > 50.	Society of Thoracic Surgeons (STS)	112000001595	ACC NCDR







Section: Chronic Lung Disease

Parent: Condition History Details

Element: 14459 Chronic Lung Disease Severity Not Documented

Coding Instruction: Indicate true if the severity of chronic lung disease is not documented.

Target Value: N/A

Technical Specification

Code: 112000001596 Code System: ACC NCDR

Short Name: ChronLungDisSeverity_ND

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Chronic Lung Disease

AND ----
Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes







Section: Diabetes Therapy

Parent: Condition History Details

Element: 14231 Diabetes Therapy

Coding Instruction: Indicate the type of treatment a patient with a diagnosis of diabetes is receiving. Indicate the

most aggressive therapy the patient presented with on admission.

Target Value: The last value between birth and the first procedure in this admission

Code: 385804009
Code System: SNOMED CT

Short Name: DiabControl Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Technical Specification

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Is Identifier: No

nit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Diabetes Mellitus

AND ----Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Diabetes Therapy

Definition	Source	Code	Code System	
		112000000322	ACC NCDR	
		112000000324	ACC NCDR	
		112000000323	ACC NCDR	
		161649006	SNOMED CT	
		112000000325	ACC NCDR	
	Definition	Definition Source	112000000322 11200000324 11200000323 161649006	







Section: Endocarditis Parent: Condition History Details

Element: 14232 Endocarditis Type

Coding Instruction: Indicate the type of endocarditis.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 56819008
Code System: SNOMED CT
Short Name: InfEndTy
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No
Is Base Element: Yes

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Endocarditis

Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Endocarditis Type - 1.3.6.1.4.1.19376.1.4.1.6.5.685

Selection	Definition	Source	Code	Code System
Treated Endocarditis	The patient has been treated previously for endocarditis and is not taking antibiotics for the infection (other than prophylactic medications). The patient is currently being treated for endocarditis.		112000001752	ACC NCDR
Active Endocarditis	The patient is currently being treated for endocarditis. This includes patients who are diagnosed and treatment begins post-op.		112000001753	ACC NCDR







Section: Myocardial Infarction

Parent: Condition History Details

Element: 13174 Myocardial Infarction Timeframe

Coding Instruction: Indicate if the timeframe of the myocardial infarction.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 22298006
Code System: SNOMED CT
Short Name: MIWhen
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 12903 Condition History Name

Operator: Equal

Value: Myocardial Infarction

Element: 14264 Condition History Occurrence

Operator: Equal Value: Yes

Prior Myocardial Infarction Timeframe - 1.3.6.1.4.1.19376.1.4.1.6.5.451

Selection	Definition	Source	Code	Code System
Prior Myocardial Infarction	Prior myocardial infarction is less than 30 days prior to	1	112000001173	ACC NCDR
Less than 30 days	the procedure.			
Prior Myocardial Infarction			112000001174	ACC NCDR
Greater than or Equal to 30				
days				



Element: 12905



Full Specifications **Data Dictionary v3.0**



Section: Procedure History

Parent: History and Risk Factors

Coding Instruction: The list of medical procedures from which the patient's history is to be determined.

Target Value: N/A

Vendor Instruction: When a Procedure History Name (12905) is selected then Procedure History Occurrence

(14268) must not be Null

Procedure History Name

Technical Specification
Code: 416940007
Code System: SNOMED CT
Short Name: ProcedHxName

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Procedure History Name - 1.3.6.1.4.1.19376.1.4.1.6.5.341

Selection	Definition	Source	Code	Code System
Aortic Valve Procedure	Any previous surgical or interventional replacement and/or repair of the aortic valve.		112000001755	ACC NCDR
Aortic Valve Balloon Valvuloplasty			77166000	SNOMED CT
Aortic Valve Repair Surgery			112816004	SNOMED CT
Aortic Valve Replacement Surgery			725351001	SNOMED CT
Aortic Valve Replacement - Transcatheter			41873006	SNOMED CT
Aortic Valve Transcatheter Intervention	Any previous interventional repair of the aortic valve. Note: Do not include surgical aortic valve repairs or transcatheter aortic valve replacements.		112000001768	ACC NCDR
Coronary Artery Bypass Graft			232717009	SNOMED CT
Implantable Cardioverter Defibrillator	Placement of an internal cardioverter defibrillator.		447365002	SNOMED CT
Mitral Valve Procedure	Any previous surgical or interventional replacement and/or repair of the mitral valve.		112000001940	ACC NCDR
Mitral Valve Annuloplasty Ring Surgery			232744004	SNOMED CT
Mitral Valve Repair Surgery			384641003	SNOMED CT
Mitral Valve Replacement Surgery			53059001	SNOMED CT
Mitral Valve Transcatheter Intervention	Any previous interventional repair of the mitral valve. Note: Do not include surgical mitral valve repairs or transcatheter mitral valve replacements.		112000001773	ACC NCDR
PCI			415070008	SNOMED CT
Permanent Pacemaker			449397007	SNOMED CT
Pulmonic Valve Procedure	Any previous surgical or interventional replacement and/or repair of the pulmonic valve.		112000001769	ACC NCDR
Tricuspid Valve Procedure	Any previous surgical or interventional replacement and/or repair of the tricuspid valve.		112000001941	ACC NCDR
Tricuspid Valve Repair Surgery			384643000	SNOMED CT
Tricuspid Valve Replacement Surgery			25236004	SNOMED CT
Tricuspid Valve Replacement - Transcatheter			112000001977	ACC NCDR
Tricuspid Valve Transcatheter Intervention	Any previous interventional repair of the tricuspid valve. Note: Do not include surgical tricuspid valve repairs or transcatheter tricuspid valve replacements		112000001779	ACC NCDR







Section: Procedure History

Parent: History and Risk Factors

Element: 14268 Procedure History Occurrence

Coding Instruction: Indicate if the patient does or does not have a history of the indicated medical procedure.

Target Value: Any occurrence between birth and the first procedure in this admission

Technical Specification

Code: 416940007 Code System: SNOMED CT Short Name: ProcHxOccur Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation Element: 12905 Procedure History Name

Operator:

Value: Any Value

Element: 14252 Procedure History Date

Coding Instruction: Indicate the date the procedure was performed.

Note(s): If the month or day of the procedure is unknown, please code 01/01/YYYY. If the specific year is unknown in the current record, the year may be estimated based on timeframes found in prior medical record documentation (Example: If the patient had "most recent procedure" documented in a record from 2011, then the year 2011 can be utilized and

coded as 01/01/2011).

Target Value: The last value between birth and the first procedure in this admission

Vendor Instruction: Procedure History Date (14252) must be Less than or Equal to Procedure Start Date and Time

Technical Specification

Code: 416940007 Code System: SNOMED CT Short Name: ProcHistDate Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: DT Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Is Identifier: No

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Aortic Valve Procedure

Data Source: User

Element: 12905 Procedure History Name

Operator: Equal

Value: Coronary Artery Bypass Graft Element: 12905 Procedure History Name

Operator: Equal

Value: Permanent Pacemaker

Element: 12905 Procedure History Name

Operator: Equal Value: PCI

Element: 12905 Procedure History Name

Operator: Equal

Value: Mitral Valve Procedure

Element: 12905 Procedure History Name

Operator: Equal

Value: Tricuspid Valve Procedure ----- AND -----

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes







Section: Aortic Valve Replacement

Parent: Procedure History Details

Element: 14335 Surgical Aortic Valve Replacement Implant ID

Coding Instruction: Indicate the implant ID of the prosthetic aortic valve.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 84683006
Code System: SNOMED CT
Short Name: SAVRImplantID
Missing Data: Report

Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Aortic Valve Replacement Surgery

Element: 14268 Procedure History Occurrence

----- AND -----

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

Element: 14519 Surgical Aortic Valve Replacement Implant Diameter

Coding Instruction: Indicate the aortic valve implant size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 84683006
Code System: SNOMED CT
Short Name: SAVRImplantDia

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:

Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 16 - 36 mm
Valid Range: 5 - 100 mm
Data Source: User

Parent/Child Validation

Element: 14335 Surgical Aortic Valve

Replacement Implant ID

Operator:

Value: Any Value



Element: 14236



Full Specifications Data Dictionary v3.0



Section: Aortic Valve Replacement

Parent: Procedure History Details

Aortic Valve Replacement Type

Coding Instruction: Indicate the type of surgical aortic valve replacement.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 725351001 Code System: SNOMED CT Short Name: PrevProcAVType

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD

Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Parent/Child Validation

Element: 14237 Aortic Valve Replacement Type

Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----Element: 12905 Procedure History Name

Operator: Equal

Value: Aortic Valve Replacement Surgery

AND Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

Aortic Valve Replacement Type - 1.3.6.1.4.1.19376.1.4.1.6.5.686

Selection	Definition	Source Co	de Code System
Stented Valve Replacement	Surgical valve replacement with a bioprosthetic stente valve.	d 112000001	758 ACC NCDR
Stentless Valve Replacement	Surgical valve replacement with a bioprosthetic stentless valve.	112000001	760 ACC NCDR







Section: Aortic Valve Replacement

Parent: Procedure History Details

Element: 14237 Aortic Valve Replacement Type Not Documented

Coding Instruction: Indicate if the surgical aortic valve replacement type was not documented.

Target Value: N/A

Technical Specification

Code: 725351001 Code System: SNOMED CT

Short Name: AVReplacementTypeND

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Aortic Valve Replacement Surgery

----- AND -----Element: 14268 Procedure History Occurrence

AND

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR







Section: Transcatheter AV Replacement

Parent: Procedure History Details

Element: 14249 Transcatheter Aortic Valve Replacement Implant ID

Coding Instruction: Indicate the model ID implanted in the transcatheter aortic valve replacement procedure.

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: TAVR Model ID

The model ID of the transcatheter valve used for transcatheter valve replacement procedure.

Source:

Technical Specification Code: 112000001766

Code System: ACC NCDR Short Name: TAVRImplantID Missing Data: Report

Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Aortic Valve Replacement - Transcatheter

AND

Element: 14268 Procedure History Occurrence
Operator: Equal
Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

Element: 14515 Transcatheter Aortic Valve Replacement Implant Diameter

Coding Instruction: Indicate the transcatheter aortic valve implant size.

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: TAVR Model ID

The model ID of the transcatheter valve used for transcatheter valve replacement procedure.

Source:

Technical Specification

Code: 112000001766
Code System: ACC NCDR
Short Name: TAVRImplantDia

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm
Valid Range: 5 - 100 mm

Data Source: User

Parent/Child Validation

Element: 14249 Transcatheter Aortic Valve

Replacement Implant ID

Operator:

Value: Any Value







Section: ICD Parent: Procedure History Details

Element: 14259 Cardiac Resynchronization Therapy Defibrillator

Coding Instruction: Indicate if the ICD includes a cardiac resynchronization therapy (CRT-D) device.

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: CRT-D

A cardiac resynchronization therapy device and defibrillator (CRT-D) has dual capabilities. It 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.

Source:

Technical Specification

Code: 112000002006
Code System: ACC NCDR
Short Name: CRTD
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Implantable Cardioverter Defibrillator

AND

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure







Section: Mitral Valve Annuloplasty

Parent: Procedure History Details

Element: 14257 Mitral Valve Annuloplasty Ring Type

Coding Instruction: Indicate the type of mitral annuloplasty ring implanted surgically.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 232744004
Code System: SNOMED CT
Short Name: PriorMVRingSurg
Missing Data: No Action

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14258 Mitral Valve Annuloplasty Ring

Type Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----Element: 12905 Procedure History Name

Omenator: Family

Operator: Equal

Value: Mitral Valve Annuloplasty Ring Surgery
----- AND ------

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND ------Element: 13171 TVT Pathway

Operator: Equal

Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Mitral Annuloplasty Ring Type - 1.3.6.1.4.1.19376.1.4.1.6.5.690

Selection	Definition	Source	Code	Code System
Circumferential Mitral Annuloplasty Ring	A circumferential mitral annuloplasty ring.		112000001772	ACC NCDR
Partial Mitral Annuloplasty	Ring A partial mitral annuloplasty ring.		112000001771	ACC NCDR







Section: Mitral Valve Annuloplasty

Parent: Procedure History Details

Element: 14258 Mitral Valve Annuloplasty Ring Type Not Documented

Coding Instruction: Indicate if the type of mitral annuloplasty ring implanted surgically was not documented.

Target Value: N/A

Technical Specification

Code: 232744004
Code System: SNOMED CT
Short Name: PriorMVRingSurgND

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 14455 Mitral Ring Implant ID

 $\label{local_continuity} \textbf{Coding Instruction:} \quad \text{Indicate the implant ID of the mitral ring or mitral band.}$

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 17107009
Code System: SNOMED CT
Short Name: MVRingImplantID
Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation
Element: 12905 Procedure History Name

Operator: Equal

Value: Mitral Valve Annuloplasty Ring Surgery

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND ------

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR







Section: Mitral Valve Annuloplasty

Parent: Procedure History Details

Element: 14533 Mitral Ring Implant Diameter

Coding Instruction: Indicate the mitral ring implant diameter size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 112000001807
Code System: ACC NCDR
Short Name: MVRingImplantDia

Missing Data: Report
Harvested: Yes (TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm

Valid Range: 5 - 100 mm Data Source: User

Parent/Child Validation

Element: 14455 Mitral Ring Implant ID

Operator:

Value: Any Value







Section: Mitral Valve Replacement

Parent: Procedure History Details

Element: 14241 Mitral Valve Replacement Type

Coding Instruction: Indicate the type of surgical mitral valve replacement.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 53059001

Code System: SNOMED CT

Short Name: PrevMVReplaceType

Missing Data: Report

Is Identifier: No

Harvested: Yes (TAVR, TMVR)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Mitral Valve Replacement Surgery

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 14242 Mitral Valve Replacement Type

Not Documented

Operator: Equal

Value: No (or Not Answered)

Mitral Valve Replacement Type - 1.3.6.1.4.1.19376.1.4.1.6.5.734

Selection	Definition	Source	Code	Code System
Mechanical			705991002	SNOMED CT
Stented			112000001758	ACC NCDR
Stentless			112000001760	ACC NCDR







Section: Mitral Valve Replacement

Parent: Procedure History Details

Element: 14242 Mitral Valve Replacement Type Not Documented

Coding Instruction: Indicate if the surgical mitral valve replacement type was not documented.

Target Value: N/A

Technical Specification

Code: 53059001 Code System: SNOMED CT

Short Name: PrevMVReplaceTypeND

Missing Data: Report

Harvested: Yes (TAVR, TMVR)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: BL Precision: Selection Type: Single Unit of Measure:

Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Mitral Valve Replacement Surgery

----- AND -----Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TAVR

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 14334 Surgical Mitral Valve Replacement Implant ID

Coding Instruction: Indicate the implant ID of the prosthetic mitral valve.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 17107009 Code System: SNOMED CT Short Name: SMVRImplantID Missing Data: Report Harvested: Yes (TMVR) Is Identifier: No Is Base Element: Yes

Is Followup No Element: Data Type: CD Precision:

Selection Type: Single (Dynamic List)

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name Operator: Equal

Value: Mitral Valve Replacement Surgery ----- AND -----

Element: 14268 Procedure History Occurrence

Operator: Equal

Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR







Section: Mitral Valve Replacement

Parent: Procedure History Details

Element: 14518 Surgical Mitral Valve Replacement Implant Diameter

Coding Instruction: Indicate the mitral valve implant size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 17107009
Code System: SNOMED CT
Short Name: SMVRImplantDia

Missing Data: Report Harvested: Yes (TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 16 - 36 mm

Valid Range: 5 - 100 mm Data Source: User

Parent/Child Validation

Element: 14334 Surgical Mitral Valve Replacement

Implant ID

Operator:

Value: Any Value







Section: Mitral Valve Transcatheter

Parent: Procedure History Details

Element: 14261 Mitral Valve Transcatheter Intervention Type

Coding Instruction: Indicate the type of transcatheter mitral valve intervention.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 112000002002
Code System: ACC NCDR
Short Name: PriorTMVRType

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Valid Range: Data Source: User

Usual Range:

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway
Operator: Equal

Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Mitral Valve Transcatheter Type - 1.3.6.1.4.1.19376.1.4.1.6.5.691

Selection	Definition	Source	Code	Code System
Leaflet Clip Procedure			112000001778	ACC NCDR
Direct Annuloplasty Intervention			112000001775	ACC NCDR
Coronary Sinus Based Intervention			112000001774	ACC NCDR
Valve in Native Valve			112000001776	ACC NCDR
Procedure				
Valve in Valve Procedure			112000001286	ACC NCDR
Other Mitral Valve			112000001777	ACC NCDR
Transcatheter Intervention	า			







Section: Mitral Valve Transcatheter

Parent: Procedure History Details

Element: 14510 Transcatheter Mitral Valve Replacement Implant ID

Coding Instruction: Indicate the transcatheter mitral valve replacement implant ID.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 17107009
Code System: SNOMED CT
Short Name: TMVRImplantID
Missing Data: Report

Harvested: Yes (TMVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: CD

Precision:
Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal

Value: TMVR

Element: 14534 Transcatheter Mitral Valve Replacement Implant Diameter

Coding Instruction: Indicate the transcatheter mitral valve replacement implant size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 112000001807
Code System: ACC NCDR
Short Name: TMVRImplantDia

Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm
Valid Range: 5 - 100 mm

Data Source: User

Parent/Child Validation

Element: 14510 Transcatheter Mitral Valve

Replacement Implant ID

Operator:

Value: Any Value







Section: Permanent Pacemaker

Parent: Procedure History Details

Element: 14260 Cardiac Resynchronization Therapy

Coding Instruction: Indicate if the pacemaker type includes cardiac resynchronization therapy (CRT).

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: Cardiac Resynchronization Therapy Pacemaker Placement

A CRT procedure is the placement of 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.

Source:

Technical Specification

Code: 704708004
Code System: SNOMED CT
Short Name: CRT
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Permanent Pacemaker

Data Source: User

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----

Element: 13171 TVT Pathway

Operator: Equal Value: TMVr

Element: 13171 TVT Pathway

Operator: Equal Value: TMVR

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure







Section: Tricuspid Valve Repair Surgery

Parent: Procedure History Details

Element: 14299 Tricuspid Valve Annuloplasty Ring

Coding Instruction: Indicate if the patient had a prior tricuspid annuloplasty ring implanted surgically.

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: Tricuspid Valve

A three-cusp valve of the heart that regulates the flow of blood between the right atrium and

the right ventricle of the heart

Source:

Technical Specification

Code: 46030003
Code System: SNOMED CT
Short Name: PreTVARing
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Tricuspid Valve Repair Surgery

Element: 14268 Procedure History Occurrence

AND

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure







Section: Tricuspid Valve Intervention

Parent: Procedure History Details

Element: 14300 Transcatheter Tricuspid Valve Intervention Type

Coding Instruction: Indicate the type of transcatheter tricuspid valve intervention.

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: Tricuspid Valve

A three-cusp valve of the heart that regulates the flow of blood between the right atrium and

the right ventricle of the heart

Source:

Technical Specification

Code: 112000001779
Code System: ACC NCDR
Short Name: PreTTVIType
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Tricuspid Valve Transcatheter Intervention

AND

AND

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure

Tricuspid Valve Intervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.735

Selection	Definition	Source	Code	Code System
Annuloplasty Ring			232782007	SNOMED CT
Other			112000001873	ACC NCDR







Section: Tricuspid Valve Replacement Surgery

Parent: Procedure History Details

Element: 14298 Surgical Tricuspid Valve Replacement Implant ID

Coding Instruction: Indicate the implant ID of the prosthetic tricuspid valve.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 703201004
Code System: SNOMED CT
Short Name: STVRImplantID
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Tricuspid Valve Replacement Surgery

AND

Element: 14268 Procedure History Occurrence

----- AND -----

Operator: Equal Value: Yes

Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14516 Surgical Tricuspid Valve Replacement Implant Diameter

Coding Instruction: Indicate the tricuspid valve implant size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 703201004
Code System: SNOMED CT
Short Name: STVRImplantDia

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm
Valid Range: 5 - 100 mm

Data Source: User

Parent/Child Validation

Element: 14298 Surgical Tricuspid Valve

Replacement Implant ID

Operator:

Value: Any Value







Section: Transcatheter TV Replacement

Parent: Procedure History Details

Element: 14301 Transcatheter Tricuspid Valve Replacement Implant ID

Coding Instruction: Indicate the implant ID of the prosthetic tricuspid valve.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 112000001810
Code System: ACC NCDR
Short Name: TTVRImplantID
Missing Data: Report

Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Precision:

Parent/Child Validation

Element: 12905 Procedure History Name

Operator: Equal

Value: Tricuspid Valve Replacement -

Transcatheter

Element: 14268 Procedure History Occurrence

Operator: Equal Value: Yes

----- AND -----Element: 13171 TVT Pathway

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14517 Transcatheter Tricuspid Valve Replacement Implant Diameter

Coding Instruction: Indicate the tricuspid valve implant size.

Target Value: The last value between birth and the first procedure in this admission

Technical Specification

Code: 112000001810
Code System: ACC NCDR
Short Name: TTVRImplantDia
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 16 - 36 mm
Valid Range: 5 - 100 mm
Data Source: User

Parent/Child Validation

Element: 14301 Transcatheter Tricuspid Valve

Replacement Implant ID

Operator:

Value: Any Value







Element: 14273 Transcatheter Valve Therapy Procedure Type Code: 112000001167 Coding Instruction: Indicate the TVT procedure performed. Code System: ACC NCDR Short Name: TVTProType Target Value: The value on current procedure Missing Data: Illegal Vendor Instruction: Transcatheter Valve Therapy Procedure Type (14273) cannot be (Transcatheter Mitral Valve

When Procedure History Name (12905) is (Mitral Valve Replacement Surgery) with Procedure

History Occurrence as (Yes) AND

Mitral Valve Transcatheter Intervention Type (14261) is (Valve in Native Valve Procedure OR

Valve in Valve Procedure)

Within an episode, a lab visit for Transcatheter Mitral Valve Repair can not happen in any

subsequent lab visit(s) for Transcatheter Mitral Valve Replacement.

Technical Specification

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Multiple

Is Identifier: No

Unit of Measure: Default Value: Null **Usual Range:**

Valid Range: Data Source: User

Data Source: User

Transcatheter Valve Therapy Procedure - 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selection	Definition	Source	Code	Code System
TAVR	Transcatheter aortic valve replacement		41873006	SNOMED CT
TMVr	Transcatheter mitral repair procedure		112000001801	ACC NCDR
TMVR	Transcatheter mitral valve replacement		112000001458	ACC NCDR
Tricuspid Valve Procedure	Transcatheter tricuspid valve procedures include eithe a transcatheter tricuspid valve replacement or transcatheter tricuspid valve repair.	ır	112000001977	ACC NCDR

Element: 13329	Procedure Room Entry Date and Time	Technical Specification
On the standard for	Indicate the date and time the patient entered the procedure room.	Code: 112000001197
Coding instruction:		Code System: ACC NCDR
Target Value:	The value on current procedure	Short Name: TVTPRocedureEntryTime
Supporting Definition:	Procedure Room Entry	Missing Data: Report
Supporting Demitton.	•	Harvested: Yes (TAVR, TMVR, TMVrpr,
	Concept associated with data elements pertaining to a patient's entry into a procedure room.	TTVP)
	Source:	Is Identifier: No
Vendor Instruction	: Procedure Room Entry Date and Time (13329) must be Less than or Equal to Procedure Start Date and Time (7000)	Is Base Element: Yes
vendor mstruction.		Is Followup No
		Element: NO
		Data Type: TS
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:

Element: 7000	Procedure Start Date and Time	Technical	Specification
Coding Instruction:	Indicate the date and time the procedure started. The time of the procedure is the time that the skin incision, vascular access, or its equivalent, was made in order to start the procedure.	Code System: AC	000142460 CC NCDR ocedureStartDateTime
	Note(s): Indicate the date/time (mm/dd/yyyy hours:minutes) using the military 24-hour clock, beginning at midnight (0000 hours).		egal es (BDS, TAVR, TMVR, //Vrpr, TTVP)
Target Value:	Any occurrence on current procedure	Is Identifier: No Is Base Element: Ye	es
Vendor Instruction:	Procedure Start Date and Time (7000) must be Less than or Equal to Discharge Date (10100)	Is Followup Element:	
		Data Type: TS Precision:	8
		Selection Type: Sill Unit of Measure:	ngle
		Default Value: Nu Usual Range:	ıll
		Valid Range: Data Source: Us	ser





Usual Range: Valid Range: Data Source: User

Section: Lab Visit Parent: Root

Element: 7005 **Technical Specification** Procedure End Date and Time Code: 1000142459 Coding Instruction: Indicate the ending date and time at which the operator completes the procedure and breaks Code System: ACC NCDR scrub at the end of the procedure. Short Name: ProcedureEndDateTime Missing Data: Illegal Note(s): Harvested: Yes (BDS, TAVR, TMVR, If more than one operator is involved in the case then use the date and time the last operator TMVrpr, TTVP) breaks scrub for the last time. Is Identifier: No Target Value: The value on current procedure Is Base Element: Yes Is Followup No Vendor Instruction: Procedure End Date and Time (7005) must be Greater than or Equal to Procedure Start Date and Time (7000) Element: Data Type: TS Procedure End Date and Time (7005) must be Less than or Equal to Discharge Date (10100) Precision: Selection Type: Single Procedure End Date and Time (7005) and Procedure Start Date and Time (7000) must not Unit of Measure: overlap on multiple procedures Default Value: Null

Technical Specification Element: 13330 Procedure Room Exit Date and Time Code: 112000001198 Coding Instruction: Indicate the date and time the patient exits the procedure room. Code System: ACC NCDR Short Name: TVTProcedureStopTime Target Value: The value on current procedure Missing Data: Report Supporting Definition: Procedure Room Exit Harvested: Yes (TAVR, TMVR, TMVrpr, Concept associated with data elements pertaining to a patient's exit from a procedure room. Is Identifier: No Source: Is Base Element: Yes Is Followup No Element: Data Type: TS Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Technical Specification Element: 13793 Mitral Leaflet Clip Procedure Code: 112000000208 Coding Instruction: Indicate if a mitral leaflet clip procedure was performed. Code System: ACC NCDR Target Value: The value on current procedure Short Name: ProcLeafClip Missing Data: Illegal Harvested: Yes (BDS, TMVrpr) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TMVr



Element: 12177



Full Specifications Data Dictionary v3.0



Section: Presentation and Evaluation Parent: Lab Visit

CAD Presentation

Coding Instruction: Indicate the patient's coronary artery disease (CAD) presentation. Choose the worst status.

Target Value: The highest value between 7 days prior to arrival and current procedure

Technical Specification Code: 112000000109 Code System: ACC NCDR Short Name: CADPresentation

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Is Identifier: No

Coronary Artery Disease Symptoms/Presentation - 1.3.6.1.4.1.19376.1.4.1.6.5.736

Selection	Definition	Source Code	Code System
No Symptoms, No Angina	The patient presents with no symptoms.	LA6111-4	LOINC
Unstable Angina	Unstable angina which includes angina at rest, ner onset or increasing angina (change in previously diagnosed pattern) within the past 2 months.	v 4557003	SNOMED CT
Stable Angina	Angina without a change in frequency or pattern for the six weeks prior to this cath lab presentation. Angina is controlled by rest and/or oral or transcutaneous medications.	or 233819005	SNOMED CT
Symptoms Unlikely to be Ischemic	Pain or symptoms that are not consistent with pain discomfort of myocardial ischemic origin within the two weeks.		ACC NCDR
STEMI	The patient presents with a STEMI within the past seven days.	401303003	SNOMED CT
Non-STEMI	The patient presents to the cath lab with an NSTEM within the past seven days.	l 401314000	SNOMED CT

Element: 14266	Heart Failure	Technical Specification
Coding Instruction:	: Indicate if there is physician documentation or report that the patient has been in a state of heart failure within the past 2 weeks.	Code: 84114007
		Code System: SNOMED CT
		Short Name: Prior2WksHF

Target Value: Any occurrence between 2 weeks prior to current procedure and current procedure

Supporting Definition: Heart Failure

Heart failure is a complex clinical syndrome that results from any structural or functional impairment of ventricular filling or ejection of blood. The cardinal manifestations of HF are dyspnea and fatigue, which may limit exercise tolerance, and fluid retention, which may lead to pulmonary and/or splanchnic congestion and/or peripheral edema. Some patients have exercise intolerance but little evidence of fluid retention, whereas others complain primarily of edema, dyspnea, or fatigue. Because some patients present without signs or symptoms of volume overload, the term "heart failure" is preferred over "congestive heart failure." There is no single diagnostic test for HF because it is largely a clinical diagnosis based on a careful history and physical examination.

Source: 2013 ACCF/AHA Guideline for the Management of Heart Failure; J Am Coll Cardiol. 2013;62(16):e147-e239. doi:10.1016/j.jacc.2013.05.019

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Is Base Element: Yes







Section: Presentation and Evaluation

Parent: Lab Visit

Element: 12163 New York Heart Association Classification

Coding Instruction: Indicate the patient's most severe dyspnea or functional class, coded as the New York Heart

Association (NYHA) classification.

Target Value: The highest value between 2 weeks prior to current procedure and current procedure

Supporting Definition: NYHA

The NYHA classes focus on exercise capacity and the symptomatic status of the disease.

Source: 2013 ACCF/AHA Guideline for the Management of Heart Failure; J Am Coll Cardiol.

2013;62(16):e147-e239. doi:10.1016/j.jacc.2013.05.019

Technical Specification

Code: 420816009
Code System: SNOMED CT
Short Name: Prior2weekNYHA

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

NYHA Functional Classification - 1.3.6.1.4.1.19376.1.4.1.6.5.8

Selection	Definition	Source	Code	Code System
Class I	Patients with cardiac disease but without resulting limitations of physical activity. Ordinary physical activit does not cause undue fatigue, palpitation, or dyspnea.	The Criteria Committee of the New York Heart y Association. Nomenclature and Criteria for Diagnosis of Diseases of the Heart and Great Vessels. 9th ed. Boston, Mass: Little, Brown & Co; 1994:253-256.	420300004	SNOMED CT
Class II	Patients with cardiac disease resulting in slight limitatio of physical activity. They are comfortable at rest. Ordinary physical activity results in fatigue, palpitation, or dyspnea.		421704003	SNOMED CT
Class III	Patients with cardiac disease resulting in marked limitation of physical activity. They are comfortable at rest. Less than ordinary activity causes fatigue, palpitation, or dyspnea.		420913000	SNOMED CT
Class IV	Patients with cardiac disease resulting in inability to carry on any physical activity without discomfort. Symptoms are present even at rest or minimal exertion If any physical activity is undertaken, discomfort is increased.		422293003	SNOMED CT

Element: 13175	Cardiogenic Shock
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Coding Instruction: Indicate if the patient has been in a state of cardiogenic shock within 24 hrs of procedure.

Target Value: Any occurrence between 24 hours prior to current procedure and up to current procedure

Supporting Definition: Cardiogenic Shock

Cardiogenic shock is defined as a sustained (>30 min) episode of systolic blood pressure <90 mm Hg and/or cardiac index <2.2 L/min per square meter determined to be secondary to cardiac dysfunction and/or the requirement for parenteral inotropic or vasopressor agents or mechanical support (eg, IABP, extracorporeal circulation, VADs) to maintain blood pressure and cardiac index above those specified levels.

Note: Transient episodes of hypotension reversed with IV fluid or atropine do not constitute cardiogenic shock. The hemodynamic compromise (with or without extraordinary supportive therapy) must persist for at least 30 min.

Source: Cannon CP, et al. 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: A Report of the ACC/AHA Task Force on Clinical Data Standards (Writing Committee to Develop Acute Coronary Syndromes and Coronary Artery Disease Clinical Data Standards). J Am Coll Cardiol. 2013;61(9):992-1025.

Technical Specification

Code: 89138009
Code System: SNOMED CT
Short Name: PriorCardioShock
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User



Element: 14267



Full Specifications **Data Dictionary v3.0**



Section: Presentation and Evaluation

Cardiac Arrest

Parent: Lab Visit

Coding Instruction: Indicate if the patient has had an episode of cardiac arrest within 24 hours of the procedure.

Target Value: Any occurrence between 24 hours prior to current procedure and up to current procedure

Supporting Definition: Cardiac Arrest

Cardiac arrest is defined as acute cardiac event documented by one of the following: ventricular fibrillation, rapid ventricular tachycardia or bradycardia rhythms with hemodynamic compromise causing loss of consciousness, pulseless rhythms (PEA), or asystole requiring cardiopulmonary resuscitation (two or more chest compressions or open chest massage, emergency temporary pacing, pericardiocentesis, institution of ECMO, or defibrillation) and without these measures death would have almost certainly resulted.

Source: Data Governance Subcommittee of the NCDR's SQOC

Technical Specification

Code: 410429000
Code System: SNOMED CT
Short Name: PriorCardArrest

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Element: 13186 Symptoms of Aortic Stenosis Present

Coding Instruction: Code yes if the patient has any symptoms of heart failure on arrival or anytime within the past

three months

For example, if a patient had symptoms within the past three months (even if there are no symptoms on arrival to the hospital), code yes. If there is documentation of symptoms (e.g. shortness of breath) but no documentation of heart failure, code yes. These indicate presence

of symptomatic aortic stenosis.

Target Value: Any occurrence between 3 months prior to arrival at this facility and start of the procedure

Technical Specification

Code: 60573004
Code System: SNOMED CT
Short Name: SxAS
Missing Data: Report

Harvested: Yes (BDS, TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TAVR ----- AND -----

Element: 13188 Symptoms of Aortic Stenosis Not

Documented

Operator: Equal

Value: No (or Not Answered)

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR







Section: Presentation and Evaluation

Parent: Lab Visit

Element: 13188 Symptoms of Aortic Stenosis Not Documented

Coding Instruction: Indicate whether there is no documentation of symptoms of aortic stenosis.

Target Value: N/A

Technical Specification

Code: 60573004
Code System: SNOMED CT
Short Name: SxASND
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13191 Five Meter Walk Test Performed

Coding Instruction: Indicate whether the five meter walk test was performed.

Note: If the five meter walk test was performed, 3 walk tests should be documented. If the patient is unable to walk for all three tests, document the tests that were completed.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Supporting Definition: Five Meter Walk Test

An outcome measure in STS the Adult Cardiac Surgery Database to predict frailty in individuals

who are candidates for cardiac surgery.

Source:

Technical Specification

Code: 112000001179
Code System: ACC NCDR
Short Name: FiveMWalkTest
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Is Identifier: No

Is Base Element: Yes

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Five Meter Walk Test Performed - 1.3.6.1.4.1.19376.1.4.1.6.5.456

Selection	Definition	Source	Code	Code System
Test Not Performed			112000001181	ACC NCDR
Test Performed			112000001180	ACC NCDR
Unable to Walk	The patient is physically unable to walk to perform this test. For example, the patient is wheelchair bound,has shortness of breath or other symptoms that are so severe, they are unable to walk.		112000001182	ACC NCDR







Section: Presentation and Evaluation

Parent: Lab Visit

Element: 13710 Six Minute Walk Test

Coding Instruction: Indicate whether the six minute walk test was performed.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 252478000
Code System: SNOMED CT
Short Name: SixMinWalkPerf

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: STS Risk Score Parent: Presentation and Evaluation

Element: 13698 Society of Thoracic Surgeons Risk Score Type

Coding Instruction: Indicate the patient's predicted risk of mortality for surgical valve replacement or repair as

determined by the heart team and based on the Society for Thoracic Surgeon's risk model.

The following STS risk scores should be documented based on the STS Adult Cardiac Surgery Risk Calculator:

TAVR: Isolated aortic valve replacement TMVR: Isolated mitral valve replacement

Mitral Leaflet Clip Procedure: mitral valve repair and isolated mitral valve replacement

Note: Currently there is not a risk score available for tricuspid procedures.

Target Value: The last value prior to the start of the first procedure

Vendor Instruction: When Society of Thoracic Surgeons Risk Score Type (13698) is Equal to (Society of Thoracic

Surgeons Risk Score for Aortic Valve Replacement) then Transcatheter Valve Therapy

Procedure Type (14273) must be Equal to (TAVR)

When Society of Thoracic Surgeons Risk Score Type (13698) is Equal to (Society of Thoracic Surgeons Risk Score for Mitral Valve Repair) then Transcatheter Valve Therapy Procedure

Type (14273) must be Equal to (TMVr)

When Society of Thoracic Surgeons Risk Score Type (13698) is Equal to (Society of Thoracic

Surgeons Risk Score for Mitral Valve Replacement) then Transcatheter Valve Therapy Procedure Type (14273) must be Equal to (TMVR,TMVr)

A Society of Thoracic Surgeons Risk Score Type (13698) may only be entered/selected once

Technical Specification

Code: 112000001412
Code System: ACC NCDR
Short Name: STSRiskScoreType

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Single Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Society of Thoracic Surgeons Risk Score Type - 1.3.6.1.4.1.19376.1.4.1.6.5.693

Selection	Definition	Source	Code	Code System
Society of Thoracic Surgeons Risk Score for Aortic Valve Replacement			112000001796	ACC NCDR
Society of Thoracic Surgeons Risk Score for Mitral Valve Repair			112000001795	ACC NCDR
Society of Thoracic Surgeons Risk Score for Mitral Valve Replacement			112000001793	ACC NCDR







Section: STS Risk Score

Parent: Presentation and Evaluation

Element: 14271 Society of Thoracic Surgeons Risk Score Measurement

Coding Instruction: Indicate the patient's predicted risk of mortality for surgical valve replacement or repair as

determined by the heart team and based on the Society for Thoracic Surgeon's risk calculator

(https://www.sts.org/resources/risk-calculator)

Target Value: The last value prior to the start of the first procedure

Technical Specification

Code: 112000001797
Code System: ACC NCDR
Short Name: STSRiskScoreValue

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

 Is Base Element:
 Yes

 Is Followup Element:
 No

 Data Type:
 PQ

 Precision:
 6,3

 Selection Type:
 Single

 Unit of Measure:
 %

 Default Value:
 Null

Usual Range: 2.000 - 15.000 % **Valid Range:** 0.000 - 100.000 %

Data Source: User

Parent/Child Validation

Element: 13698 Society of Thoracic Surgeons

Risk Score Type

Operator:

Value: Any Value







Section: Shared Decision Making

Parent: Presentation and Evaluation

Element: 14732 Shared Decision Making

Coding Instruction: Indicate if shared decision making was performed for the procedure.

Target Value: The value on current procedure

Supporting Definition: Shared Decision Making

Shared decision making occurs when a health care provider and a patient work together to

make a health care decision that is best for the patient.

The optimal decision takes into account evidence-based information about available options, the provider's knowledge and experience, and the patient's values and preferences.

Source: AHRQ.gov

Technical Specification

Code: 112000002041 Code System: ACC NCDR Short Name: SDM_Proc

Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes

Is Followup No Element: Data Type: BL Precision:

Selection Type: Single

Unit of Measure: **Default Value: Usual Range:** Valid Range: Data Source: User

Element: 14733 Shared Decision Making Tool Used

Coding Instruction: Indicate if a shared decision making tool was used.

Target Value: The value on current procedure

Technical Specification

Code: 415806002 Code System: SNOMED CT Short Name: SDM_Tool Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: **Default Value: Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14732 Shared Decision Making

Operator: Equal Value: Yes







Section: Shared Decision Making

Parent: Presentation and Evaluation

Element: 14734 Shared Decision Making Tool Name

Coding Instruction: Indicate what tool was used.

If the tool used is not in the drop-down list, please contact NCDR@acc.org to have a selection

added.

Target Value: The value on current procedure

Technical Specification

Code: 405083000
Code System: SNOMED CT
Short Name: SDM_Tool_Name

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value:
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14733 Shared Decision Making Tool

Used
Operator: Equal
Value: Yes

Shared Decision Making Tools - 1.3.6.1.4.1.19376.1.4.1.6.5.765

Selection	Definition	Source	Code	Code System
Other Shared Decision Making		100000351	ACC NCDR	
Tool				



Element: 13843



Full Specifications Data Dictionary v3.0



Section: KCCQ12	Parent: Presentation and Evaluation

Kansas City Cardiomyopathy Questionnaire 12 Performed Coding Instruction: Indicate if the baseline Kansas City Cardiomyopathy Questionnaire (KCCQ-12) was performed.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001540 Code System: ACC NCDR Short Name: KCCQ12_Performed

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User

Element: 13846 Kansas City Cardiomyopathy Questionnaire 12 Question 1a

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 1a.

Heart Failure Limitation - Showering/bathing

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

Technical Specification

Code: 112000001541 Code System: ACC NCDR Short Name: KCCQ12_1a Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyonathy Questionnaire 1a thru 1c - 1 3 6 1 4 1 19376 1 4 1 6 5 570

Italiaaa Oity Garaio	tansas oney our domy opacity & desironnaire 1a tina 10 - 1.5.6.1.4.1.1.5576.1.4.1.0.5.576				
Selection	Definition	Source	Code	Code System	
1 - Extremely Limited	·		100001173	ACC NCDR	
2 - Quite a Bit Limited			100001171	ACC NCDR	
3 - Moderately Limited	d		100001170	ACC NCDR	
4 - Slightly Limited			100014042	ACC NCDR	
5 - Not at All Limited			100001167	ACC NCDR	
6 - Limited for Other I			100014041	ACC NCDR	
or Did Not Do These	Activities				







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13848 Kansas City Cardiomyopathy Questionnaire 12 Question 1b

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 1b

Heart Failure Limitation - Walking 1 block on level ground

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001542 Code System: ACC NCDR

Short Name: KCCQ12_1b Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 1a thru 1c - 1.3.6.1.4.1.19376.1.4.1.6.5.570

Selection	Definition	Source	Code	Code System
1 - Extremely Limited			100001173	ACC NCDR
2 - Quite a Bit Limited			100001171	ACC NCDR
3 - Moderately Limited	i		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Not at All Limited			100001167	ACC NCDR
6 - Limited for Other F			100014041	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13849 Kansas City Cardiomyopathy Questionnaire 12 Question 1c

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 10

Heart Failure Limitation - Hurrying or jogging

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001543
Code System: ACC NCDR
Short Name: KCCQ12_1c

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 1a thru 1c - 1.3.6.1.4.1.19376.1.4.1.6.5.570

Selection	Definition	Source	Code	Code System
1 - Extremely Limited			100001173	ACC NCDR
2 - Quite a Bit Limited			100001171	ACC NCDR
3 - Moderately Limited	d		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Not at All Limited			100001167	ACC NCDR
6 - Limited for Other or Did Not Do These			100014041	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13851 Kansas City Cardiomyopathy Questionnaire 12 Question 2

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 2

Symptom Frequency - swelling in legs

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001544
Code System: ACC NCDR
Short Name: KCCQ12_2

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 2 - 1.3.6.1.4.1.19376.1.4.1.6.5.571

Selection	Definition	Source	Code	Code System
1 - Every Morning			112000001553	ACC NCDR
2 - Three or More Times P Week But Not Everyday	er		112000001554	ACC NCDR
3 - One to Two Times Per Week			112000001555	ACC NCDR
4 - Less Than Once a We	ek		112000001556	ACC NCDR
5 - Never Over the Past T Weeks	wo		112000001557	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13853 Kansas City Cardiomyopathy Questionnaire 12 Question 3

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 3

Symptom Frequency - fatigue

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001545
Code System: ACC NCDR
Short Name: KCCQ12_3

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 3 and 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.572

Selection	Definition	Source	Code	Code System
1 - All the Time			112000001818	ACC NCDR
2 - Several Times Per Day			112000001559	ACC NCDR
3 - At Least Once Per Day			112000001560	ACC NCDR
4 - Three or More Times Per Week But Not Everyday			112000001554	ACC NCDR
5 - One to Two Times Per Week			112000001555	ACC NCDR
6 - Less Than Once a Week			112000001556	ACC NCDR
7 - Never Over the Past Two Weeks)		112000001557	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13855 Kansas City Cardiomyopathy Questionnaire 12 Question 4

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 4

Symptom Frequency - shortness of breath

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001546 Code System: ACC NCDR

Short Name: KCCQ12_4 Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 3 and 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.572

Selection	Definition	Source	Code	Code System
1 - All the Time			112000001818	ACC NCDR
2 - Several Times Per Day			112000001559	ACC NCDR
3 - At Least Once Per Day			112000001560	ACC NCDR
4 - Three or More Times Per Week But Not Everyday			112000001554	ACC NCDR
5 - One to Two Times Per Week			112000001555	ACC NCDR
6 - Less Than Once a Week			112000001556	ACC NCDR
7 - Never Over the Past Two Weeks)		112000001557	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13857 Kansas City Cardiomyopathy Questionnaire 12 Question 5

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 5

Symptom Frequency - sleep sitting up due to shortness of breath

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001547 Code System: ACC NCDR

Short Name: KCCQ12_5 Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 5 - 1.3.6.1.4.1.19376.1.4.1.6.5.704

Selection	Definition	Source	Code	Code System
1 - Every Night			112000001819	ACC NCDR
2 - Three or More Times Per Week But Not Everyday			112000001554	ACC NCDR
3 - One to Two Times Per Week			112000001555	ACC NCDR
4 - Less Than Once a Week			112000001556	ACC NCDR
5 - Never Over the Past Two Weeks			112000001557	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13859 Kansas City Cardiomyopathy Questionnaire 12 Question 6

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 6

Quality of Life - effect on enjoyment of life due to heart failure

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001548 Code System: ACC NCDR

Short Name: KCCQ12_6
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 6 - 1.3.6.1.4.1.19376.1.4.1.6.5.573

Selection	Definition	Source	Code	Code System
1 - It Has Extremely Limi Enjoyment of Life	ited My		100014049	ACC NCDR
2 - It Has Limited My Enj of Life Quite a Bit	oyment		100014050	ACC NCDR
3 - It Has Moderately Lin My Enjoyment of Life	nited		100014051	ACC NCDR
4 - It Has Slightly Limited Enjoyment of Life	i Му		100014052	ACC NCDR
5 - It Has Not Limited My Enjoyment of Life at All			100014053	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13861 Kansas City Cardiomyopathy Questionnaire 12 Question 7

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 7

Quality of life - remaining life with heart failure

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001549 Code System: ACC NCDR Short Name: KCCQ12_7

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 7 - 1.3.6.1.4.1.19376.1.4.1.6.5.574

Selection	Definition	Source	Code	Code System
1 - Not At All Satisfied			112000001561	ACC NCDR
2 - Mostly Dissatisfied			112000001562	ACC NCDR
3 - Somewhat Satisfie			112000001563	ACC NCDR
4 - Mostly Satisfied			112000001564	ACC NCDR
5 - Completely Satisfie	d		112000001565	ACC NCDR

Element: 13863 Kansas City Cardiomyopathy Questionnaire 12 Question 8a

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 8a.

Social limitation - hobbies, recreational activities

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001550
Code System: ACC NCDR
Short Name: KCCQ12_8a
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

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Selection	Definition	Source	Code	Code System	
1 - Severely Limited		,	112000001566	ACC NCDR	
2 - Limited Quite a Bit			112000001567	ACC NCDR	
3 - Moderately Limited	i		100001170	ACC NCDR	
4 - Slightly Limited			100014042	ACC NCDR	
5 - Did Not Limit at All			112000001569	ACC NCDR	
6 - Does Not Apply or Do for Other Reasons			112000001570	ACC NCDR	







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13865 Kansas City Cardiomyopathy Questionnaire 12 Question 8b

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 8

Social limitation - working or doing household chores

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001551
Code System: ACC NCDR
Short Name: KCC012, 8b

Short Name: KCCQ12_8b Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

Selection	Definition	Source	Code	Code System
1 - Severely Limited			112000001566	ACC NCDR
2 - Limited Quite a Bit			112000001567	ACC NCDR
3 - Moderately Limited			100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Did Not Limit at All			112000001569	ACC NCDR
6 - Does Not Apply or			112000001570	ACC NCDR







Section: KCCQ12 Parent: Presentation and Evaluation

Element: 13867 Kansas City Cardiomyopathy Questionnaire 12 Question 8c

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 80

Social limitation - visiting family or friends

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 112000001552 Code System: ACC NCDR Short Name: KCCQ12_8c

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

Selection	Definition	Source	Code	Code System
1 - Severely Limited			112000001566	ACC NCDR
2 - Limited Quite a Bit			112000001567	ACC NCDR
3 - Moderately Limited			100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Did Not Limit at All			112000001569	ACC NCDR
6 - Does Not Apply or Did Not Do for Other Reasons	t		112000001570	ACC NCDR

Element: 14310 KCCQ Overall Summary Score

Coding Instruction: (Auto Calculated) This field is auto-populated by your application.

Kansas City Cardiomyopathy Questionnaire (KCCQ-12) Overall Summary Score.

Note(s): The 12 patient responses are reduced into four summary scores (Physical Limitation Score, Symptom Frequency Score, Quality of Life Score, Social Limitation Score). The four

summary scores are used to calculate the Overall Summary Score.
For more information, please refer to the KCCQ-12 Scoring Instructions document

provided by the STS/ACC TVT Registry.

Target Value: The value on start of current procedure

Technical Specification

Code: 112000001540
Code System: ACC NCDR
Short Name: KCCQ12_Overall

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: NUM
Precision: 5,2
Selection Type: Single
Unit of Measure:
Default Value: Null

Valid Range:
Data Source: Computed

Usual Range:

Parent/Child Validation

Element: 13843 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes







Section: Five Meter Walk Test

Parent: Presentation and Evaluation

Element: 13199 Five Meter Walk Test Counter

Coding Instruction: The software assigned five meter walk test counter should start at one and be incremented by

one for each test performed, in chronological order, during the clinical encounter. The five meter walk test number should be assigned sequentially in ascending order. Do not skip

numbers.

Note: If the five meter walk test was performed, 3 walk tests should be documented. If the $\,$

patient is unable to walk for all three tests, document the tests that were completed.

Target Value: N/A

Supporting Definition: Five Meter Walk Test

An outcome measure in STS the Adult Cardiac Surgery Database to predict frailty in individuals

who are candidates for cardiac surgery.

Source:

Technical Specification

Code: 112000002003
Code System: ACC NCDR
Short Name: FiveMWTCounter

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Element:

Data Type: CTR

Precision:

Selection Type: Single Unit of Measure: Default Value: Null Usual Range:

Valid Range:
Data Source: Automatic

Parent/Child Validation

Element: 13191 Five Meter Walk Test Performed

Operator: Equal

Value: Test Performed

Element: 13201 Five Meter Walk Test Time

Coding Instruction: Indicate the value of the five meter walk test in seconds.

Target Value: The value on current admission

Supporting Definition: Five Meter Walk Test

An outcome measure in STS the Adult Cardiac Surgery Database to predict frailty in individuals

who are candidates for cardiac surgery.

Source:

Technical Specification

Code: 112000001184
Code System: ACC NCDR
Short Name: FiveMWTTime
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: sec
Default Value: Null
Usual Range: 1 - 100 sec

Valid Range: 1 - 500 sec Data Source: User

Parent/Child Validation

Element: 13191 Five Meter Walk Test Performed

Operator: Equal

Value: Test Performed







Section: Six Minute Walk Test

Parent: Presentation and Evaluation

Element: 13711 Six Minute Walk Test Date

Coding Instruction: Indicate the date the six minute walk test was performed.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 252478000
Code System: SNOMED CT
Short Name: SixMinWalkDate

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13710 Six Minute Walk Test

Operator: Equal Value: Yes

Element: 13712 Six Minute Walk Test Total Distance

Coding Instruction: Indicate the total distance, in feet, the patient walked.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification
Code: 112000001422
Code System: ACC NCDR

Short Name: SixMinWalkDist
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Parent/Child Validation

Element: 13710 Six Minute Walk Test

Operator: Equal Value: Yes







Section: Six Minute Walk Test

Parent: Presentation and Evaluation

Element: 14262 Six Minute Walk Test Reason Not Performed

Coding Instruction: Indicate the reason why the six minute walk test was not performed.

Target Value: The last value between 90 days prior to the start of the current procedure and the start of

procedure

Technical Specification

Code: 252478000
Code System: SNOMED CT
Short Name: SixMinWalkReason

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13710 Six Minute Walk Test

Operator: Equal Value: No

Six Minute Walk Test Reason Not Performed - 1.3.6.1.4.1.19376.1.4.1.6.5.544

Selection	Definition	Source	Code	Code System
Non-Cardiac Reason			112000001418	ACC NCDR
Cardiac Reason			112000001419	ACC NCDR
Patient Not Willing to Wa	lk		112000001420	ACC NCDR
Not Performed by Site			112000001421	ACC NCDR







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Technical Specification Element: 6030 Hemoglobin Code: 718-7 Coding Instruction: Indicate the hemoglobin (Hgb) value in g/dL. Code System: LOINC Short Name: HGB Note(s): Missing Data: Report This may include POC (Point of Care) testing results or results obtained prior to arrival at this Harvested: Yes (BDS, TAVR, TMVR, facility. TMVrpr, TTVP) Target Value: The last value within 30 days prior to the first procedure in this admission Is Identifier: No Is Base Element: Yes Supporting Definition: Hemoglobin Is Followup No Hemoglobin (Hb or Hgb) is the iron-containing oxygen-transport metalloprotein in the red blood Element: cells. It carries oxygen from the lungs to the rest of the body (i.e. the tissues) where it Data Type: PQ releases the oxygen to burn nutrients and provide energy. Hemoglobin concentration Precision: 4,2 measurement is among the most commonly performed blood tests, usually as part of a complete blood count. If the concentration is below normal, this is called anemia. Anemias are Selection Type: Single classified by the size of red blood cells: "microcytic" if red cells are small, "macrocytic" if they Unit of Measure: g/dL are large, and "normocytic" if otherwise. Dehydration or hyperhydration can greatly influence Default Value: Null measured hemoglobin levels. Usual Range: 5.00 - 20.00 g/dL Source: http://s.details.loinc.org/LOINC/718-7.html?sections=Simple Valid Range: 1.00 - 50.00 g/dL Data Source: User

Parent/Child Validation
Element: 6031 Hemoglobin Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 6031 Hemoglobin Not Drawn

Coding Instruction: Indicate if the hemoglobin was not drawn.

Target Value: The last value within 30 days prior to the first procedure in this admission

Technical Specification

Code: 718-7
Code System: LOINC
Short Name: HGBND
Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User



Element: 6036



Full Specifications **Data Dictionary v3.0**



Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Technical Specification Element: 6035 Sodium Code: 2950-4 Coding Instruction: Indicate the sodium (Na) level, in mEq/L. Code System: LOINC Short Name: Sodium Target Value: The last value within 30 days prior to the first procedure in this admission Missing Data: Report Supporting Definition: Sodium Harvested: Yes (TAVR, TMVR, TMVrpr, Sodium is an essential nutrient that regulates blood volume, blood pressure, osmotic equilibrium TTVP) and electrolyte balance. Sodium chloride is the principal source of sodium in the diet, and is Is Identifier: No used for seasoning and as a preservative. Increased levels of sodium intake can cause Is Base Element: Yes hypertension and reportedly leads to 7.6 million premature deaths worldwide. Sodium is also Is Followup No important in neuron function and osmoregulation between cells and the extracellular fluid. Element: Source: http://s.details.loinc.org/LOINC/2950-4.html?sections=Simple Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mEg/L Default Value: Null Usual Range: 120 - 150 mEq/L Valid Range: 1 - 300 mEq/L Data Source: User Parent/Child Validation

Parent/Child Validation 6036 Sodium Not Drawn

Element: 6036 Operator: Equal

Value: No (or Not Answered)

Sodium Not Drawn Technical Specification

Coding Instruction: Indicate if the sodium level was not drawn.

Target Value: The last value within 30 days prior to the first procedure in this admission

Code: 2950-4
Code System: LOINC
Short Name: SodiumND
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Technical Specification Element: 6050 Creatinine Code: 2160-0 Coding Instruction: Indicate the creatinine (Cr) level mg/dL. Code System: LOINC Short Name: PreProcCreat Note(s): Missing Data: Report This may include POC (Point of Care) testing results or results obtained prior to arrival at this Harvested: Yes (BDS, TAVR, TMVR. TMVrpr, TTVP) Target Value: The last value between 30 days prior to the procedure and the current procedure Is Identifier: No Is Base Element: Yes **Supporting Definition: Creatinine** Is Followup No Creatinine or creatine anhydride, is a breakdown product of creatine phosphate in muscle. The Element: loss of water molecule from creatine results in the formation of creatinine. It is transferred to Data Type: PQ the kidneys by blood plasma, whereupon it is eliminated by glomerular filtration and partial Precision: 4,2 tubular excretion. Creatinine is usually produced at a fairly constant rate and measuring its serum level is a simple test. A rise in blood creatinine levels is observed only with marked Selection Type: Single damage to functioning nephrons; therefore this test is not suitable for detecting early kidney Unit of Measure: mg/dL disease. Creatine and creatinine are metabolized in the kidneys, muscle, liver and pancreas. Default Value: Null Source: http://s.details.loinc.org/LOINC/2160-0.html?sections=Simple Usual Range: 0.10 - 5.00 mg/dL Valid Range: 0.10 - 30.00 mg/dL Data Source: User

Parent/Child Validation

Element: 6051 Creatinine Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 6051 Creatinine Not Drawn

Coding Instruction: Indicate if a creatinine level was not drawn.

Target Value: N/A

Technical Specification

Code: 2160-0 Code System: LOINC

Short Name: PreProcCreatND

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Element: 6055 Bilirubin (Total)

Coding Instruction: Indicate the total bilirubin (mg/dL)

Note(s):

This may include POC (Point of Care) testing results.

Target Value: The last value between 30 days prior to the procedure and the current procedure

Supporting Definition: Bilirubin (Total)

Bilirubin is the brownish yellow breakdown product of normal red blood cell, specifically heme, catabolism. Bilirubin is excreted in bile, and its levels are elevated in certain diseases including bile obstruction, hepatitis, cirrhosis, liver or pancreatic tumor, hemolysis, certain medications and inherited disorders. Levels of bilirubin in amniotic fluid are indicative of the severity of fetal hemolysis as in Rh disease. It is responsible for the brownish yellow color of bruises and in

jaundice.

Source: http://s.details.loinc.org/LOINC/42719-5.html?sections=Simple

Technical Specification

Code: 42719-5
Code System: LOINC
Short Name: Bilirubin
Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: mg/dL
Default Value: Null

Usual Range: 0.05 - 1.50 mg/dL **Valid Range:** 0.01 - 30.00 mg/dL

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND -----

Element: 6056 Bilirubin Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 6056 Bilirubin Not Drawn

Coding Instruction: Indicate if the total Bilirubin was not drawn.

Target Value: N/A

Technical Specification

Code: 42719-5
Code System: LOINC
Short Name: BilirubinND
Missing Data: Report

Harvested: Yes (TAVR, TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Operator: Equal
Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal



Element: 14210



Full Specifications **Data Dictionary v3.0**



Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Coding Instruction: Indicate the total albumin (in g/dL).

Albumin

Target Value: The last value between 30 days prior to the procedure and the current procedure

Technical Specification

Code: 52454007
Code System: SNOMED CT
Short Name: Albumin
Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Precision: 3,1
Selection Type: Single
Unit of Measure: g/dL
Default Value: Null
Usual Range: 3.5 - 5.0 g/dL
Valid Range: 1.0 - 10.0 g/dL
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14211 Albumin Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 14211 Albumin Not Drawn

Coding Instruction: Indicate true if the total albumin was not drawn

Target Value: N/A

Technical Specification

Code: 52454007
Code System: SNOMED CT
Short Name: Albumin_ND
Missing Data: Report

Harvested: Yes (TAVR, TTVP)
Is Identifier: No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Is Base Element: Yes

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal



Element: 13213



Full Specifications **Data Dictionary v3.0**



Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Coding Instruction: Indicate the pre-procedure platelet count in platelets per microliter.

Target Value: The last value between 30 days prior to the procedure and the current procedure

Supporting Definition: Platelet Count

A laboratory test used to determine of the number of platelets in a blood sample.

Source: NCI Thesaurus.

Platelet Count

Technical Specification

Code: 777-3
Code System: LOINC
Short Name: PlateletCt
Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)

 $\begin{array}{lll} \textbf{Usual Range:} & 150,000 - 400,000 \; \mu L \\ \textbf{Valid Range:} & 1,000 - 900,000 \; \mu L \\ \end{array}$

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND -----

Element: 13214 Platelet Count Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 13214 Platelet Count Not Drawn

Coding Instruction: Indicate if a platelet count was not drawn prior to the procedure.

Target Value: N/A

Supporting Definition: Platelet Count

A laboratory test used to determine of the number of platelets in a blood sample.

Source: NCI Thesaurus.

Technical Specification

Code: 777-3
Code System: LOINC
Short Name: PlateletCtND
Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Element: 13203

Coding Instruction: Indicate the international normalized ratio (INR) if the patient is on routine warfarin or coumadin

Target Value: The last value between 30 days prior to the procedure and the current procedure

Supporting Definition: International Normalized Ratio (INR)

The INR is specifically intented for evaluating protime results on patients stabilized on long term oral anticoagulant therapy. The INR is not appropriate to evalulate hemostatic function in patients with liver disease, for screening for hereditary factor deficiencies or acquired vitamin K deficiencies, or for routine preoperative screening; this should be evaluated on the normal range in seconds. INR is calculated by the equation, INR = (PTR) raised to the power of ISI, where ISI = International Sensitivity Index (assigned to each reagent thromboplastin). PTR = prothrombin time ratio (pat PT/pop mean PT). Computation of the INR of specific thromboplastin reagent should allow for uniformity in prothrombin time testing regardless of the reagent

system or instrumentation used.

Source: http://s.details.loinc.org/LOINC/6301-6.html?sections=Simple

Technical Specification

Code: 34714-6 Code System: LOINC Short Name: INRtvt Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 3,1 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range: 0.9 - 1.3** Valid Range: 0.5 - 30.0

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----

Element: 6046 International Normalized Ratio Not

Drawn Operator: Equal

Value: No (or Not Answered)

International Normalized Ratio Not Drawn Element: 6046

Coding Instruction: Indicate if INR was not drawn.

Target Value: N/A

Technical Specification

Code: 34714-6 Code System: LOINC Short Name: INRND Missing Data: Report

Is Identifier: No

Harvested: Yes (TAVR, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal



Element: 14280



Full Specifications **Data Dictionary v3.0**



Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Coding Instruction: Indicate the B-type natriuretic peptide (BNP) value.

BNP

Target Value: The last value between birth and the first procedure in this admission

Supporting Definition: Natriuretic peptide B

Brain natriuretic peptide (BNP) is an active fragment (1-32) of ProBNP which is produced by myocardial cells. It increases in both right-sided and left-sided heart failure as well as in systolic and diastolic heart failure. Thus, it is used to diagnose and manage heart failure. When a patient is taking recombinant PBN (Natricor), BNP will reflect serum levels. NT-ProBNP, an inactive fragment (1-78) of ProBNP is used to assess the degree of failure. Both of these polypeptides have roughly the same predictive power. NT-ProBNP is commonly called ProBNP.

Source: http://s.details.loinc.org/LOINC/42637-9.html?sections=Simple

Technical Specification

Code: 42637-9 Code System: LOINC

Short Name: PreProc_BNPValue

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

> Usual Range: 5 - 1,000 pg/mL Valid Range: 1 - 10,000 pg/mL

Data Source: User

Parent/Child Validation

Element: 13205 B-Type Natriuretic Peptide Not

Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Element: 13205 B-Type Natriuretic Peptide Not Drawn

Coding Instruction: Indicate if a pre-procedure B-type natriuretic peptide (BNP) was not collected.

Target Value: N/A

Technical Specification

Code: 42637-9

Code System: LOINC

Short Name: PreProcBNPNotDrawn

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Element: 14279 N-Terminal Pro B-Type Natriuretic Peptide Value

Coding Instruction: Indicate the N-Terminal Pro B-Type Natriuretic Peptide (NT-proBNP) Value.

Target Value: The last value between 6 months prior to procedure and the start of the current procedure

Supporting Definition: N-Terminal Pro B-Type Natriuretic Peptide Value

ProBNP is the 108 amino acid pro-hormone of BNP (Brain Naturetic Peptide) that is produced mainly in the left ventricle. The prohormone splits into two polypeptides- the biologically active but shorter BNP (77-108) and the longer N terminal (1-76) fragment called NT-proBNP. Commercial assays are available for NT-proBNP because of its usefulness in predicting cardiovascular risk. In one study, it was the single best predictor of survival among patients with the acute coronary syndrome. It also declines with successful treatment of left ventricular dysfunctionand heart failure and is used by some to track the success of such treatment. No commercial assays exist for proBNP (the whole peptide)- though the trade name for one companies NT-proBNP is "proBNP" -- a misnomer. We include proBNP as the a related name for NT-proBNP so that people who call it proBNP will find it in LOINC.

Source: Regenstrief Help

Source: http://s.details.loinc.org/LOINC/33762-6.html?sections=Simple

Technical Specification

Code: 33762-6 Code System: LOINC

Short Name: PreProcedureNTBNP

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 5,0 Selection Type: Single Unit of Measure: pg/mL Default Value: Null

> Usual Range: 5 - 30,000 pg/mL Valid Range: 5 - 30,000 pg/mL

Data Source: User

Parent/Child Validation

Element: 13206 N-Terminal Pro B-Type Natriuretic

Peptide Not Drawn

Operator: Equal

Value: No (or Not Answered)

----- AND Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure Clinical Data

Parent: Presentation and Evaluation

Element: 13206 N-Terminal Pro B-Type Natriuretic Peptide Not Drawn

Coding Instruction: Indicate if a pre-procedure N-terminal pro B-type natriuretic peptide (NT-proBNP) was not

collected

Target Value: N/A

Technical Specification

Code: 33762-6

Code System: LOINC

Short Name: PreProcNTBNPNotDrawn

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure ECG and Pulmonary Function

Parent: Presentation and Evaluation

Technical Specification Element: 13216 Forced Expiratory Volume in One Second Predicted Code: 19925-7 Coding Instruction: Indicate the FEV1 % predicted from the most recent pulmonary function test prior to procedure. Code System: LOINC Short Name: FEV1 Target Value: The last value between 12 months prior to arrival and start of the first procedure Missing Data: Report Supporting Definition: FEV1 Harvested: Yes (TAVR, TMVR, TMVrpr, A test of lung function, the FEV1 is the volume exhaled during the first second of a forced TTVP) expiratory maneuver started from the level of total lung capacity. It is the most frequently used Is Identifier: No index for assessing bronchoconstriction or bronchodilatation. Is Base Element: Yes Is Followup No FEV1% predicted is defined as FEV1% of the patient divided by the average FEV1% in the Element: population for any person of similar age, sex and body composition. Data Type: PQ Source: NCI Thesaurus Precision: 3,0

Parent/Child Validation

Element: 13217 Forced Expiratory Volume in One

Second Predicted Not Performed

Operator: Equal

Value: No (or Not Answered)

Selection Type: Single
Unit of Measure: %
Default Value: Null
Usual Range: 25 - 100 %
Valid Range: 1 - 150 %
Data Source: User

Element: 13217 Forced Expiratory Volume in One Second Predicted Not Performed

Coding Instruction: Indicate whether % predicted Forced Expiratory Volume (FEV1) was not performed or the patient did not have a pulmonary function test prior to the procedure.

ations did not have a pullionary function test prior to the proof

Target Value: N/A Supporting Definition: FEV1

A test of lung function, the FEV1 is the volume exhaled during the first second of a forced expiratory maneuver started from the level of total lung capacity. It is the most frequently used index for assessing bronchoconstriction or bronchodilatation.

FEV1% predicted is defined as FEV1% of the patient divided by the average FEV1% in the

population for any person of similar age, sex and body composition.

Source: NCI Thesaurus

Technical Specification

Code: 19925-7 Code System: LOINC Short Name: FEV1ND Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null

Usual Range: Valid Range: Data Source: User







Section: Pre-Procedure ECG and Pulmonary Function

Parent: Presentation and Evaluation

Element: 13218 Diffusing Capacity of the Lungs for Carbon Monoxide Predicted

Coding Instruction: Indicate the % predicted diffusing capacity of the lungs for carbon monoxide (DLCO) value obtained for the patient. Choose the value that represents the lowest % predicted whether or

not it is the simple DLCO or the DLCO/VA.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Supporting Definition: DLCO

A measurement of carbon monoxide (CO) transfer from inspired gas to pulmonary capillary

blood.

Source: NCI Thesaurus

Technical Specification

Code: 112000001185 Code System: ACC NCDR Short Name: DLCOPred Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: % Default Value: Null **Usual Range: 10 - 150 %**

Valid Range: 1 - 200 % Data Source: User

Parent/Child Validation

Element: 13219 Diffusing Capacity of the Lungs

for Carbon Monoxide Not Performed

Operator: Equal

Value: No (or Not Answered)

Element: 13219 Diffusing Capacity of the Lungs for Carbon Monoxide Not Performed

Coding Instruction: Indicate if a lung diffusion test (DLCO) was not performed.

Target Value: N/A Supporting Definition: DLCO

A measurement of carbon monoxide (CO) transfer from inspired gas to pulmonary capillary

Source: NCI Thesaurus

Technical Specification

Code: 112000001185 Code System: ACC NCDR Short Name: DLCOND Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User







Section: Pre-Procedure ECG and Pulmonary Function

Parent: Presentation and Evaluation

Element: 5055 Non-Ventricular Paced QRS duration

Coding Instruction: Indicate the duration of the non-ventricular paced or intrinsic QRS complex, in milliseconds, that was derived from the surface electrocardiogram (ECG). Surface ECGs are obtained from the

surface of the body and do not include intracardiac ECGs.

Note(s)

If no ECG is available, a pre-procedure 6 inch cardiac rhythm strip or clinician documentation

may be utilized to obtain this information.

Target Value: The last value within 30 days prior to the first procedure in this admission

Technical Specification

Code: 251208001
Code System: SNOMED CT
Short Name: NVPQRS
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: msec
Default Value: Null
Usual Range: 20 - 250 msec

Valid Range: 10 - 300 msec

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Element: 5045 Only Ventricular Paced QRS

Complexes Present

Operator: Equal

Value: No (or Not Answered)







Section: Pre-Procedure ECG and Pulmonary Function

Parent: Presentation and Evaluation

Element: 5045 Only Ventricular Paced QRS Complexes Present

Coding Instruction: Indicate if there were only ventricular paced QRS complexes present.

Note(s):

If the patient has some intrinsic ventricular complexes present, code "No".

If no ECG is available, a pre-procedure 6 inch cardiac rhythm strip or clinician documentation

may be utilized to obtain this information.

Target Value: The last value within 30 days prior to the first procedure in this admission

Technical Specification

Code: 100001120
Code System: ACC NCDR
Short Name: VPQRS
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure Medication(s)

Parent: Presentation and Evaluation

Element: 13699 Anticoagulants Administered

Coding Instruction: Indicate whether anticoagulants were administered.

Target Value: Any occurrence between 24 hours prior to current procedure and up to current procedure

Technical Specification

Code: 112000001416
Code System: ACC NCDR
Short Name: PreProcAnticoag

Missing Data: Report

Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup

Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Valid Range: Data Source: User

Usual Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Pre-procedure Medication Administration - 1.3.6.1.4.1.19376.1.4.1.6.5.44

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Element: 13643 Positive Inotropes Administered

Coding Instruction: Indicate if positive inotropes was administered.

For patients requiring IV inotropic support, indicate positive inotropes only.

Target Value: Any occurrence between 24 hours prior to current procedure and up to current procedure

Technical Specification

Code: 112000001358
Code System: ACC NCDR
Short Name: PreOpInotropes
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Pre-procedure Medication Administration - 1.3.6.1.4.1.19376.1.4.1.6.5.44

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13220 Diagnostic Catheterization Performed Technical Specification

Coding Instruction: Indicate whether diagnostic cardiac catheterization was performed.

Target Value: The last value between 12 months prior to arrival and start of the first procedure Missing Data: Report

Rechnical Specification

Code: 41976001

Code System: SNOMED CT

Short Name: DxCathPer

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:
Data Source: User

Element: 13222 Diagnostic Catheterization Date

Coding Instruction: Indicate the date the diagnostic catheterization was performed.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 41976001
Code System: SNOMED CT
Short Name: DxCathDt
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13220 Diagnostic Catheterization

Performed

Operator: Equal Value: Yes







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13381 Number of Diseased Vessels

Coding Instruction: Indicate the number of diseased major native coronary vessel systems: LAD system,

circumflex system, and/or right system with >= 50% narrowing of any vessel preoperatively.

Notes:

1. Do not include coronary artery bypass grafts.

2. Left main disease (>=50%) is counted as TWO vessels (LAD and Circumflex, which may include a Ramus Intermedius). For example, left main and RCA would count as three total.

Target Value: The highest value between birth and start of the procedure

Technical Specification

Code: 112000000201
Code System: ACC NCDR
Short Name: NumDisV
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Is Identifier: No

Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13382 Number of Diseased Vessels Not

Documented Operator: Equal

Value: No (or Not Answered)

Number of Diseased Vessels - 1.3.6.1.4.1.19376.1.4.1.6.5.380

Selection	Definition	Source	Code	Code System
None			100001231	ACC NCDR
One			11200000788	ACC NCDR
Two			112000000790	ACC NCDR
Three			112000000792	ACC NCDR

Element: 13382 Number of Diseased Vessels Not Documented

Coding Instruction: Indicate true if the number of diseased vessels was not documented in the medical record.

Target Value: N/A

Technical Specification

Code: 112000000201
Code System: ACC NCDR
Short Name: NumDisVND
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Usual Range: Valid Range: Data Source: User

Default Value: Null







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13260 Left Main Stenosis Greater Than or Equal to 50 Percent

Coding Instruction: Indicate whether the patient has left main coronary disease. Left main coronary disease is

present when there is >= 50% compromise of vessel diameter pre-operatively.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Supporting Definition: Left Main Stenosis

Stenosis of the left main coronary artery.

Source:

Technical Specification

Code: 112000001186
Code System: ACC NCDR
Short Name: LMainDis
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:
Data Source: User

Parent/Child Validation

Element: 13261 Left Main Stenosis Not

Documented Operator: Equal

Value: No (or Not Answered)

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Element: 13261	Left Main Stenosis Not Documented	Technical Specification
On the sales at second as	Indicate whether the OV standing of the 1-th and a second second at the second	Code: 112000001186
Coding Instruction:	Indicate whether the % stenosis of the left main coronary artery was not documented	

Target Value: N/A

Supporting Definition: Left Main Stenosis

Stenosis of the left main coronary artery.

Source:

Code: 112000001180
Code System: ACC NCDR
Short Name: LMainDisND
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Data Source: User

Usual Range: Valid Range:







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Proximal Left Anterior Descending Artery Disease Greater or Equal to 70 **Element: 13301**

percent

Coding Instruction: Indicate whether the percent luminal narrowing of the proximal left anterior descending artery

at the point of maximal stenosis is greater than or equal to 70%. Target Value: The last value between 12 months prior to arrival and start of the first procedure

Supporting Definition: LAD Stenosis

Narrowing of the left anterior descending coronary artery.

Technical Specification

Code: 28248000 Code System: SNOMED CT Short Name: ProxLAD Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Technical Specification

Element: 13302 Proximal Left Anterior

Descending Artery Disease Greater or Equal to 70 percent Not Documented

Operator: Equal

Value: No (or Not Answered)

Valid Range: Data Source: User

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

F I	Proximal Left Anterior Descending Artery Disease Greater or Equal to 70	Technical Specification
Element: 13302	percent Not Documented	Code: 28248000
On the standard on		Code System: SNOMED CT
Coding instruction:	Indicate whether the % stenosis of the proximal left anterior descending coronary artery was not documented.	Short Name: ProxLADND
	not documented.	Missing Data: Report
		Harvested: Yes (BDS, TAVR, TMVR,
Target Value:	N/A	TMVrpr, TTVP)
Supporting Definition:	LAD Stenosis	Is Identifier: No
3	Narrowing of the left anterior descending coronary artery.	Is Base Element: Yes
		Is Followup No
	Source:	Element:
		Data Type: BL
		Precision:
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13496 Syntax Score

Coding Instruction: Indicate the syntax score documented in the medical record. The syntax score is required for

patients with left main disease and/or 3 vessel disease in native coronary arteries.

SYNTAX (Synergy between PCI with TAXUS drug-eluting stent and Cardiac Surgery) Score: a

grading tool used to determine the complexity of CAD in native vessels.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 10001424796

Code System: ACC NCDR
Short Name: Syntax
Missing Data: Report

Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes

Is Followup Element: No Data Type: CD

Precision: Selection Type: Single

Unit of Measure:

Default Value: Null

Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13497 Syntax Score Not Documented

Operator: Equal

Value: No (or Not Answered)

Syntax Score Tiers - 1.3.6.1.4.1.19376.1.4.1.6.5.504

Selection	Definition	Source	Code	Code System
Low Syntax Score (<22)	Low Syntax Score(<22)		10001424799	ACC NCDR
Intermediate Syntax Score (232)	22- Intermediate Syntax Score (22-32)		10001424798	ACC NCDR
High Syntax Score (>= 33)	High Syntax Score (>= 33)		10001424797	ACC NCDR

Element: 13497 Syntax Score Not Documented

Coding Instruction: Indicate if the syntax score was not documented in the medical record.

Target Value: N/A

Technical Specification

Code: 10001424796
Code System: ACC NCDR
Short Name: SyntaxND
Missing Data: Report

Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
No

Element: NO
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13713 Cardiac Output

Coding Instruction: Indicate the cardiac output in L/min, documented by pre-procedure diagnostic cardiac cath

findings

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 82799009
Code System: SNOMED CT
Short Name: CardiacOutput

Missing Data: Report
Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: L/min
Default Value: Null
Usual Range: 2.0 - 8.0

Usual Range: 2.0 - 8.0 L/min Valid Range: 0.1 - 10.0 L/min

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Element: 13714 Cardiac Output Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13714 Cardiac Output Not Documented

Coding Instruction: Indicate if the cardiac output was not documented.

Target Value: N/A

Technical Specification

Code: 82799009
Code System: SNOMED CT
Short Name: CardiacOutput_ND

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy Procedure Type







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13715 Pulmonary Capillary Wedge Pressure

Coding Instruction: Indicate the pulmonary capillary wedge pressure, in mm Hg.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 118433006 Code System: SNOMED CT Short Name: PCWP Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: PQ Precision: 2,0 Selection Type: Single Unit of Measure: mm[Hg] Default Value: Null

Usual Range: 6 - 12 mm[Hg] Valid Range: 1 - 75 mm[Hg] Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----Element: 13716 Pulmonary Capillary Wedge

Pressure Not Documented

Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13716 Pulmonary Capillary Wedge Pressure Not Documented

Coding Instruction: Indicate if the pulmonary capillary wedge pressure was not documented.

Target Value: N/A

Technical Specification

Code: 118433006
Code System: SNOMED CT
Short Name: PCWP_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13719 Pulmonary Artery Mean Pressure

Coding Instruction: Indicate the pulmonary artery mean pressure, in mm Hg.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001423
Code System: ACC NCDR
Short Name: PAPMean
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 5 - 25 mm[Hg]

Usual Range: 5 - 25 mm[Hg]
Valid Range: 1 - 99 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Element: 13720 Pulmonary Artery Mean Pressure

Not Documented

Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13720 Pulmonary Artery Mean Pressure Not Documented

Coding Instruction: Indicate the pulmonary artery mean pressure, in mm Hg.

Target Value: N/A

Technical Specification

Code: 112000001423 Code System: ACC NCDR Short Name: PAPMean_ND Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13717 Pulmonary Artery Systolic Pressure

Coding Instruction: Indicate the pulmonary artery systolic pressure, in mm Hg.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 250768007 Code System: SNOMED CT Short Name: PAPSys Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mm[Hg] Default Value: Null Usual Range: 10 - 35 mm[Hg] Valid Range: 1 - 150 mm[Hg]

Data Source: User

Parent/Child Validation Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

----- AND -----Element: 13718 Pulmonary Artery Systolic

Pressure Not Documented

Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13718 Pulmonary Artery Systolic Pressure Not Documented

Coding Instruction: Indicate true if the pulmonary artery systolic pressure is not documented

Target Value: N/A

Technical Specification

Code: 250768007
Code System: SNOMED CT
Short Name: PAPSys_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14291 Pulmonary Vascular Resistance

Coding Instruction: Indicate the pulmonary vascular resistance in Woods units (mm Hg/L/min).

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 276901002
Code System: SNOMED CT
Short Name: PVR
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: Wood units

Usual Range: 0.10 - 10.00 Wood units **Valid Range:** 0.10 - 25.00 Wood units

Data Source: User

Default Value: Null

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14289 Pulmonary Vascular Resistance

Not Documented

Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 14289 Pulmonary Vascular Resistance Not Documented

Coding Instruction: Indicate if the pulmonary vascular resistance was not documented.

Target Value: N/A

Technical Specification

Code: 276901002 Code System: SNOMED CT

Short Name: PVRND Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No

Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null

Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14272 Right Atrial Pressure

Coding Instruction: Indicate the mean right atrial pressure (RAP) in mm Hg.

This can also documented as the central venous pressure (CVP).

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 276755008 Code System: SNOMED CT Short Name: RAP Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 2,0 Selection Type: Single Unit of Measure: mm[Hg] Default Value: Null

> Usual Range: 1 - 10 mm[Hg] Valid Range: 0 - 35 mm[Hg] Data Source: User

> > Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

----- AND -----

Element: 13829 Right Atrial Pressure Not

Documented Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13829 Right Atrial Pressure Not Documented

Coding Instruction: Indicate if the mean right atrial pressure pre-procedure, was not documented.

Target Value: N/A

Technical Specification

Code: 276755008
Code System: SNOMED CT
Short Name: MeanRAP_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13303 Right Ventricular Systolic Pressure

Coding Instruction: Indicate the right ventricular systolic pressure in mm Hg recorded prior to the start of the

procedure. Note: If more than one RVSP documented, code the highest value.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSP
Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 15 - 30 mm[Hg]

Usual Range: 15 - 30 mm[Hg]
Valid Range: 1 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13304 Right Ventricular Systolic

Pressure Not Documented

Operator: Equal







Section: Pre-Procedure Diagnostic Cath Findings

Parent: Presentation and Evaluation

Element: 13304 Right Ventricular Systolic Pressure Not Documented

Coding Instruction: Indicate if the right ventricular systolic pressure was not documented.

Target Value: N/A

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSYSND
Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure







Section: Pre-Procedure CTA Findings

Parent: Presentation and Evaluation

Element: 13422 Aortic Valve Annulus Assessment Method

Coding Instruction: Indicate the method used to assess the aortic valve annulus size.

Note: If the annulus was assessed with more than one method, code the findings based on computed tomography angiography (CTA). If CTA was not performed, code the measurement based on the assessment method (echo or other method) used to assess the annulus size to

determine the size of the prosthetic valve implanted during the procedure.

Target Value: The value on current procedure

Supporting Definition: AV Annulus Assessment Method

The imaging modality method used to assess the aortic valve annulus.

Source:

Technical Specification

Code: 112000001238

Code System: ACC NCDR

Short Name: AVDAnnulusSizeMethod

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes

Is Followup Element: No Data Type: CD Precision:

Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Imaging Modalities - 1.3.6.1.4.1.19376.1.4.1.6.5.486

Selection	Definition	Source	Code	Code System
Computed Tomography			418272005	SNOMED CT
Angiography Transthoracic Echo (TTE)			433236007	SNOMED CT
Transesophageal			105376000	SNOMED CT
Echocardiogram (TEE)				
Other			100000351	ACC NCDR

Element: 13428 Aortic Valve Annulus Minimum Diameter

Coding Instruction: Indicate the minimum diameter of the aortic valve annulus, in mm.

Note: Document aortic valve annulus measurements that are available, preferably measured

from a CT.

Target Value: The value on current procedure

Technical Specification

Code: 112000001804
Code System: ACC NCDR
Short Name: AVAnnulusDia
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: mm
Default Value: Null

Usual Range: 10.0 - 40.0 mm Valid Range: 5.0 - 80.0 mm Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure CTA Findings

Parent: Presentation and Evaluation

Element: 13429 Aortic Valve Annulus Maximum Diameter

Coding Instruction: Indicate the maximum diameter of the aortic valve annulus, in mm.

Note: Document aortic valve annulus measurements that are available, preferably measured

from a CT.

Target Value: The value on current procedure

Technical Specification

Code: 112000001241
Code System: ACC NCDR
Short Name: AVAnnulusMaxDia

Missing Data: Report Harvested: Yes (TAVR)

> Usual Range: 10.0 - 40.0 mm Valid Range: 5.0 - 80.0 mm

Data Source: User

Default Value: Null

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13438 Aortic Valve Annulus Area

Coding Instruction: Indicate the area of the aortic valve annulus, in mm2.

Note: Document aortic valve annulus measurements that are available, preferably measured

from a CT.

Target Value: The value on current procedure

Technical Specification

Code: 112000001251
Code System: ACC NCDR
Short Name: AVAnnulusArea

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 4,1
Selection Type: Single
Unit of Measure: mm2
Default Value: Null

Usual Range: 100.0 - 600.0 mm2 **Valid Range:** 100.0 - 999.0 mm2

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Pre-Procedure CTA Findings

Parent: Presentation and Evaluation

Element: 13439 Aortic Valve Annulus Perimeter

Coding Instruction: Indicate the perimeter of the aortic valve annulus, in mm.

Note: Document aortic valve annulus measurements that are available, preferably measured

from a CT.

Target Value: The value on current procedure

Technical Specification

Code: 112000001252
Code System: ACC NCDR
Short Name: AVAnnulusPeri

Missing Data: Report Harvested: Yes (TAVR)

Default Value: Null
Usual Range: 50.0 - 90.0 mm
Valid Range: 10.0 - 100.0 mm

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13423 Aortic Valve Calcification Severity

Coding Instruction: Indicate the degree of calcification on the aortic valve, documented by CT.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 18115005
Code System: SNOMED CT
Short Name: AVCalc
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13437 Aortic Valve Calcification

Severity Not Documented

Operator: Equal

Value: No (or Not Answered)
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Aortic Valve Calcification - 1.3.6.1.4.1.19376.1.4.1.6.5.489

Selection	Definition	Source	Code	Code System
None			112000001127	ACC NCDR
Minimal			112000001247	ACC NCDR
Moderate/Severe			112000001249	ACC NCDR







Section: Pre-Procedure CTA Findings

Parent: Presentation and Evaluation

Element: 13437 Aortic Valve Calcification Severity Not Documented

Coding Instruction: Indicate if the degree of calcification on the aortic valve was not documented.

Target Value: N/A

Technical Specification

Code: 18115005
Code System: SNOMED CT
Short Name: AVCalcND
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Left Ventricular Ejection

Parent: Pre-Procedure Echocardiogram Findings

Technical Specification Element: 13305 Left Ventricular Ejection Fraction Code: 10230-1 Coding Instruction: Indicate the percentage of the blood emptied from the left ventricle at the end of the Code System: LOINC Short Name: LVEFMeasure Target Value: The last value between 12 months prior to arrival and start of the first procedure Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, Supporting Definition: Most Recent LVEF % TMVrpr, TTVP) The left ventricular ejection fraction is the percentage of blood emptied from the left ventricle at Is Identifier: No the end of contraction. Is Base Element: Yes Is Followup No Source: ACC Clinical Data Standards, Society for Thoracic Surgeons Adult Cardiac Surgery Database (STS) Element: Data Type: PQ Precision: 2,0 Selection Type: Single Unit of Measure: % Default Value: Null

Parent/Child Validation

Element: 13306 Left Ventricular Ejection Fraction

Not Assessed

Usual Range: 5 - 90 % Valid Range: 1 - 99 % Data Source: User

Operator: Equal

Value: No (or Not Answered)

Element: 13306 Left Ventricular Ejection Fraction Not Assessed

Coding Instruction: Indicate whether the left ventricular ejection fraction was not assessed or not measured.

Target Value: N/A

Technical Specification

Code: 100001027
Code System: ACC NCDR
Short Name: LVEFNA
Missing Data: Report

 $\textbf{Harvested:} \ \ \text{Yes (BDS, TAVR, TMVR,} \\$

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User







Section: Left Ventricular Dimension

Parent: Pre-Procedure Echocardiogram Findings

Element: 13721 Left Ventricular Internal Systolic Dimension

Coding Instruction: Indicate the left ventricular internal systolic dimension in cm.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001424
Code System: ACC NCDR
Short Name: LVIDs
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,1
Selection Type: Single
Unit of Measure: cm
Default Value: Null
Usual Range: 2.5 - 4.5 cm
Valid Range: 1.0 - 9.0 cm
Data Source: User

Parent/Child Validation

Element: 13722 Left Ventricular Internal Systolic

Dimension Not Measured

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13722 Left Ventricular Internal Systolic Dimension Not Measured

Coding Instruction: Indicate if the left ventricular internal systolic dimension was not measured.

Target Value: N/A

Technical Specification

Code: 112000001424
Code System: ACC NCDR
Short Name: LVIDs_NM
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Left Ventricular Dimension

Parent: Pre-Procedure Echocardiogram Findings

Element: 13723 Left Ventricular Internal Diastolic Dimension

Coding Instruction: Indicate the left ventricular internal diastolic dimension in cm.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

 Code:
 112000001425

 Code System:
 ACC NCDR

 Short Name:
 LVIDd

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: cm
Default Value: Null
Usual Range: 3.5 - 5.5 cm
Valid Range: 1.0 - 10.0 cm
Data Source: User

Parent/Child Validation

Element: 13724 Left Ventricular Internal Diastolic

Dimension Not Measured

Operator: Equal

Value: No (or Not Answered)

----- AND -----

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

value: IMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13724 Left Ventricular Internal Diastolic Dimension Not Measured

Coding Instruction: Indicate if the left ventricular internal diastolic dimension was not measured.

Target Value: N/A

Technical Specification

Code: 112000001425
Code System: ACC NCDR
Short Name: LVIDd_NM
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Left Ventricular Dimension

Parent: Pre-Procedure Echocardiogram Findings

Element: 13725 Left Ventricular End Systolic Volume

Coding Instruction: Indicate the left ventricular end systolic volume in ml documented by echocardiogram.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 250931004
Code System: SNOMED CT
Short Name: LVESV
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mL
Default Value: Null
Usual Range: 10 - 150 mL
Valid Range: 1 - 300 mL
Data Source: User

Parent/Child Validation

Element: 13727 Left Ventricular End Systolic

Volume Not Measured

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13727 Left Ventricular End Systolic Volume Not Measured

Coding Instruction: Indicate if the left ventricular end systolic volume was not measured.

Target Value: N/A

Technical Specification

Code: 250931004
Code System: SNOMED CT
Short Name: LVESV_NM
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Left Ventricular Dimension

Parent: Pre-Procedure Echocardiogram Findings

Element: 13726 Left Ventricular End Diastolic Volume

Coding Instruction: Indicate the left ventricular end diastolic volume in ml, documented by echocardiogram. Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 250932006 Code System: SNOMED CT Short Name: LVEDV Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mL Default Value: Null Usual Range: 40 - 250 mL Valid Range: 1 - 400 mL Data Source: User

Parent/Child Validation

Element: 13728 Left Ventricular End Diastolic

Volume Not Measured

Operator: Equal

Value: No (or Not Answered)

----- AND -----

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13728 Left Ventricular End Diastolic Volume Not Measured

Coding Instruction: Indicate if the left ventricular end diastolic volume was not measured.

Target Value: N/A

Technical Specification

Code: 250932006 Code System: SNOMED CT Short Name: LVEDV_NM Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type



Element: 13729



Full Specifications **Data Dictionary v3.0**



Section: Left Atrial Volume

Parent: Pre-Procedure Echocardiogram Findings

Coding Instruction: Indicate the left atrial volume in ml documented by echocardiogram.

Left Atrial Volume

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification
Code: 112000001426
Code System: ACC NCDR

Short Name: LAVol Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Precision: 3,0
Selection Type: Single
Unit of Measure: mL
Default Value: Null
Usual Range: 10 - 90 mL
Valid Range: 1 - 500 mL
Data Source: User

Parent/Child Validation

Element: 13730 Left Atrial Volume Not Measured

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13730 Left Atrial Volume Not Measured

Coding Instruction: Indicate if the left atrial volume was not measured.

Target Value: N/A

Technical Specification

Code: 112000001426
Code System: ACC NCDR
Short Name: LAVol_NM
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Left Atrial Volume

Parent: Pre-Procedure Echocardiogram Findings

Element: 13731 Left Atrial Volume Index

Coding Instruction: Indicate the left atrial volume index in mL/m2, documented by echocardiogram.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001427
Code System: ACC NCDR
Short Name: LAVolIndex
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: ml/m2
Default Value: Null
Usual Range: 10 - 90 ml/m2
Valid Range: 1 - 250 ml/m2

Parent/Child Validation

Element: 13732 Left Atrial Volume Index Not

Measured

Data Source: User

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13732 Left Atrial Volume Index Not Measured

Coding Instruction: Indicate if the left atrial volume index was not measured.

Target Value: N/A

Technical Specification

Code: 112000001427
Code System: ACC NCDR
Short Name: LAVolIndex_NM
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13442 Aortic Valve Disease Etiology

Coding Instruction: Indicate primary etiology of aortic valve disease.

Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure

Supporting Definition: Aortic Valve Disease Etiology

The cause of aortic valve disease.

Source:

Technical Specification

Code: 112000001253
Code System: ACC NCDR
Short Name: VDAoEt
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Aortic Valve Disease Etiology - 1.3.6.1.4.1.19376.1.4.1.6.5.493

Selection	Definition	Source	Code	Code System
Degenerative			112000001254	ACC NCDR
Endocarditis			56819008	SNOMED CT
Rheumatic			58718002	SNOMED CT
Other			100000351	ACC NCDR

Element: 13468 Aortic Valve Morphology

Coding Instruction: Indicate the morphology of the aortic valve.

If a patient was born with a tricuspid valve with two leaflets that are fused, code tricuspid.

Target Value: The value at birth

Supporting Definition: Aortic Valve Disease

A disorder characterized by a defect in aortic valve function or structure.

Source:

Technical Specification

Code: 8722008

Code System: SNOMED CT
Short Name: AVMorphology
Missing Data: Report

Harvested: Yes (BDS, TAVR) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Aortic Valve Disease Morphology - 1.3.6.1.4.1.19376.1.4.1.6.5.495

Selection	Definition	Source	Code	Code System
Bicuspid Aortic Valve			72352009	SNOMED CT
Tricuspid Valve			46030003	SNOMED CT
Other			100000351	ACC NCDR







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13469 Ascending Aorta Size

Coding Instruction: Indicate the size, in cm, of the ascending aorta.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Supporting Definition: Ascending Aorta Measurement

Quantitative measurement of the ascending aorta.

Source:

Technical Specification

Code: 112000001258 Code System: ACC NCDR Short Name: AASize Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: PQ Precision: 3,1 Selection Type: Single Unit of Measure: cm Default Value: Null Usual Range: 0.2 - 8.0 cm Valid Range: 0.0 - 12.0 cm

Data Source: User

Parent/Child Validation

----- AND -----

Element: 13468 Aortic Valve Morphology

Operator: Equal

Value: Bicuspid Aortic Valve

Element: 13470 Ascending Aorta Size Not

Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13470 Ascending Aorta Size Not Documented

Coding Instruction: Indicate if the size of the ascending aorta was not documented in the medical record.

Target Value: N/A

Supporting Definition: Ascending Aorta Measurement

Quantitative measurement of the ascending aorta.

Technical Specification

Code: 112000001258 Code System: ACC NCDR Short Name: AASizeND Missing Data: Report Harvested: Yes (TAVR) Is Identifier: No

Is Base Element: Yes Is Followup Element: No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 13468 Aortic Valve Morphology

Operator: Equal

Value: Bicuspid Aortic Valve







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13471 Aortic Valve Annular Calcification

Coding Instruction: Indicate if annular calcification is present on the aortic valve.

Code yes if echo reports document calcification in the aortic valve leaflets, aorta adjacent to the AV, leaflets or the left ventricular outflow tract (LVOT), or if echo reports document AV

calcific degeneration.

Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 18115005
Code System: SNOMED CT
Short Name: AVAnnularCalc

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13477 Aortic Valve Regurgitation

Coding Instruction: Indicate the severity of aortic valve regurgitation.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 60234000
Code System: SNOMED CT
Short Name: VDInsufA
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System	
None			112000001910	ACC NCDR	
Trace/Trivial			112000001911	ACC NCDR	
Mild			112000000380	ACC NCDR	
Moderate			112000000381	ACC NCDR	
Severe			112000000382	ACC NCDR	







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13307 **Technical Specification** Aortic Stenosis Code: 60573004 Coding Instruction: Indicate whether aortic stenosis is present. Code System: SNOMED CT Short Name: VDStenA Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Element: 13481 Aortic Valve Area

Coding Instruction: Indicate the smallest aortic valve area (in cm squared) obtained from an echocardiogram or

cath report.

Target Value: The lowest value between 12 months prior to start of procedure and start of procedure

Technical Specification

Code: 112000001280
Code System: ACC NCDR
Short Name: VDAoVA
Missing Data: Report

Valid Range: Data Source: User

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null
Usual Range: 0.20 - 4.00 cm2

Valid Range: 0.20 - 4.00 cm2

Valid Range: 0.05 - 5.00 cm2

Data Source: User

Parent/Child Validation
Element: 13307 Aortic Stenosis

Operator: Equal Value: Yes

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13674 Aortic Valve Mean Gradient

Coding Instruction: Indicate the highest MEAN gradient (in mm Hg) across the aortic valve.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001398 Code System: ACC NCDR Short Name: VDGradA Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mm[Hg] Default Value: Null Usual Range: 5 - 50 mm[Hg]

Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13307 Aortic Stenosis

Operator: Equal Value: Yes

AND Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13700 Low Flow

Coding Instruction: Indicate if there was low flow, which is defined as a stroke volume index <35 ml/m2.

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 21762000 Code System: SNOMED CT Short Name: SVI Missing Data: Report Harvested: Yes (TAVR) Is Identifier: No Is Base Element: Yes

Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation Element: 13674 Aortic Valve Mean Gradient

Operator: Less Than

Value: 40

----- AND -----Element: 13701 Low Flow Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System	
No			100013073	ACC NCDR	
Yes			100013072	ACC NCDR	







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13701 Low Flow Not Documented

Coding Instruction: Indicate if the stroke volume index was not documented.

Target Value: N/A

Technical Specification

Code: 112000001830
Code System: ACC NCDR
Short Name: SVI_ND
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13674 Aortic Valve Mean Gradient

Operator: Less Than Value: 40

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13702 Aortic Valve Peak Gradient

Coding Instruction: Indicate the aortic valve peak gradient in mm Hg.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001413
Code System: ACC NCDR
Short Name: AVPeakGrad
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup

Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 5 - 70 mm[Hg]
Valid Range: 0 - 200 mm[Hg]
Data Source: User

Parent/Child Validation

Element: 13307 Aortic Stenosis

Operator: Equal Value: Yes

Element: 14273 Transcatheter Valve Therapy

Procedure Type







Section: Aortic Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13703 Aortic Valve Peak Velocity

Coding Instruction: Indicate the aortic valve peak velocity, in meters per second, as determined by continuous

wave (CW) spectral velocity recording on echocardiography.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001414
Code System: ACC NCDR
Short Name: AVDPeakVelocity

Missing Data: Report Harvested: Yes (TAVR)

Usual Range: 1.0 - 4.0 m/sec Valid Range: 1.0 - 8.0 m/sec

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type



Element: 13704



Full Specifications Data Dictionary v3.0



Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Mitral Valve Disease Coding Instruction: Indicate whether mitral valve disease is present.

If there was no documentation of mitral valve disease, code no.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification Code: 11851006

Code System: SNOMED CT Short Name: MVD Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR.

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Valid Range: Data Source: User

Element: 13672 Mitral Regurgitation

Coding Instruction: Indicate the severity of regurgitation through the mitral valve.

Code the highest value or most severe regurgitation when a range is reported.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 48724000 Code System: SNOMED CT Short Name: PreprocMR Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 13704 Mitral Valve Disease

Operator: Equal Value: Yes

Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.728

Regulgitation Severity - 1.3.6.1.4.1.19376.1.4.1.0.3.726				
Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Moderate-Severe			1000142345	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13733 Paravalvular Mitral Regurgitation

Coding Instruction: Indicate the severity of paravalvular mitral regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: VDInsufMPara
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

----- AND -----

Element: 13734 Paravalvular Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Mild

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Moderate

Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13672 Mitral Regurgitation

Operator: Equal
Value: Trace/Trivial

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13734 Paravalvular Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular mitral regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: VDInsufMPara_ND

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Mild

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Moderate

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13735 Central Mitral Regurgitation

Coding Instruction: Indicate the severity of central mitral regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001433 Code System: ACC NCDR Short Name: VDInsuffMCentral

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Parent/Child Validation

Element: 13672 Mitral Regurgitation

Data Source: User

Operator: Equal Value: Mild

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Moderate

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

----- AND -----Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVR

----- AND -----Element: 13736 Central Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

valve regulgitation deventy 4 - 1.5.6.1.4.1.15076.1.4.1.1.55760				
Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13736 Central Regurgitation Not Documented

Coding Instruction: Indicate whether the severity of central regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001433 Code System: ACC NCDR

Short Name: VDInsuffMCentral_ND

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Florenty 13672 Mitrol Degraphitation

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Moderate

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13672 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13672 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13737 Effective Regurgitant Orifice Area

Coding Instruction: Indicate the effective regurgitant orifice area (EROA), in cm2.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001437 Code System: ACC NCDR Short Name: VDMitEOA Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 2,1 Selection Type: Single Unit of Measure: cm2 Default Value: Null **Usual Range:** 0.1 - 5.0 cm2 Valid Range: 0.1 - 5.0 cm2 Data Source: User

Parent/Child Validation

Flement: 13704 Mitral Valve Disease

Operator: Equal Value: Yes

AND Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13738 Effective Regurgitant Orifice Area Method of Assessment

Coding Instruction: Indicate the method used to assess the effective regurgitant orifice area. If multiple methods

are available, code the 3D planimetry method first, then PISA.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001437 Code System: ACC NCDR Short Name: VDMitEOA_MoA Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision:

Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13737 Effective Regurgitant Orifice Area

Operator:

Value: Any Value

Effective Regurgitant Orifice Area Method of Assessment - 1 3 6 1 4 1 19376 1 4 1 6 5 547

Lifective Regulgita	Trective Regulgitant Offfice Area Method of Assessment - 1.3.6.1.4.1.1937 6.1.4.1.0.3.347			
Selection	Definition	Source	Code	Code System
3D Planimetry			112000001438	ACC NCDR
Proximal Isovelocity Surface Area		112000001439	ACC NCDR	
Quantitative Doppler		112000001440	ACC NCDR	
Other			100000351	ACC NCDR



Element: 13308



Full Specifications **Data Dictionary v3.0**



Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

 $\begin{tabular}{ll} \textbf{Coding Instruction:} & Indicate whether mitral stenosis is present. \\ \end{tabular}$

Mitral Stenosis

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 79619009
Code System: SNOMED CT
Short Name: VDStenM
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Operator: Equal Value: Yes

Element: 13316 Mitral Valve Area

Coding Instruction: Indicate the smallest mitral valve area in centimeters squared.

Target Value: The lowest value between 12 months prior to start of procedure and start of procedure

Supporting Definition: Mitral Valve Area

Measurement of mitral valve area.

Source:

Technical Specification

Code: 251012002
Code System: SNOMED CT
Short Name: VDMVA
Missing Data: Report

Element: 13704 Mitral Valve Disease

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 3.00 - 6.00 cm2 **Valid Range:** 0.05 - 12.00 cm2

Data Source: User

Parent/Child Validation

Element: 13704 Mitral Valve Disease

Operator: Equal Value: Yes







Section: Mitral Valve Disease

Parent: Pre-Procedure Echocardiogram Findings

Element: 13317 Mitral Valve Mean Gradient

Coding Instruction: Indicate the highest mean gradient (in mm Hg) across the mitral valve.

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Supporting Definition: Mitral Valve Mean Gradient

The average gradient across the mitral valve occurring during the entire systole.

Source: Baumgartner, H. et. al (2009). Echocardiographic assessment of valve stenosis:

EAE/ASE

recommendations for clinical practice.

Technical Specification

Code: 112000001191
Code System: ACC NCDR
Short Name: VDGradM
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

 Is Base Element:
 Yes

 Is Followup Element:
 No

 Data Type:
 PQ

 Precision:
 3,0

 Selection Type:
 Single

 Unit of Measure:
 mm[Hg]

 Default Value:
 Null

 Usual Range:
 5 - 50 mm[Hg]

Is Identifier: No

Valid Range: 5 - 50 mm[Hg]
Valid Range: 0 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13704 Mitral Valve Disease
Operator: Equal
Value: Yes







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13490 Technical Specification Mitral Valve Disease Etiology Code: 11851006 Coding Instruction: Indicate the etiology of mitral valve disease. Code System: SNOMED CT Short Name: MVDEtio Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure Missing Data: Report Supporting Definition: Mitral Valve Disease Harvested: Yes (BDS, TAVR, TMVR, A disorder characterized by a defect in mitral valve function or structure.

Source: NCI Thesaurus

Vendor Instruction: When Mitral Valve Disease Etiology (13490) is Equal to (None) then Transcatheter Valve Therapy Procedure Type (14273) must be not Equal to (TMVR,TMVr)

Cannot select option None with any other option: Functional MR (Secondary), Degenerative MR

(Primary), Post Inflammatory, Endocarditis or Other

TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision:

Selection Type: Multiple

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Mitral Valve Disease Etiology - 1.3.6.1.4.1.19376.1.4.1.6.5.548

Selection	Definition	Source	Code	Code System
Functional MR (Secondary)	Typically the valve structures (i.e leaflets and che tendinae) are normal in functional mitral regurgitate but a variety of diseases (such as a prior myocarcinfarction or cardiomyopathy) compromises the leability to coapt (i.e. form a tight seal when closed) results in mitral regurgitation.	on, dial aflets	112000001276	ACC NCDR
Degenerative MR (Primary)	Degenerative mitral valve disease is due to multiple conditions that lead to abnormal leaflets and/or ch that result and mitral regurgitation. The leaflets ma prolapse or flail into the left atrium.	ordae	112000001277	ACC NCDR
Post Inflammatory			112000001441	ACC NCDR
Endocarditis			56819008	SNOMED CT
Other			100000351	ACC NCDR
None			100001231	ACC NCDR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13740 Functional Mitral Valve Regurgitation Type

Coding Instruction: Indicate the type of functional mitral regurgitation.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Supporting Definition: Functional Mitral Valve Regurgitation

Typically the valve structures (i.e.. leaflets and chord tendinae) are normal in functional mitral regurgitation, but a variety of diseases (such as a prior myocardial infarction or cardiomyopathy) compromises the leaflets ability to coapt (i.e. form a tight seal when closed)

and results in mitral regurgitation.

Source:

Technical Specification

Code: 112000001276 Code System: ACC NCDR Short Name: FMRType Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Functional MR (Secondary)

Element: 13741 Functional Mitral Valve

----- AND -----Regurgitation Type Not Documented

Operator: Equal

Value: No (or Not Answered)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Functional Mitral Valve Regurgitation - 1.3.6.1.4.1.19376.1.4.1.6.5.549

Selection	Definition	Source Code	Code System
Ischemic Acute, Post Infarction	The patient has a new onset of mitral regurgitation that occurs within weeks of having of having a myocardial infarction.		ACC NCDR
Ischemic Chronic		112000001443	ACC NCDR
Non-Ischemic Dilated Cardiomyopathy		195021004	SNOMED CT
Restrictive Cardiomyopathy		415295002	SNOMED CT
Hypertrophic Cardiomyopathy		233873004	SNOMED CT
Pure Annular Dilation with Normal Left Ventricular Systolic Function		112000001444	ACC NCDR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13741 Functional Mitral Valve Regurgitation Type Not Documented

Coding Instruction: Indicate whether the type of functional mitral regurgitation was not documented.

Target Value: N/A

Supporting Definition: Functional Mitral Valve Regurgitation

Typically the valve structures (i.e.. leaflets and chord tendinae) are normal in functional mitral regurgitation, but a variety of diseases (such as a prior myocardial infarction or cardiomyopathy) compromises the leaflets ability to coapt (i.e. form a tight seal when closed)

and results in mitral regurgitation.

Source:

Technical Specification

Code: 112000001276
Code System: ACC NCDR
Short Name: FMRType_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Functional MR (Secondary)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13742 Leaflet Prolapse

Coding Instruction: Indicate if there was leaflet prolapse.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001445
Code System: ACC NCDR
Short Name: MVDLeafPro
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Degenerative MR (Primary)

AND ----
Element: 13745 Leaflet Prolapse Not Documented

Operator: Equal

Value: No (or Not Answered)
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Valve Leaflet Type - 1.3.6.1.4.1.19376.1.4.1.6.5.550

Selection	Definition	Source	Code	Code System
None			100001231	ACC NCDR
Anterior Leaflet			112000001449	ACC NCDR
Posterior Leaflet			112000001450	ACC NCDR
Bileaflet			112000001446	ACC NCDR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13745 Leaflet Prolapse Not Documented

Coding Instruction: Indicate if leaflet prolapse was not documented.

Target Value: N/A

Technical Specification

Code: 112000001445
Code System: ACC NCDR
Short Name: MVDLeafPro_ND

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Degenerative MR (Primary)
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR



Element: 13743



Full Specifications **Data Dictionary v3.0**



Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Coding Instruction: Indicate if there was leaflet flail.

Leaflet Flail

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001447
Code System: ACC NCDR
Short Name: MVDLeafFlail
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Degenerative MR (Primary)

Element: 13746 Leaflet Flail Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Valve Leaflet Type - 1.3.6.1.4.1.19376.1.4.1.6.5.550

Selection	Definition	Source	Code	Code System
None			100001231	ACC NCDR
Anterior Leaflet			112000001449	ACC NCDR
Posterior Leaflet			112000001450	ACC NCDR
Bileaflet			112000001446	ACC NCDR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13746 Leaflet Flail Not Documented

Coding Instruction: Indicate if leaflet flail was not documented.

Target Value: N/A

Technical Specification

Code: 112000001447
Code System: ACC NCDR
Short Name: MVDLeafFlail_ND

Missing Data: Report

Is Identifier: No

Harvested: Yes (TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Degenerative MR (Primary)
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13748 Inflammatory Mitral Valve Disease Type

Coding Instruction: Indicate type of inflammatory mitral valve disease.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001451
Code System: ACC NCDR
Short Name: InflamType
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Post Inflammatory
AND

Element: 13753 Inflammatory Mitral Valve Disease

Type Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Inflammatory Mitral Valve Disease Type - 1.3.6.1.4.1.19376.1.4.1.6.5.551

Selection	Definition	Source	Code	Code System
Collagen Vascular Dis	ease		398049005	SNOMED CT
Drug Induced			112000001454	ACC NCDR
Idiopathic			112000001453	ACC NCDR
Prior Radiation Therap	у		112000001455	ACC NCDR
Rheumatic Fever			58718002	SNOMED CT







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13753 Inflammatory Mitral Valve Disease Type Not Documented

Coding Instruction: Indicate if the type of inflammatory mitral valve disease was not documented.

Target Value: N/A

Technical Specification

Code: 112000001451
Code System: ACC NCDR
Short Name: InflamType_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13490 Mitral Valve Disease Etiology

Operator: Equal

Value: Post Inflammatory
AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13744 Leaflet Tethering

Coding Instruction: Indicate if there was leaflet tethering.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification
Code: 112000001448
Code System: ACC NCDR
Short Name: MVDLeafTeth

Missing Data: Report
Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

AND ----
Element: 13747 Leaflet Tethering Not Documented

Operator: Equal

Value: No (or Not Answered)

Valve Leaflet Type - 1.3.6.1.4.1.19376.1.4.1.6.5.550

Turvo Eduliot Typo Tidiot 14 Titologood				
Selection	Definition	Source	Code	Code System
None			100001231	ACC NCDR
Anterior Leaflet			112000001449	ACC NCDR
Posterior Leaflet			112000001450	ACC NCDR
Bileaflet			112000001446	ACC NCDR

Element: 13747 Leaflet Tethering Not Documented

Coding Instruction: Indicate if leaflet tethering was not documented.

Target Value: N/A

Technical Specification

Code: 112000001448
Code System: ACC NCDR
Short Name: MVDLeafTeth_ND

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal

Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13749 Mitral Valve Annular Calcification

Coding Instruction: Indicate if there was mitral annular calcification.

Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 251002009
Code System: SNOMED CT
Short Name: MVDAnnular
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

AND -----

Element: 13750 Mitral Valve Annular Calcification

Not Documented

Operator: Equal

Value: No (or Not Answered)

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Element: 13750 Mitral Valve Annular Calcification Not Documented

Coding Instruction: Indicate if mitral annular calcification was not documented.

Target Value: N/A

Technical Specification

Code: 251002009
Code System: SNOMED CT
Short Name: MVCalcND
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr) **Is Identifier:** No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr







Section: Mitral Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13751 Mitral Leaflet Calcification

Coding Instruction: Indicate if there was mitral leaflet calcification.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001452
Code System: ACC NCDR
Short Name: MLeafCalc
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

----- AND -----

Element: 13752 Mitral Leaflet Calcification Not

Documented
Operator: Equal

Value: No (or Not Answered)

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR

Element: 13752 Mitral Leaflet Calcification Not Documented

Coding Instruction: Indicate if mitral calcification was not documented.

Target Value: N/A

Technical Specification

Code: 112000001452
Code System: ACC NCDR
Short Name: MLeafCalc_ND
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr) Is Identifier: No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal

Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr







Section: Tricuspid Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13806 Tricuspid Valve Disease Etiology

Coding Instruction: Indicate the etiology of tricuspid valve disease.

Target Value: Any occurrence between 12 months prior to the procedure and start of the procedure

Supporting Definition: Tricuspid Valve

A three-cusp valve of the heart that regulates the flow of blood between the right atrium and

the right ventricle of the heart

Source:

Technical Specification

Code: 46030003 Code System: SNOMED CT Short Name: TVDisEtio Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Tricuspid Valve Disease Etiology - 1.3.6.1.4.1.19376.1.4.1.6.5.563

Selection	Definition	Source Code	Code System
Primary	Valve structures are abnormal and the abnormalities cause the valve disease.	112000001509	ACC NCDR
Secondary	Valve structures are normal but other conditions (e.g myocardial infarction or cardiomyopathy) compromise the valve's ability to function normally.		ACC NCDR
Pacemaker Induced		112000001511	ACC NCDR
Other		100000351	ACC NCDR

Element: 13318	Tricuspid Valve Regurgitation	Technical Specification
		Code: 111287006
Coding Instruction:	: Indicate whether there is evidence of tricuspid valve regurgitation. Enter level of valve function	Code System: SNOMED CT
	associated with highest risk (i.e., worst performance).	Short Name: PreprocTR
	If there was no documentation of tricuspid valve disease, code none.	Missing Data: Report
	in the first the decementation of the depth takes discuss, seed from	Harvested: Yes (BDS, TAVR, TMVR,

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range:

Data Source: User

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Tricuspid Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13807 Tricuspid Valve Diastolic Gradient

Coding Instruction: Indicate the tricuspid valve diastolic gradient in mm Hg. This can also be called the TV inflow

gradien

Target Value: The highest value between 12 months prior to the procedure and start of the procedure

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: TVDGrad
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 1 - 15 mm[Hg] Valid Range: 1 - 50 mm[Hg] Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND

Element: 13810 Tricuspid Valve Diastolic Gradient Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13810 Tricuspid Valve Diastolic Gradient Not Documented

Coding Instruction: Indicate if the tricuspid valve diastolic gradient was not documented.

Target Value: N/A

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: TVDGradND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Tricuspid Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13808 Tricuspid Valve Annulus Size

Coding Instruction: Indicate the tricuspid valve annulus size in mm. Document the size using end-diastolic, 4

chamber view is preferred (in mm).

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001513
Code System: ACC NCDR
Short Name: TVAnnulus

Missing Data: Report Harvested: Yes (TTVP) Is Identifier: No

 Is Base Element:
 Yes

 Is Followup Element:
 No

 Data Type:
 PQ

 Precision:
 2,0

Precision: 2,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 15 - 60 mm
Valid Range: 1 - 80 mm

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13809 Tricuspid Valve Annulus Size Not

Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13809 Tricuspid Valve Annulus Size Not Documented

Coding Instruction: Indicate if the tricuspid valve annulus size was not documented.

Target Value: N/A

Technical Specification

Code: 112000001513
Code System: ACC NCDR
Short Name: TVAnnulusND
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Tricuspid Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13811 End Diastolic Mid Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic mid-RV diameter, using the 4 chamber view (in cm).

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001514
Code System: ACC NCDR
Short Name: MidRVDia
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,1
Selection Type: Single
Unit of Measure: cm
Default Value: Null
Usual Range: 1.0 - 7.0 cm
Valid Range: 0.1 - 9.9 cm
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND ------ Element: 13812 End Diastolic Mid Right Ventricle

Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13812 End Diastolic Mid Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the end-diastolic mid-RV diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001514
Code System: ACC NCDR
Short Name: MidRVDiaND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL

Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Tricuspid Valve Disease Etiology

Parent: Pre-Procedure Echocardiogram Findings

Element: 13813 End Diastolic Basal Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic basal RV diameter, using the 4 chamber view (in cm).

Target Value: The last value between 12 months prior to arrival and start of the first procedure

Technical Specification

Code: 112000001515 Code System: ACC NCDR Short Name: BasalDia Missing Data: Report

Harvested: Yes (TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 2,1 Selection Type: Single Unit of Measure: cm Default Value: Null Usual Range: 1.0 - 7.0 cm

Valid Range: 0.1 - 9.9 cm Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----Element: 13814 End Diastolic Basal Right

Ventricle Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13814 End Diastolic Basal Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the basal diastolic mid-RV diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001515 Code System: ACC NCDR Short Name: BasalDiaND Missing Data: Report Harvested: Yes (TTVP) Is Identifier: No Is Base Element: Yes Is Followup No

Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Pre-Procedure Dobutamine Challenge

Parent: Presentation and Evaluation

 Element: 13319
 Dobutamine Challenge Performed

 Coding Instruction:
 Indicate if a dobutamine challenge was performed.

 A dobutamine challenge is a type of stress echocardiography that can distinguish between true-severe versus pseudo-severe aortic stenosis.

 Target Value:
 Any occurrence between 12 months prior to arrival and start of the first procedure

 Supporting Definition:
 Dobutamine Stress Echocardiography

Code: 703338002
Code System: SNOMED CT
Short Name: DobutChal
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Technical Specification

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13320 Flow Reserve Present

Coding Instruction: Indicate if coronary flow reserve was documented on the dobutamine challenge.

Flow reserve on dobutamine echocardiogram is indicated by an increase in the stroke volume

A pharmacologic stress echocardiography technique to detect coronary artery disease and

index by ≥20%.

myocardial ischemia.

Source:

Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure

Supporting Definition: Dobutamine Stress Echocardiography Findings

The results or findings of dobutamine stress echocardiogram.

Source:

Technical Specification

Code: 112000001193
Code System: ACC NCDR
Short Name: FlowRes
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13319 Dobutamine Challenge Performed

Operator: Equal Value: Yes







Section: Pre-Procedure Dobutamine Challenge

Parent: Presentation and Evaluation

Element: 13321 Aortic Stenosis Type

Coding Instruction: Indicate the type of aortic stenosis documented on dobutamine challenge. Physicians may use

different criteria to differentiate, characterize and document truly severe aortic or pseudo-

severe aortic stenosis

The 2017 AUC for Severe Aortic Stenosis guideline differentiates "truly severe aortic stenosis"

with an AVA <=1.0 cm2 and Vmax >4 m/sec at any flow rate.

Target Value: Any occurrence between 12 months prior to arrival and start of the first procedure

Supporting Definition: Dobutamine Stress Echocardiography Findings

The results or findings of dobutamine stress echocardiogram.

Technical Specification

Code: 112000002013

Code System: ACC NCDR Short Name: ASType

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No Is Base Element: Yes

Is Followup No Element:

Data Type: CD Precision:

Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User

> > Parent/Child Validation

Element: 13319 Dobutamine Challenge Performed

Operator: Equal Value: Yes

----- AND -----

Element: 13325 Aortic Stenosis Type Not

Documented Operator: Equal

Value: No (or Not Answered)

Aortic Stenosis Type - 1.3.6.1.4.1.19376.1.4.1.6.5.462

Selection	Definition	Source	Code	Code System
Truly Severe Aortic Stenos	is		112000001194	ACC NCDR
Pseudo-Severe Aortic			112000001195	ACC NCDR
Stenosis				

Element: 13325 Aortic Stenosis Type Not Documented

Coding Instruction: Indicate if the type of aortic stenosis is not documented on dobutamine challenge.

Target Value: N/A

Supporting Definition: Dobutamine Stress Echocardiography Findings

The results or findings of dobutamine stress echocardiogram.

Source:

Technical Specification

Code: 112000002013

Code System: ACC NCDR Short Name: ASTypeND Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No. Is Base Element: Yes Is Followup No Element:

Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Dobutamine Challenge Performed Element: 13319

Operator: Equal Value: Yes







Section: Procedure Information Parent: Lab Visit

Element: 7065 **Technical Specification** Concomitant Procedures Performed Code: 100001271 Coding Instruction: Indicate if another procedure was performed concurrently. Code System: ACC NCDR Short Name: ConcomProc Target Value: The value on current procedure Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null

Element: 7066 Concomitant Procedures Performed Type

Coding Instruction: Indicate the type of procedure performed in conjunction with the TVT procedure.

Note(s):

The procedure(s) collected in your application is controlled by Procedure Master file. This file is maintained by the TVT Registry and will be made available on the internet for downloading and

importing/updating into your application.

Target Value: The value on current procedure

Technical Specification

Code: 100013075
Code System: ACC NCDR
Short Name: ConcomProcType

Missing Data: Report

Usual Range: Valid Range: Data Source: User

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Multiple (Dynamic List)

Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 7065 Concomitant Procedures

Performed

Operator: Equal Value: Yes

Concomitant Procedures Type - 2.16.840.1.113883.3.3478.6.4.10

Selection	Definition	Source	Code	Code System
Left Atrial Appendage Occlusion			233032004	SNOMED CT
Peripheral Intervention			100001272	ACC NCDR
Procedure Type Not Listed			10001424810	ACC NCDR
PCI			415070008	SNOMED CT
Permanent Pacemaker			449397007	SNOMED CT
Balloon Mitral Valvuloplasty			112000001951	ACC NCDR
BASILICA	Bioprosthetic Aortic Scallop Intentional Laceration to prevent latrogenic Coronary Artery obstruction (BASILICA) is a procedure that prevents coronary artery obstruction during transcatheter aortic valve replacement (TAVR).		112000001952	ACC NCDR
Alcohol Septal Ablation			437746009	SNOMED CT
LAMPOON	Laceration of the Anterior Mitral Valve Leaflet to Prevent Left Ventricular Outflow Tract Obstruction During Transcatheter Mitral Valve Replacement		112000001953	ACC NCDR



Element: 7025



Full Specifications Data Dictionary v3.0



Section: Procedure Information	Parent: Lab Visit

Procedure Status Coding Instruction: Indicate the status of the procedure.

Target Value: The value on current procedure

Vendor Instruction: When a Transcatheter Valve Therapy Procedure Type (14273) is selected Procedure Status

(7025) cannot be Null

Technical Specification Code: 100001218 Code System: ACC NCDR Short Name: ProcStatus Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Is Identifier: No

Procedure Status - 1.3.6.1.4.1.19376.1.4.1.6.5.226

Selection	Definition	Source	Code	Code System
Elective Procedure	The patient's cardiac function has been stable in the days or weeks prior to the procedure. The procedure could be deferred without increased risk of compromised cardiac outcome.	Society of Thoracic Surgeons (STS)	71388002:260870009=103390000	SNOMED CT
Urgent Procedure			71388002:260870009=103391001	SNOMED CT
Emergency Procedure	Patients requiring emergency operations will have ongoing, refractory (difficult, complicated, and/or unmanageable) unrelenting cardiac compromise, with or without hemodynamic instability, and not responsive to any form of therapy except cardiac surgery. An emergency operation is one in which there should be no delay in providing operative intervention.		112000001278	ACC NCDR
Salvage Procedure			112000001279	ACC NCDR





Parent: Lab Visit



Section: Procedure Information

Element: 13499 Heart Team Reason for Procedure

Coding Instruction: Indicate the heart team's reason for the transcatheter valve replacement procedure.

Note: If the heart team did not document a risk category, consider patients with a predicted risk of 30-day mortality based on the risk model developed by the Society of Thoracic Surgeons as

noted below:

Low risk is considered <3% Intermediate risk is considered 3-7%. High risk is considered >=8%.

Extreme risk includes technically inoperable, co-morbid and debilitated patients.

Target Value: The value on current procedure

Technical Specification

Code: 112000001281
Code System: ACC NCDR
Short Name: OperatorReason

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Transcatheter Valve Therapy Procedure Risk Types - 1.3.6.1.4.1.19376.1.4.1.6.5.505

Selection	Definition	Source	Code	Code System
Extreme Risk			112000001282	ACC NCDR
High Risk			112000001283	ACC NCDR
Intermediate Risk			112000001284	ACC NCDR
Low Risk			112000001285	ACC NCDR

Element: 13504 Heart Team Evaluation of Suitability for Surgical Replacement

Coding Instruction: Indicate if, as part of the Heart Team patient assessment, both an Interventional Cardiologist

AND a Cardiothoracic Surgeon evaluated the patient face to face for the suitability for open

heart valve replacement surgery and documented the evaluation in the medical record.

Target Value: The value on current procedure

Technical Specification

 Code:
 112000001291

 Code System:
 ACC NCDR

 Short Name:
 EvalAVRSuit

 Missing Data:
 Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal



Element: 12871



Full Specifications Data Dictionary v3.0



Section: Procedure Information Parent: Lab Visit

Procedure Location

Coding Instruction: Indicate the location where the procedure was performed.

Target Value: The value on current procedure

Supporting Definition: Procedure Location

The area of the healthcare facility where the procedure was performed.

Technical Specification Code: 112000000623

Code System: ACC NCDR Short Name: ProcedureLocation

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes Is Followup Element: No Data Type: CD

Is Identifier: No

Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Procedure Location - 1.3.6.1.4.1.19376.1.4.1.6.5.327

Selection	Definition	Source	Code	Code System
Cardiac Catheterization Laboratory	on		112000000616	ACC NCDR
Hybrid Catheterization Laboratory Suite	1		112000001266	ACC NCDR
Hybrid Operating Room	m Suite		112000001265	ACC NCDR
Other			100000351	ACC NCDR



Element: 13331



Full Specifications Data Dictionary v3.0



Section: Procedure Information Parent: Lab Visit

Anesthesia Type Coding Instruction: Indicate the type of anesthesia used for the procedure.

Target Value: The highest value on current procedure

Supporting Definition: Anesthesia

Anesthesia is defined as the loss of sensation resulting from pharmacologic depression of nerve function. There are several types of anesthesia including neuraxial, general, or peripheral nerve block. Monitored Anesthesia Care is a specific type of anesthesia service that may be provided when neuraxial anesthesia, general anesthesia, or peripheral nerve block is not utilized.

Source: Anesthesia Quality Institute (2018). 2018 AQI NACOR data element conceptual

definition. Retrieved from

http://www.aqihq.org/files/AQI_NACOR_DATA_ELEMENT_DEFINITIONS_v3%

202018_FINAL.pdf

Technical Specification

Code: 399248000 Code System: SNOMED CT Short Name: AnesthesiaType

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Valid Range: Data Source: User

Anesthesia Type - 1.3.6.1.4.1.19376.1.4.1.6.5.463

Selection	Definition	Source	Code	Code System
General Anesthesia	General Anesthesia is a drug-induced loss of consciousness during which patients are not arousable, even by painful stimulation. The ability to independently maintain ventilatory function is often impaired. Patients often require assistance in maintaining a patent airway, and positive pressure ventilation may be required because of depressed spontaneous ventilation or drug-induced depression of neuromuscular function. Cardiovascular function may be impaired.	Excerpted from Continuum of Depth of Sedation: Definition of General Anesthesia and Levels of Sedation/Analgesia approved on October 13, 1999 and last amended October 15, 2014 of the American Society of Anesthesiologists. A copy of the full text can be obtained from ASA, 1061 American Lane Schaumburg, IL 60173-4973 or online at www.asahq.org.	420653000	SNOMED CT
Deep sedation/Analgesia	Deep Sedation/Analgesia is a drug-induced depression of consciousness during which patients cannot be easily aroused but respond purposefully following repeated or painful stimulation. The ability to independently maintain ventilatory function may be impaired. Patients may require assistance in maintaining a patent airway, and spontaneous ventilation may be inadequate. Cardiovascular function is usually maintained.		426155000	SNOMED CT
Moderate Sedation/Analgesia (Conscious Sedation)	Moderate Sedation/Analgesia ("Conscious Sedation") is a drug-induced depression of consciousness during which patients respond purposefully to verbal commands, either alone or accompanied by light tactile stimulation. No interventions are required to maintain a patent airway, and spontaneous ventilation is adequate. Cardiovascular function is usually maintained. Reflex withdrawal from a painful stimulus is NOT considered a purposeful response.		314271007	SNOMED CT
Minimal Sedation/Anxiolysis	Minimal Sedation (Anxiolysis) is a drug-induced state during which patients respond normally to verbal commands. Although cognitive function and physical coordination may be impaired, airway reflexes, and ventilatory and cardiovascular functions are unaffected.		427255001	SNOMED CT







Section: Procedure Information

Parent: Lab Visit

Element: 13505 Procedure Aborted

Coding Instruction: Indicate whether the procedure was cancelled or aborted after the patient entered the procedure room. A procedure is aborted when the procedure is terminated before device

deployment is attempted. Once device deployment is attempted, the procedure is considered

failed. In this scenario, code device successfully deployed=no.

For mitral leaflet clip procedures, a procedure is considered aborted when the steerable guide

cath was never introduced into the patient.

Target Value: The value on current procedure

Technical Specification

Code: 112000000515
Code System: ACC NCDR
Short Name: TVTProcedureAbort

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup
Element:

Data Type: BL
Precision:

Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User







Section: Procedure Information Parent: Lab Visit

Element: 13506 Reason for Aborting Procedure

Coding Instruction: Indicate the reason why the procedure was canceled or aborted.

Target Value: The value on current procedure

Technical Specification

Code: 112000001292

Code System: ACC NCDR

Short Name: ProcedureAbortReason

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13505 Procedure Aborted

Operator: Equal Value: Yes

Transcatheter Valve Therapy Procedure Aborted Reasons - 1.3.6.1.4.1.19376.1.4.1.6.5.554

Selection	Definition	Source	Code	Code System
Access Related	The procedure was aborted because of difficulties at the procedure access site.		112000001460	ACC NCDR
Navigation Issue After Successful Access	The procedure was aborted because of navigation issues after successful access. Examples include inability to advance through ilio-femoral system due to vessel size/tortuousity/calcification/disease; inability to navigate aorta; and inability to cross aortic valve.		112000001461	ACC NCDR
New Clinical Findings	The procedure was aborted because of new clinical findings (that are not access or navigation related issues). Examples include (but are not limited to) an annulus too large or small, thrombus or vegetation on valve, valve not felt to be severely stenosed or diseased as suspected.		112000001462	ACC NCDR
Device or Delivery System Malfunction	The procedure was aborted because of a malfunction of either the device or delivery system prior to when the operator attempted to deploy the device.	1	112000001463	ACC NCDR
Patient Clinical Status	The procedure was aborted because of a change in the patient's clinical status. Examples include (but are not limited to) a patient becoming hemodynamically unstable during the procedure, a patient having an adverse medication or other reaction, or a patient experiencing another complication prior to completion of the procedure.		112000001464	ACC NCDR
Consent Issue	The procedure was aborted because the patient/famil or physician changed their decision to perform the procedure after the start of the case.	ly	112000001465	ACC NCDR
Transseptal Access Related	The procedure was aborted because of difficulties crossing the septum.		112000001466	ACC NCDR
System Issue	The procedure was aborted because of equipment (not device) malfunction (such as x-ray system equipment malfunction), or a situation where an emergency surgical case causes the transcatheter case to be aborted and rescheduled after the patient was in the room but prior to starting the case.		112000001467	ACC NCDR
Other			100000351	ACC NCDR







Section: Procedure Information

Parent: Lab Visit

Element: 13757 Procedure Aborted Action

Coding Instruction: Indicate the reason or action taken as a result of the aborted TVT procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001468 Code System: ACC NCDR

Short Name: ProcedureAbortAction

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Is Identifier: No

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13505 Procedure Aborted

Operator: Equal Value: Yes

Transcatheter Valve Therapy Procedure Aborted Action - 1.3.6.1.4.1.19376.1.4.1.6.5.555

Selection	Definition	Source	Code	Code System
Conversion to Open	Heart		112000001327	ACC NCDR
Surgery				
Scheduled Open He	art		112000001473	ACC NCDR
Surgery				
Rescheduled Transo	catheter		112000001470	ACC NCDR
Procedure				
Converted to Clinical	Trial		112000001472	ACC NCDR
Balloon Valvuloplast	у		112000001469	ACC NCDR
Converted to Medica	l Therapy		112000001471	ACC NCDR
Other			100000351	ACC NCDR

Element: 13542	Conversion to Open Heart Surgery	Technical Specification
		Code: 112000001327
Coding Instruction:	Indicate if conversion to open heart surgical access was required.	Code System: ACC NCDR

Target Value: The value on current procedure

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

Short Name: ConvSurgAccess

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Procedure Information

Parent: Lab Visit

Element: 13543 Reason for Conversion to Open Heart Surgery

Coding Instruction: Indicate the reason for conversion to open heart surgical access.

Target Value: The value on current procedure

Technical Specification

Code: 112000001327 Code System: ACC NCDR

Short Name: ConvSurgAccessReason

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
No
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Multiple

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13542 Conversion to Open Heart

Surgery
Operator: Equal
Value: Yes

Reason for Conversion to Open Heart Surgery - 1.3.6.1.4.1.19376.1.4.1.6.5.513

Selection	Definition	Source	Code	Code System
Valve Dislodged to Aorta	1		112000001328	ACC NCDR
Valve Dislodged to Left			112000001329	ACC NCDR
Ventricle				
Annulus Rupture			112000001331	ACC NCDR
Ventricular Rupture			112000001330	ACC NCDR
Aortic Dissection			308546005	SNOMED CT
Coronary Occlusion			63739005	SNOMED CT
Access Related			112000001460	ACC NCDR
Cardiac Tamponade			35304003	SNOMED CT
Inability to Position Device	e		112000001479	ACC NCDR
Device Embolization			112000001324	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR

Element: 7422 Mechanical Ventricular Support

Coding Instruction: Indicate if the patient required mechanical ventricular support.

Target Value: Any occurrence on current procedure

Technical Specification

Code: 100014009
Code System: ACC NCDR
Short Name: MechVentSupp

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Procedure Information

Parent: Lab Visit

Element: 7423 Mechanical Ventricular Support Device

Coding Instruction: Indicate the mechanical ventricular support device used.

Note(s):

The device that should be collected in your application are controlled by a Mechanical Ventricular Support Master file. This file is maintained by the NCDR and will be made available

on the internet for downloading and importing/updating into your application.

If more than one device is used, code the device with the highest level of support.

Target Value: Any occurrence on current procedure

Technical Specification

Code: 100001278
Code System: ACC NCDR
Short Name: MVSupportDevice

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 7422 Mechanical Ventricular Support

Operator: Equal Value: Yes

Mechanical Ventricular Support Device - 2.16.840.1.113883.3.3478.6.1.24

Selection	Definition	Source	Code	Code System
Cardiopulmonary Support (CPS)			1000142428	ACC NCDR
Extracorporeal membrane oxygenation (ECMO)			233573008	SNOMED CT
Impella: Left Ventricular Support			100014011	ACC NCDR
Impella: Right Ventricular Support			112000000188	ACC NCDR
Intra-aortic balloon pump (IABP)			442807006	SNOMED CT
Isolated Right Ventricular Support			112000000546	ACC NCDR
Left ventricular assist device (LVAD)			232967006	SNOMED CT
Right Ventricular Assist Device (RVAD)			360065002	SNOMED CT
Percutaneous Heart Pump (PHP)			1000142429	ACC NCDR
TandemHeart			100014010	ACC NCDR
Other			100000351	ACC NCDR







Section: Procedure Information

Parent: Lab Visit

Element: 7424 Mechanical Ventricular Support Timing

Coding Instruction: Indicate when the mechanical ventricular support device was placed.

Target Value: Any occurrence on current procedure

Technical Specification

Code: 100014009
Code System: ACC NCDR
Short Name: MVSupportTiming

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:
Data Source: User

Parent/Child Validation

Element: 7422 Mechanical Ventricular Support

Operator: Equal Value: Yes

Mechanical Ventricular Support Timing - 1.3.6.1.4.1.19376.1.4.1.6.5.524

Selection	Definition	Source	Code	Code System
In place at start of proced	ure		100001280	ACC NCDR
Inserted during procedure prior to intervention	and		100001281	ACC NCDR
Inserted after intervention begun	has		100013042	ACC NCDR
Post Procedure			112000001347	ACC NCDR

Element: 13579 Cardiopulmonary Bypass Used Technical Code: 636

Coding Instruction: Indicate if cardiopulmonary bypass or coronary perfusion was used during the procedure.

Target Value: Any occurrence on current procedure

Technical Specification

Code: 63697000
Code System: SNOMED CT
Short Name: CPB
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Usual Range: Valid Range: Data Source: User

Default Value: Null







Section: Procedure Information

Parent: Lab Visit

Element: 13580 Cardiopulmonary Bypass Status

Coding Instruction: Indicate if the use of cardiopulmonary bypass was elective or emergent.

Target Value: The value on current procedure

Technical Specification

Code: 63697000
Code System: SNOMED CT
Short Name: CPBStatus
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13579 Cardiopulmonary Bypass Used

Operator: Equal Value: Yes

Cardiopulmonary Procedure Status - 1.3.6.1.4.1.19376.1.4.1.6.5.766

 Selection
 Definition
 Source
 Code
 Code System

 Elective Procedure
 71388002:260870009=103390000
 SNOMED CT

 Emergency Procedure
 112000001278
 ACC NCDR

Element: 13581 Cardiopulmonary Bypass Time

Coding Instruction: Indicate the total number of minutes that systemic return is diverted into the cardiopulmonary

bypass (CPB) circuit and returned to the systemic system. This time period (Cardiopulmonary Bypass Time) includes all periods of cerebral perfusion and sucker bypass. This time period (Cardiopulmonary Bypass Time) excludes any circulatory arrest and modified ultrafiltration periods. If more than one period of CPB is required during the procedure, the sum of all the CPB

periods will equal the total number of CPB minutes.

Target Value: The total between start of procedure and end of procedure

Technical Specification Code: 364669000

Code System: SNOMED CT Short Name: PerfusTm Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: min
Default Value: Null
Usual Range: 1 - 300 min
Valid Range: 1 - 999 min
Data Source: User

Parent/Child Validation

Element: 13579 Cardiopulmonary Bypass Used

Operator: Equal Value: Yes





Parent: Lab Visit



Section: Procedure Information

Element: 13525 Delivery System Successfully Removed

Coding Instruction: Indicate if the delivery system was successful removed.

Target Value: The value on current procedure

Technical Specification

Code: 112000001312
Code System: ACC NCDR
Short Name: DeliveryRemoved

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13644 Positive Inotropes Administered

Coding Instruction: Indicate if positive inotropes was administered.

For patients requiring IV inotropic support, indicate positive inotropes only.

Target Value: Any occurrence between start of procedure and end of procedure

Technical Specification
Code: 112000001358

Code System: ACC NCDR
Short Name: ProcInotropesAdmin

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Procedure Medications Administered - 1.3.6.1.4.1.19376.1.4.1.6.5.415

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR







Section: Operator Information Parent: Procedure Information

Element: 14476 TVT Operator First Name

Coding Instruction: Indicate the first name of operator.

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

Target Value: The value on current procedure

Vendor Instruction: A TVT Operator - combination First Name (14476), Last Name (14478) and NPI (14479) - may

only be entered/selected once

Technical Specification

Code: 112000001955 Code System: ACC NCDR

Short Name: TVT_Oper_FirstName

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup

Element: NO
Data Type: FN
Precision: 50
Selection Type: Single
Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Element: 14478 TVT Operator Last Name

Coding Instruction: Indicate the last name of operator.

Note(s):

If the name exceeds 50 characters, enter the first 50 characters only.

Target Value: The value on current procedure

Technical Specification

Code: 112000001955 Code System: ACC NCDR

Short Name: TVT_Oper_LastName

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: LN

Precision: 50
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Element: 14477 TVT Operator Middle Name

Coding Instruction: Indicate the middle name of operator.

Note(s):

It is acceptable to specify the middle initial.

If there is no middle name given, leave field blank.

If there are multiple middle names, enter all of the middle names sequentially.

If the name exceeds 50 characters, enter the first 50 letters only.

Target Value: The value on current procedure

Technical Specification

Code: 112000001955
Code System: ACC NCDR

Short Name: TVT_Oper_MidName

Missing Data: Report

Data Source: User

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: MN
Precision: 50
Selection Type: Single
Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Operator Information

Parent: Procedure Information

Element: 14479 TVT Operator NPI

Coding Instruction: Indicate the National Provider Identifier (NPI) of the operator who is performing the procedure.

NPI's, assigned by the Centers for Medicare and Medicaid Services (CMS), are used to

uniquely identify physicians for Medicare billing purposes.

Target Value: The value on current procedure

Code: 112000001955 Code System: ACC NCDR

Technical Specification

Short Name: TVT_Oper_NPI
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: NUM
Precision: 10
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User







Sect	ion:	Radia	tion and	Con	trast

Parent: Procedure Information

Technical Specification Element: 14278 Dose Area Product Code: 100000994 Coding Instruction: Indicate the total fluoroscopy dose to the nearest integer. The value recorded should include Code System: ACC NCDR the total dose for the lab visit. Short Name: FluoroDoseDAP2 Target Value: The total between start of current procedure and end of current procedure Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, Supporting Definition: Dose Area Product TTVP) Dose Area Product is the integral of air kerma (the energy extracted from an x-ray beam per Is Identifier: No unit mass of air in a small irradiated air volume; for diagnostic x-rays, the dose delivered to that Is Base Element: Yes volume of air) across the entire x-ray beam emitted from the x-ray tube. It is a surrogate Is Followup No measure of the amount of energy delivered to the patient. Element: Data Type: PQ Also known as KAP (Kerma Area Product). Precision: 7,0 Source: Miller DL, et al. Radiation doses in interventional radiology procedures. (J Vasc Selection Type: Single Interv Radiol 2003; 14:711-727.) Unit of Measure: Gy-cm², dGy-cm², cGy-cm², mGy⋅cm², µGy⋅M² Default Value: Null Usual Range: 1 - 700 Gy·cm² 10 - 7,000 dGy·cm² 100 - 70,000 cGy·cm² 100 - 70,000 μGy·M² 1,000 - 700,000 mGy·cm² Valid Range: 1 - 5,000 Gy·cm² 10 - 50,000 dGy·cm² 100 - 500,000 cGy·cm² 100 - 500,000 μGy·M² 1,000 - 5,000,000 mGy·cm²

Technical Specification Element: 7210 Cumulative Air Kerma Code: 228850003 Coding Instruction: Indicate the total radiation dose (Cumulative Air Kerma, or Reference Air Kerma) recorded to Code System: SNOMED CT the nearest milligray (mGy) or gray (Gy). The value recorded should include the total dose for Short Name: FluoroDoseKerm the lab visit. Cumulative air kerma is the total air kerma accrued from the beginning of an Missing Data: Report examination or procedure and includes all contributions from fluoroscopic and radiographic Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP) Target Value: The total between start of current procedure and end of current procedure Is Identifier: No Supporting Definition: Cumulative (Reference) Air kerma Cumulative air kerma (also known as reference, reference dose, cumulative dose, or Element: cumulative dose at a reference point) is the air kerma accumulated at a specific point in space (the patient entrance reference point) relative to the gantry of the fluoroscopy system. The quantity, kerma, originated from the acronym, KERMA, for Kinetic Energy Released per unit

Source: Miller DL, et al. Radiation doses in interventional radiology procedures. (J Vasc

Interv Radiol 2003:14:711-727.)

Is Base Element: Yes Is Followup No Data Type: PQ Precision: 5,0 Selection Type: Single Unit of Measure: mGy, Gy Default Value: Null Usual Range: 1 - 10 Gy 1 - 10,000 mGy

Data Source: User

Valid Range: 1 - 50 Gy 1 - 50,000 mGy

Data Source: User







Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mL
Default Value: Null
Usual Range: 5 - 300 mL
Valid Range: 0 - 999 mL
Data Source: User

Section: Radiation and	Contrast Parent: Procedure In	formation	
Element: 7214	Fluoroscopy Time	Technic	al Specification
Coding Instruction:	Indicate the total fluoroscopy time recorded to the nearest 0.1-minute. The time recorde should include the total time for the lab visit.		
Target Value:	The total between start of current procedure and end of current procedure	Missing Data: Harvested:	
		Is Identifier: Is Base Element: Is Followup	No Yes
		Element: Data Type: Precision:	PQ
		Selection Type: Unit of Measure: Default Value:	min
		Usual Range: Valid Range:	0.1 - 30.0 min 0.1 - 300.0 min
		Data Source:	
lement: 7215	Contrast Volume		80242-1
Coding Instruction:	Indicate the volume of contrast (ionic and non-ionic) used in milliliters (ml). The volume reshould be the total volume for the lab visit.	code: Code System: Short Name:	LOINC
Target Value:	The total between start of current procedure and end of current procedure	Missing Data: Harvested:	Report Yes (TAVR, TMVR, TMVrpr







Section: Post Implant Mitral Valve Data

Parent: Procedure Information

Element: 14274 Mitral Regurgitation

Coding Instruction: Indicate the severity of regurgitation through the mitral valve.

Note(s):

Code the highest value or most severe regurgitation when a range is reported.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 48724000
Code System: SNOMED CT
Short Name: Intraproc_Post_MR

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None	,		112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR

Flement: 13762	Mitral Valve Mean Gradient

 $\textbf{Coding Instruction:} \quad \text{Indicate the mean gradient (in mm Hg) across the mitral valve}.$

Target Value: The last value between the implant and the end of current procedure

Supporting Definition: Mitral Valve Mean Gradient

The average gradient across the mitral valve occurring during the entire systole.

Source: Baumgartner, H. et. al (2009). Echocardiographic assessment of valve stenosis:

EAE/ASE

recommendations for clinical practice.

Technical Specification

Code: 112000001191
Code System: ACC NCDR

Short Name: MVR_Post_MeanMVGrad

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ

Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 5 - 50 mm[Hg] **Valid Range:** 0 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: TAVR	Parent: Procedure Information
Section: TAVK	Parcente Proceedure Intormation

Element: 13498 Primary Transcatheter Aortic Valve Replacement Procedure Indication

Coding Instruction: Indicate the primary indication for the transcatheter aortic valve replacement. If more than one

indication is present, choose the most significant.

Target Value: The highest value between 2 months prior to current procedure and current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (TAVR) then Primary Transcatheter Aortic Valve Replacement Procedure Indication (13498) cannot be Null

Technical Specification
Code: 112000000482
Code System: ACC NCDR

Short Name: PrimTAVRProcInd Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Transcatheter Aortic Valve Replacement Primary Procedure Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.738

Selection	Definition	Source	Code	Code System
Aortic Regurgitation			60234000	SNOMED CT
Aortic Stenosis			60573004	SNOMED CT

Element: 13500	Valve In Valve Procedure
	Indicate whether a "valve-in-valve" procedure was performed on previously implanted bioprosthetic valve.
	Code no if the procedure is being performed in a native aortic valve. Code yes if the procedure is being performed in a previously implanted bioprosthetic valve.
Target Value:	The value on current procedure
Supporting Definition:	Valve in Valve Procedure
	A procedure where a prosthetic valve is placed in a prosthetic valve that was previously implanted.
	Source:
Vendor Instruction:	Valve In Valve Procedure (13500) must be (Yes) when Procedure History Name (12905) is (AV Replacement Surgery OR AV Replacement -Transcatheter) and Procedure History Occurrence (14268) is (Yes)
	When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (TAVR) then Valve In Valve Procedure (13500) cannot be Null

Technical Specification

Code: 112000001286
Code System: ACC NCDR
Short Name: ValveInValve
Missing Data: Report
Harvested: Yes (BDS, TAVR)

Harvested: Yes (BDS, TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: TAVR Parent: Procedure Information

Element: 13501 Bioprosthetic Valve Fracture Attempted

Coding Instruction: Indicate if bioprosthetic valve fracture (BVF) with high pressure balloon dilation was attempted

on the previously implanted bioprosthetic valve.

Note 1: If pre-implant valvuloplasty or post-implant post dilatation with lower pressure

inflations (e.g. a hand inflation up to 4 atm), code no.

Note 2: If the previously implanted bioprosthetic valve was fractured during the procedure

(even though BVF was not planned), code yes.

Target Value: The value on current procedure

Supporting Definition: Bioprosthetic Valve Fracture

Bioprosthetic Valve Fracture (BVF) is a technique that uses a high pressure dilatation with intent to purposefully fracture or crack the ring of the previously implanted bioprosthetic valve and allow the new implanted valve to more fully expand. This technique requires balloon

pressures of up to 20 atm.

Source: STS/ACC TVT Registry

Technical Specification

Code: 112000001287 Code System: ACC NCDR

Short Name: BVFAttempt Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes

Is Followup Element: No Data Type: BL Precision:

Selection Type: Single Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Flement: 13500 Valve In Valve Procedure

Operator: Equal Value: Yes

Element: 13502 Bioprosthetic Valve Fracture Timing

Coding Instruction: Indicate the timing of the bioprosthetic valve fracture.

Note: If BVF was attempted both pre and post valve implant, code both.

Target Value: The value on current procedure

Technical Specification

Code: 112000001287
Code System: ACC NCDR
Short Name: BVFTiming
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13501 Bioprosthetic Valve Fracture

Attempted
Operator: Equal
Value: Yes

Timing - 1.3.6.1.4.1.19376.1.4.1.6.5.729

Selection	Definition	Source	Code	Code System
Pre Implant			112000001912	ACC NCDR
Post Implant			112000001913	ACC NCDR



Element: 13503



Full Specifications **Data Dictionary v3.0**



Section: TAVR Parent: Procedure Information

Coding Instruction: Indicate if the valve was observed to be fractured. Documentation can include any of the

following:

(1) Fluoroscopically by either visualizing the waist of the balloon release and/or the fractured

valve ring (if the valve ring is radiopaque);

Valve Observed to be Fractured

(2) By an audible snap, or

(3) By a sudden drop in the balloon pressure in the absence of balloon rupture.

Target Value: The value on current procedure

Technical Specification

Code: 112000001290
Code System: ACC NCDR
Short Name: ValveFractured
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 13501 Bioprosthetic Valve Fracture

Attempted

Data Source: User

Operator: Equal Value: Yes

Element: 13507 Valve Sheath Access Site

Coding Instruction: Indicate the access site for the valve sheath.

Target Value: The value on current procedure

Technical Specification

Code: 112000001293
Code System: ACC NCDR
Short Name: TVTAccessSite
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal

Value: TAVR

Valve Sheath Access Site - 1.3.6.1.4.1.19376.1.4.1.6.5.506

Selection	Definition	Source	Code	Code System
Axillary Artery			67937003	SNOMED CT
Carotid			32062004	SNOMED CT
Direct Aortic			112000001957	ACC NCDR
Femoral Artery			7657000	SNOMED CT
Iliac			112000000893	ACC NCDR
Subclavian Artery			36765005	SNOMED CT
Transapical			112000001295	ACC NCDR
Transcaval			112000001299	ACC NCDR
Transseptal via Femoral	l Vein		112000001296	ACC NCDR
Other			100000351	ACC NCDR







Section: TAVR Parent: Procedure Information

Element: 13508 Valve Sheath Access Site Method

Coding Instruction: Indicate the access method used to deliver the valve sheath.

Target Value: The value on current procedure

Technical Specification

Code: 112000001300
Code System: ACC NCDR
Short Name: TVTAccessMethod

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Valve Sheath Access Site Method - 1.3.6.1.4.1.19376.1.4.1.6.5.507

Selection	Definition	Source	Code	Code System
Percutaneous Approac	ch		103388001	SNOMED CT
Cutdown			112000001301	ACC NCDR
Mini Sternotomy			112000001303	ACC NCDR
Mini Thoracotomy			112000001302	ACC NCDR
Other			100000351	ACC NCDR

Element: 13509 Valve Sheath Delivery Size

Coding Instruction: Indicate the size, in french, of the valve sheath delivery system.

Target Value: The value on current procedure

Technical Specification

Code: 112000001304

Code System: ACC NCDR

Short Name: ValveSheathDelivery

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: Fr
Default Value: Null
Usual Range: 14 - 32 Fr
Valid Range: 5 - 40 Fr
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: TAVR Parent: Procedure Information

Element: 13510 Embolic Protection Deployed

Coding Instruction: Indicate if embolic protection was used during the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001305
Code System: ACC NCDR
Short Name: EmbProt
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13511 Embolic Protection Device

Coding Instruction: Indicate the embolic protection device used during the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001306
Code System: ACC NCDR
Short Name: EmbProtDevice
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Embolic Protection Deployed

Element: 13510 Operator: Equal Value: Yes







Section: TAVR Parent: Procedure Information

Element: 14304 Aortic Valve Regurgitation

Coding Instruction: Indicate the severity of aortic valve regurgitation.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 60234000
Code System: SNOMED CT
Short Name: AVR_Post_AR

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TAVR

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR

Element: 14303 Aortic Valve Mean Gradient

Coding Instruction: Indicate the highest documented mean gradient (in mm Hg) across the aortic valve.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 112000001398 Code System: ACC NCDR

Short Name: PostImplant_AVMeanGrad

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Element: NO
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 5 - 50 mm[Hg]

Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: TAVR Devices Parent: TAVR

Element: 13524 Transcatheter Aortic Valve Replacement Device Counter

Coding Instruction: This is a software-assigned value. The counter will start at one and be incremented by one for

each device or system used.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.851

Code System: ACC NCDR
Short Name: TAVRDevCounter

Missing Data: Illegal

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CTR
Precision: 3
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range: 1 - 999
Data Source: Automatic

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 13505 Procedure Aborted

Operator: Equal Value: No

Element: 14485 Transcatheter Aortic Valve Replacement Device ID

Coding Instruction: Indicate the device ID of the aortic valve.

Target Value: The value on current procedure

Technical Specification

Code: 112000001805
Code System: ACC NCDR
Short Name: TAVRDeviceID

Missing Data: Illegal

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13524 Transcatheter Aortic Valve Replacement Device Counter

Operator:

Value: Any Value







Section: TAVR Devices Parent: TAVR

Element: 14532 Transcatheter Aortic Valve Replacement Device Diameter

Coding Instruction: Indicate the transcatheter aortic valve replacement device diameter (in mm).

Target Value: The value on current procedure

Technical Specification

Code: 112000001805
Code System: ACC NCDR
Short Name: TAVRDeviceDia

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm
Valid Range: 5 - 100 mm

Data Source: User

Parent/Child Validation

Element: 14485 Transcatheter Aortic Valve

Replacement Device ID

Operator:

Value: Any Value

Element: 13534 Device Capture and Repositioning Performed

Coding Instruction: Indicate if device capture and repositioning was performing during the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001318 Code System: ACC NCDR

Short Name: TVTDeviceRepositioning

Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13535 Device Capture and Repositioning

Performed Not Applicable

Operator: Equal

Value: No (or Not Answered)
----- AND

Element: 13524 Transcatheter Aortic Valve

Replacement Device Counter

Operator:

Value: Any Value

Boolean w/Unknown - 1.3.6.1.4.1.19376.1.4.1.6.5.444

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR







Section: TAVR Devices Parent: TAVR

Element: 13535 Device Capture and Repositioning Performed Not Applicable

Coding Instruction: Indicate if performing a device capture and repositioning was not applicable.

Target Value: N/A

Technical Specification

Code: 63653004 Code System: SNOMED CT

Short Name: TVTDeviceRepositioningNA

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range:

Valid Range: Data Source: User

Parent/Child Validation
Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Replacement Device Counter

Operator:

Value: Any Value

Element: 13536 Transcatheter Aortic Valve Replacement Device Implanted Successfully

Coding Instruction: Indicate if there is correct positioning of a single prosthetic heart valve in the proper anatomical

ocation

Target Value: The value on current procedure

Technical Specification

Code: 112000001805 Code System: ACC NCDR

Short Name: TAVRDeviceImplantSuccessful

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13524 Transcatheter Aortic Valve

Replacement Device Counter

Operator:

Value: Any Value







Section: TAVR Devices Parent: TAVR

Element: 13539 Reason Transcatheter Aortic Valve Replacement Device Not Implanted

Successfully

Coding Instruction: Indicate the reason the device was not implanted successfully.

Target Value: The value on current procedure

Technical Specification

Code: 112000002014 Code System: ACC NCDR

Short Name: TAVR_Unsuccessful

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 13536 Transcatheter Aortic Valve

Replacement Device Implanted

Successfully
Operator: Equal

Value: No

Transcatheter Valve Therapy Reason Device Not Implanted Successfully - 1.3.6.1.4.1.19376.1.4.1.6.5.512

Selection	Definition	Source	Code	Code System
Device Embolization			112000001324	ACC NCDR
Improper Device Positionin	ng		112000001325	ACC NCDR
Improper Device Sizing			112000001326	ACC NCDR
Other			100000351	ACC NCDR

Element: 14286 Transcatheter Aortic Valve Replacement Device Serial Number

Coding Instruction: Indicate the device transcatheter aortic valve replacement device serial number.

Target Value: The value on current procedure

Technical Specification

Code: 112000001805
Code System: ACC NCDR
Short Name: TAVRDeviceSN

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: ST
Precision: 30
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13536 Transcatheter Aortic Valve

Replacement Device Implanted

Successfully

Operator: Equal Value: Yes







Section: TAVR Devices Parent: TAVR

Element: 14572 Transcatheter Aortic Valve Unique Device ID

Coding Instruction: Indicate the full unique device identifier (UDI) for the implanted device.

Target Value: The value on current procedure

Supporting Definition: Unique Device Identifier (UDI)

An identifier that is the main (primary) lookup for a medical device product and meets the requirements to uniquely identify a device through its distribution and use. This value is

supplied to the FDA by the manufacturer.

Source: US FDA

Technical Specification

Code: 2.16.840.1.113883.3.3719

Code System: ACC NCDR Short Name: TAV_UDI Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: ST
Precision: 150
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13536 Transcatheter Aortic Valve

Replacement Device Implanted

Successfully

Operator: Equal Value: Yes







Section: TMVr Parent: Procedure Information

Element: 13792 Mitral Leaflet Clip Procedure Indication

Coding Instruction: Indicate the indication(s) for the mitral leaflet clip procedure.

Target Value: The last value on current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (TMVr) then Mitral

Leaflet Clip Procedure Indication (13792) cannot be Null

Technical Specification

Code: 112000000482
Code System: ACC NCDR
Short Name: MRRIndication
Missing Data: Report

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: Data Type: CD
Precision:
Selection Type: Multiple
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Mitral Leaflet Clip Procedure Indications - 1.3.6.1.4.1.19376.1.4.1.6.5.558

(Cirrhosis or MELD score >12) End-Stage Porcelain Aorta Predicted STS MV Repair ROM Predicted		1120000019	944 ACC NCDR
Frailty Hostile Chest Severe Pulmonary Hypertension Severe Liver Disease The patier (Cirrhosis or MELD score >12) End-Stage Porcelain Aorta Predicted STS MV Repair ROM Predicted			ACC NODIC
Severe Pulmonary Hypertension Severe Liver Disease The patier (Cirrhosis or MELD score >12) End-Stage Porcelain Aorta Predicted STS MV Repair ROM Predicted		248279	007 SNOMED CT
Hypertension Severe Liver Disease The patier (Cirrhosis or MELD score >12) End-Stage Porcelain Aorta Predicted STS MV Repair ROM Predicted		112000001	489 ACC NCDR
(Cirrhosis or MELD score >12) End-Stage Porcelain Aorta Predicted STS MV Repair ROM Predicted		112000001	490 ACC NCDR
Predicted STS MV Repair ROM Predicted	at has a history of cirrhosis or a "Model For Liver Disease" (MELD) score >12 points.	112000001	482 ACC NCDR
•		112000001	175 ACC NCDR
Percent mitral valv	STS Mitral Valve Repair Operative Mortality 6% for a patient deemed likely to undergo e repair surgery.	112000001	483 ACC NCDR
Replacement ROM Greater Mortality R	STS Mitral Valve Replacement Operative isk is >=8% for a patient deemed likely to itral valve replacement surgery.	112000001-	484 ACC NCDR
RVD with Severe TR Right Venture Regurgitat	tricular Dysfunction with Severe Tricuspid ion.	112000001	486 ACC NCDR
Major Bleeding Diathesis		112000001	487 ACC NCDR
Chemotherapy for Malignancy		112000001	491 ACC NCDR
AIDS Acquired I	mmune Deficiency Syndrome	624790	008 SNOMED CT
Immobility		112000001	492 ACC NCDR
High Risk of Aspiration		112000001	488 ACC NCDR
Severe Dementia		112000001	914 ACC NCDR
IMA at High Risk of Injury Internal Ma	ammary Artery at High Risk of Injury.	112000001	494 ACC NCDR
Other			







Section: TMVr Parent: Procedure Information

Element: 13794 Guiding Catheter Access Site

Coding Instruction: Indicate the leaflet clip guiding catheter access site.

Target Value: The value on current procedure

Technical Specification

Code: 112000001495
Code System: ACC NCDR
Short Name: LeafAccess
Missing Data: Report

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Guiding Catheter Access Site - 1.3.6.1.4.1.19376.1.4.1.6.5.560

Selection	Definition	Source	Code	Code System
Right Femoral Vein			767174009	SNOMED CT
Left Femoral Vein			767173003	SNOMED CT
Jugular Vein			63190004	SNOMED CT
Other Vein			100000351	ACC NCDR

Element: 13795 Steerable Guide Cath Device ID

Coding Instruction: Indicate the steerable guide cath device ID utilized.

Target Value: The value on current procedure

Technical Specification

Code: 112000001496
Code System: ACC NCDR
Short Name: SGCDeviceID
Missing Data: Report
Harvested: Yes (TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr







Section: TMVr Parent: Procedure Information

Element: 13796 Steerable Guide Catheter Serial Number

Coding Instruction: Indicate the manufacturer serial number for the steerable guide used during the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001496
Code System: ACC NCDR
Short Name: MRR_GuideSerNo

Missing Data: Report Harvested: Yes (TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: ST
Precision: 30
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13795 Steerable Guide Cath Device ID

Operator:

Value: Any Value







Section: Mitral Leaflet Devices Parent: TMVr

Element: 13533 Mitral Repair Device Counter

Coding Instruction: This is a software-assigned value. The counter will start at one and be incremented by one for

each device or system used.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.851

Code System: ACC NCDR
Short Name: MRepairDevCounter

Missing Data: Illegal

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CTR
Precision: 3
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range: 1 - 999
Data Source: Automatic

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 13505 Procedure Aborted

Operator: Equal Value: No

Element: 13797 Mitral Repair Device ID

Coding Instruction: Indicate all mitral repair device IDs utilized.

Target Value: The value on current procedure

Technical Specification

Code: 112000002005
Code System: ACC NCDR
Short Name: MRepairDeviceID

Missing Data: Illegal

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13533 Mitral Repair Device Counter

Operator:

Value: Any Value







Section: Mitral Leaflet Devices Parent: TMVr

Element: 13798 Mitral Repair Serial Number

Coding Instruction: Indicate the serial number of the mitral repair device.

Target Value: The value on current procedure

Technical Specification

Code: 112000002005
Code System: ACC NCDR
Short Name: MRepairNum
Missing Data: Report

Harvested: Yes (BDS, TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: ST
Precision: 30
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Is Identifier: No

Parent/Child Validation

Element: 13799 Mitral Repair Device Implanted

Successfully

Data Source: User

Operator: Equal Value: Yes

Element: 14574 Mitral Repair Unique Device ID

Coding Instruction: Indicate the full unique device identifier (UDI) for the implanted device.

Target Value: The value on current procedure
Supporting Definition: Unique Device Identifier (UDI)

An identifier that is the main (primary) lookup for a medical device product and meets the requirements to uniquely identify a device through its distribution and use. This value is

supplied to the FDA by the manufacturer.

Source: US FDA

Technical Specification

Code: 2.16.840.1.113883.3.3719

Code System: ACC NCDR Short Name: MRepair_UDI Missing Data: Report

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: ST
Precision: 150
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13799 Mitral Repair Device Implanted

Successfully
Operator: Equal

Value: Yes







Section: Mitral Leaflet Devices Parent: TMVr

Element: 13800 Mitral Valve Repair Location

Coding Instruction: Indicate the location on the mitral valve where the leaflet clip was attached.

Target Value: The value on current procedure

Technical Specification

Code: 112000002050 Code System: ACC NCDR Short Name: MRR_Loc Missing Data: Report

Harvested: Yes (BDS, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13533 Mitral Repair Device Counter

Operator:

Value: Any Value

Mitral Leaflet Clip Procedure Location - 1.3.6.1.4.1.19376.1.4.1.6.5.709

Selection	Definition	Source	Code	Code System
A1/P1	The mitral leaflet clip was attached to the A1P1 position on the anterior and posterior mitral valve leaflets.		112000001847	ACC NCDR
A2/P2	The mitral leaflet clip was attached to the on the anterior and posterior mitral valve.	•	112000001848	ACC NCDR
A3/P3	The mitral leaflet clip was attached to the on the anterior and posterior mitral valves.	•	112000001849	ACC NCDR
Other Location	Mitral leaflet clip was attached to a loca anterior and posterior mitral leaflets that otherwise specified.		112000001850	ACC NCDR

Element: 13799 Mitral Repair Device Implanted Successfully

Coding Instruction: Indicate if the mitral repair device was successfully deployed.

Target Value: The value on current procedure

Technical Specification

Code: 112000002015 Code System: ACC NCDR

Short Name: MRR_LeafletClipDeploy

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TMVrpr)

Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Data Source: User

Parent/Child Validation

Element: 13533 Mitral Repair Device Counter

Operator:

Value: Any Value







Section: Mitral Leaflet Devices

Parent: TMVr

Element: 13801 Reason Mitral Repair Device Not Implanted Successfully

Coding Instruction: Indicate the reason why the mitral repair device was not deployed.

Target Value: The value on current procedure

Technical Specification

Code: 112000002014

Code System: ACC NCDR

Short Name: MRR_LeafletClipReasonNotDeploy

Missing Data: Report Harvested: Yes (TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13799 Mitral Repair Device Implanted

Successfully

Operator: Equal Value: No

Mitral Leaflet Clip Reason Not Deployed - 1.3.6.1.4.1.19376.1.4.1.6.5.561

Selection I	Definition	Source	Code	Code System
Adverse Event			112000001505	ACC NCDR
Device Malfunction			112000001504	ACC NCDR
Inability to Grasp Leaflets			112000001501	ACC NCDR
Inability to Reduce Mitral			112000001502	ACC NCDR
Regurgitation				
Mitral Valve Injury			112000001503	ACC NCDR
Mitral Valve Stenosis			79619009	SNOMED CT
Other			100000351	ACC NCDR

Element: 13802 Mitral Leaflet Clip Deployed then Removed

Coding Instruction: Indicate if the leaflet clip was removed after it was deployed.

Target Value: The value on current procedure

Technical Specification

Code: 112000002005
Code System: ACC NCDR
Short Name: MRR_ClipRemoved

Missing Data: Report
Harvested: Yes (TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13799 Mitral Repair Device Implanted

Successfully

Operator: Equal Value: Yes







Section: TMVR Parent: Procedure Information

Element: 13754 Transcatheter Mitral Valve Replacement Type

Coding Instruction: Indicate the transcatheter mitral valve replacement procedure type.

Target Value: The value on current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (TMVR) then

Transcatheter Mitral Valve Replacement Type (13754) cannot be Null

Transcatheter Mitral Valve Replacement Type (13754) cannot be (Native Valve) When

Procedure History Name (12905) is (Mitral Valve Replacement Surgery) with Procedure History

Occurrence as (Yes)

AND

Mitral Valve Transcatheter Intervention Type (14261) is (Valve in Native Value Procedure OR

Valve in Valve Procedure)

Technical Specification

Code: 112000001458
Code System: ACC NCDR
Short Name: TMVRType
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Transcatheter Mitral Valve Replacement Types - 1.3.6.1.4.1.19376.1.4.1.6.5.739

Selection	Definition	Source	Code	Code System
Native Valve			112000001456	ACC NCDR
Valve-in-Valve			112000001286	ACC NCDR
Valve-in-Ring			112000001938	ACC NCDR

Element: 13755 Mitral Valve Annular Calcification

Coding Instruction: Indicate if there was mitral annular calcification.

Target Value: The value on current procedure

Technical Specification

Code: 251002009
Code System: SNOMED CT
Short Name: MVDAnnular_Native

Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13754 Transcatheter Mitral Valve

Replacement Type

Operator: Equal

Value: Native Valve







Section: TMVR Parent: Procedure Information

Element: 14480 TMVR Bioprosthetic Valve Fracture Attempted

Coding Instruction: Indicate if bioprosthetic valve fracture (BVF) with high pressure balloon dilation was attempted

on the previously implanted bioprosthetic valve.

Note 1: If pre-implant valvuloplasty or post-implant post dilatation with lower pressure

inflations (e.g. a hand inflation up to 4 atm), code no.

Note 2: If the previously implanted bioprosthetic valve was fractured during the procedure

(even though BVF was not planned), code yes.

Target Value: The value on current procedure

Supporting Definition: Bioprosthetic Valve Fracture

Bioprosthetic Valve Fracture (BVF) is a technique that uses a high pressure dilatation with

intent to purposefully fracture or crack the ring of the previously implanted bioprosthetic valve and allow the new implanted valve to more fully expand. This technique requires balloon

pressures of up to 20 atm.

Source: STS/ACC TVT Registry

Technical Specification

Code: 112000001287 Code System: ACC NCDR

Short Name: TMVR_BVFAttempt

Missing Data: Report

Harvested: Yes (TMVR)

Is Identifier: No Is Base Element: Yes

Is Followup No Element:

Data Type: BL

Precision:

Selection Type: Single Unit of Measure:

Default Value: Null

Usual Range:

Valid Range:

Data Source: User

Parent/Child Validation

Element: 13754 Transcatheter Mitral Valve

Replacement Type

Operator: Equal

Value: Valve-in-Valve

Element: 13754 Transcatheter Mitral Valve

Replacement Type

Operator: Equal Value: Valve-in-Ring

Element: 14481 TMVR Bioprosthetic Valve Fracture Timing

Coding Instruction: Indicate the timing of the bioprosthetic valve fracture.

Note: If BVF was attempted both pre and post valve implant, code both.

Target Value: The value on current procedure

Technical Specification

Code: 112000001287 Code System: ACC NCDR

Short Name: TMVR_BVFTiming Missing Data: Report

Harvested: Yes (TMVR)

Is Identifier: No

Is Base Element: Yes

Is Followup Element:

Data Type: CD

Precision:

Selection Type: Multiple

Unit of Measure:

Default Value: Null

Usual Range:

Valid Range:

Data Source: User

Parent/Child Validation

Element: 14480 TMVR Bioprosthetic Valve Fracture Attempted

Operator: Equal

Value: Yes

Timing - 1.3.6.1.4.1.19376.1.4.1.6.5.729

Selection	Definition	Source	Code	Code System
Pre Implant			112000001912	ACC NCDR
Post Implant			112000001913	ACC NCDR



Element: 14482



Full Specifications **Data Dictionary v3.0**



Coding Instruction: Indicate if the valve was observed to be fractured. Documentation can include any of the

following

(1) Fluoroscopically by either visualizing the waist of the balloon release and/or the fractured

valve ring (if the valve ring is radiopaque);

TMVR Valve Observed to be Fractured

(2) By an audible snap, or

(3) By a sudden drop in the balloon pressure in the absence of balloon rupture.

Target Value: The value on current procedure

Technical Specification

Code: 112000001290

Code System: ACC NCDR

Short Name: TMVR_ValveFractured

Missing Data: Report Harvested: Yes (TMVR)

Harvested: Yes (TM)
Is Identifier: No

Is Base Element: Yes

Is Followup Element: No Data Type: BL Precision:

Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14480 TMVR Bioprosthetic Valve

Fracture Attempted

Operator: Equal Value: Yes

Element: 13756 Transcatheter Mitral Valve Replacement Primary Procedure Indication

Coding Instruction: Indicate the primary procedure indication for the TMVR procedure. If more than one indication

is present, choose the most significant.

Target Value: The highest value between 2 months prior to current procedure and current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (TMVR) then

Transcatheter Mitral Valve Replacement Primary Procedure Indication (13756) cannot be Null

Technical Specification

Code: 112000000482
Code System: ACC NCDR
Short Name: TMVRProcedureInd

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal

Value: TMVR

Transcatheter Mitral Valve Replacement Primary Procedure Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.553

Selection	Definition	Source	Code	Code System
Mitral Stenosis			112000001459	ACC NCDR
Mitral Regurgitation			48724000	SNOMED CT







Section: TMVR Parent: Procedure Information

Element: 13758 Mitral Valve Replacement - Procedure Access Site

Coding Instruction: Indicate the access site used to perform the mitral procedure.

Target Value: The last value on current procedure

Technical Specification

Code: 112000001474
Code System: ACC NCDR
Short Name: MVAccessSite
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Transcatheter Mitral Valve Replacement Procedure Access Site - 1.3.6.1.4.1.19376.1.4.1.6.5.556

Selection	Definition	Source	Code	Code System
Transseptal via Femo	oral Vein		112000001296	ACC NCDR
Transapical			112000001295	ACC NCDR
Direct Left Atrium			112000001475	ACC NCDR
Other			100000351	ACC NCDR

Element: 13759 Preimplant Balloon Inflation Performed

Coding Instruction: Indicate if pre-implant balloon inflation was performed.

Target Value: The value on current procedure

Technical Specification

Code: 112000001476
Code System: ACC NCDR
Short Name: MVR_MVPreBalloon

Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: TMVR Parent: Procedure Information

Element: 13760 Significant Hemodynamic Deterioration After Inflation

Coding Instruction: Indicate if significant hemodynamic deterioration occurred after inflation. The patient would

valve can cause severe mitral regurgitation.

experience hypotension and pulmonary congestion because balloon inflation of the stenotic

Target Value: The value on current procedure

Technical Specification

Code: 112000001477 Code System: ACC NCDR Short Name: MVR_MVHemDet

Missing Data: Report Harvested: Yes (TMVR)

Is Identifier: No Is Base Element: Yes Is Followup No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range:

Parent/Child Validation

Element: 13759 Preimplant Balloon Inflation

Performed

Valid Range: Data Source: User

Operator: Equal Value: Yes

Element: 13761 Post Implant Balloon Inflation Performed

Coding Instruction: Indicate if post-implant balloon inflation was performed.

Target Value: The value on current procedure

Technical Specification

Code: 112000001478 Code System: ACC NCDR Short Name: MVR_MVPostBalloon

Missing Data: Report Harvested: Yes (TMVR) Is Identifier: No

Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: TMVR Devices Parent: TMVR

Element: 13532 Mitral Valve Device Counter

Coding Instruction: This is a software-assigned value. The counter will start at one and be incremented by one for

each device or system used.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.851

Code System: ACC NCDR Short Name: MVDevCounter Missing Data: Illegal

Harvested: Yes (BDS, TMVR)

Is Base Element: Yes Is Followup Element: No Data Type: CTR Precision: 3 Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: 1 - 999 Data Source: Automatic

Is Identifier: No

Parent/Child Validation

Element: 13505 Procedure Aborted

Operator: Equal Value: No

AND Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14484 Transcatheter Mitral Valve Replacement Device ID

Coding Instruction: Indicate the device ID of the mitral valve.

Target Value: The value on current procedure

Technical Specification

Code: 112000001807 Code System: ACC NCDR Short Name: TMVRDeviceID

Missing Data: Illegal

Harvested: Yes (BDS, TMVR)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision:

Selection Type: Single (Dynamic List)

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 13532 Mitral Valve Device Counter Operator:

Value: Any Value







Section: TMVR Devices Parent: TMVR

Element: 14521 Transcatheter Mitral Valve Replacement Device Diameter

Coding Instruction: Indicate the transcatheter mitral valve replacement device diameter (in mm).

Target Value: The value on current procedure

Technical Specification

Code: 112000001807
Code System: ACC NCDR
Short Name: TMVRDeviceDia

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 10 - 36 mm

Valid Range: 5 - 100 mm Data Source: User

Parent/Child Validation

Element: 14484 Transcatheter Mitral Valve

Replacement Device ID

Operator:

Value: Any Value

Element: 14288 Transcatheter Mitral Valve Replacement Device Serial Number

Coding Instruction: Indicate the transcatheter mitral valve replacement device serial number.

Target Value: The value on current procedure

Technical Specification

Code: 112000001807 Code System: ACC NCDR

Short Name: TMVReplacementDeviceSN

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TMVR)

Is Base Element: Yes
Is Followup
Element: No
Data Type: ST
Precision: 30
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13538 Mitral Valve Device Implanted

Successfully

Operator: Equal Value: Yes







Section: TMVR Devices Parent: TMVR

Element: 14573 Transcatheter Mitral Valve Unique Device ID

Coding Instruction: Indicate the full unique device identifier (UDI) for the implanted device

Target Value: The value on current procedure

Supporting Definition: Unique Device Identifier (UDI)

An identifier that is the main (primary) lookup for a medical device product and meets the requirements to uniquely identify a device through its distribution and use. This value is

supplied to the FDA by the manufacturer.

Source: US FDA

Technical Specification

Code: 2.16.840.1.113883.3.3719

Code System: ACC NCDR Short Name: TMV_UDI Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: ST
Precision: 150
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 13538 Mitral Valve Device Implanted

Successfully

Operator: Equal Value: Yes

Element: 13538 Mitral Valve Device Implanted Successfully

Coding Instruction: Indicate if there is correct positioning of a single prosthetic heart valve in the proper anatomical

location.

Target Value: The value on current procedure

Technical Specification

Code: 17107009 Code System: SNOMED CT

Short Name: MVDeviceImplantSuccessful

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TMVR)

Is Base Element: Yes
Is Followup
Element: No
Bata Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13532 Mitral Valve Device Counter

Operator:

Value: Any Value







Section: TMVR Devices Parent: TMVR

Element: 13541 Reason Mitral Valve Device Not Implanted Successfully

Coding Instruction: Indicate the reason the device was not implanted successfully.

Target Value: The value on current procedure

Technical Specification

Code: 112000002014
Code System: ACC NCDR
Short Name: MV_Unsuccessful

Missing Data: Report Harvested: Yes (TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13538 Mitral Valve Device Implanted

Successfully

Operator: Equal Value: No

Transcatheter Valve Therapy Reason Device Not Implanted Successfully - 1.3.6.1.4.1.19376.1.4.1.6.5.512

Selection	Definition	Source	Code	Code System
Device Embolization			112000001324	ACC NCDR
Improper Device Posit	tioning		112000001325	ACC NCDR
Improper Device Sizin	ng		112000001326	ACC NCDR
Other			100000351	ACC NCDR







Section: TTVP Parent: Procedure Information

Element: 13815 Tricuspid Valve Procedure Type

Coding Instruction: Indicate the type of transcatheter tricuspid valve intervention.

Target Value: The value on current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (Tricuspid Valve

Procedure) then Tricuspid Valve Procedure Type (13815) cannot be Null

Technical Specification

Code: 232778005
Code System: SNOMED CT
Short Name: TVProcType
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal

Value: Tricuspid Valve Procedure

Tricuspid Valve Procedure Type - 1.3.6.1.4.1.19376.1.4.1.6.5.564

Selection	Definition	Source	Code	Code System
Annular Reduction			112000001516	ACC NCDR
Direct Leaflet			112000001517	ACC NCDR
Tricuspid Valve Replac	cement		25236004	SNOMED CT

Element: 13816 Tricuspid Valve Replacement Location

Coding Instruction: Indicate the location of the tricuspid valve replacement.

Target Value: The value on current procedure

Vendor Instruction: Tricuspid Valve Replacement Location (13816) must not be Equal to (Native Valve) when

Procedure History Name (12905) is (Tricuspid Valve Replacement OR Tricuspid Valve Replacement - Transcatheter) and the Procedure History Occurrence (14268) is (Yes)

Technical Specification

Code: 25236004

Code System: SNOMED CT
Short Name: TVLocation
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13815 Tricuspid Valve Procedure Type

Operator: Equal

Value: Tricuspid Valve Replacement

Tricuspid Valve Replacement Location - 1.3.6.1.4.1.19376.1.4.1.6.5.565

Selection	Definition	Source	Code	Code System
Inferior and Superior Vena Cava		112000001522	ACC NCDR	
Inferior Vena Cava			64131007	SNOMED CT
Native Valve			112000001519	ACC NCDR
Surgical Ring			112000001521	ACC NCDR
Surgical Valve			112000001520	ACC NCDR







Section: TTVP Parent: Procedure Information

Element: 13817 Tricuspid Valve Repair or Replacement Procedure Indication

Coding Instruction: Indicate the primary procedure indication for the tricuspid procedure.

Target Value: The value on current procedure

Vendor Instruction: When Transcatheter Valve Therapy Procedure Type (14273) is Equal to (Tricuspid Valve

Procedure) then Tricuspid Valve Repair or Replacement Procedure Indication (13817) cannot

e Null

Technical Specification

Code: 112000000482
Code System: ACC NCDR
Short Name: TVProcedureInd

Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Tricuspid Valve Repair or Replacement Procedure Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.566

Selection	Definition	Source	Code	Code System
Tricuspid Valve Regurgitation		111287006	SNOMED CT	
Tricuspid Valve Stenosis 49		49915006	SNOMED CT	
Both Tricuspid Stenosis and at			112000001829	ACC NCDR
least Moderate Tricus	spid			
Regurgitation				

Element: 13838 Tricuspid Valve Procedure Access Site

Coding Instruction: Indicate the access site used to perform the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 112000001474
Code System: ACC NCDR
Short Name: TVAccess
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Tricuspid Valve Replacement Procedure Access Site - 1.3.6.1.4.1.19376.1.4.1.6.5.567

Selection	Definition	Source	Code	Code System
Femoral Vein			83419000	SNOMED CT
Jugular Vein			63190004	SNOMED CT
Right Atrium			73829009	SNOMED CT
Other			100000351	ACC NCDR







Section: TTVP Parent: Procedure Information

Element: 13839 Transvenous Right Ventricular Lead Present

Coding Instruction: Indicate if a transvenous right ventricular lead is present.

Target Value: The value on current procedure

Technical Specification

Code: 112000001526
Code System: ACC NCDR
Short Name: RVLead
Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Valid Range: Data Source: User

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13840 Right Ventricular Lead Strategy

Coding Instruction: Indicate the strategy to manage the right ventricular lead.

Target Value: The value on current procedure

Technical Specification

Code: 112000001529
Code System: ACC NCDR
Short Name: RVLeadStrat
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13839 Transvenous Right Ventricular

Lead Present

Operator: Equal

Value: Yes

Right Ventricular Lead Strategy - 1.3.6.1.4.1.19376.1.4.1.6.5.568

Selection	Definition	Source	Code	Code System
Jailed by Transcathe	eter Valve		112000001528	ACC NCDR
Lead Removed Prior	r to Valve		112000001527	ACC NCDR
Implant				







Section: TTVP Parent: Procedure Information

Element: 13841 Change in Lead Function Technical Specification

Coding Instruction: Indicate if jailing the right ventricular lead led to a change in lead function.

Target Value: The value on current procedure

Code: 112000001529
Code System: ACC NCDR
Short Name: RVLeadFx
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13840 Right Ventricular Lead Strategy

Operator: Equal

Value: Jailed by Transcatheter Valve







Section: TTVP Pre-Implant Parent: TTVP

Element: 13819 Preimplant Superior Vena Cava Pressure

Coding Instruction: Indicate the pressure in the superior vena cava prior to the device implant.

Target Value: The value between start of procedure and prior to the intervention

Technical Specification

Code: 112000001524 Code System: ACC NCDR Short Name: SVDPre Missing Data: Report

Harvested: Yes (TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Data Type: PQ

Precision: 2,0 Selection Type: Single Unit of Measure: mm[Hg] Default Value: Null

Usual Range: 1 - 10 mm[Hg] Valid Range: 0 - 35 mm[Hg] Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava

AND

Element: 13820 Preimplant Superior Vena Cava Pressure Not Documented

Operator: Equal Value: No (or Not Answered)

Element: 13820 Preimplant Superior Vena Cava Pressure Not Documented

Coding Instruction: Indicate if the pressure in the superior vena cava pre-implant was not documented.

Target Value: N/A

Technical Specification

Code: 112000001524 Code System: ACC NCDR Short Name: SVDPreND Missing Data: Report Harvested: Yes (TTVP) Is Identifier: No Is Base Element: Yes

Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location Operator: Equal

Valid Range: Data Source: User

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava







Section: TTVP Pre-Implant Parent: TTVP

Element: 13823 Preimplant Inferior Vena Cava Pressure

Coding Instruction: Indicate the pressure in the inferior vena cava prior to device implant.

Target Value: The value between start of procedure and prior to the intervention

Technical Specification

Code: 112000001525
Code System: ACC NCDR
Short Name: IVCPre
Missing Data: Report

Usual Range: 1 - 10 mm[Hg] Valid Range: 0 - 35 mm[Hg] Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava

AND

Element: 13825 Preimplant Inferior Vena Cava

Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13825 Preimplant Inferior Vena Cava Pressure Not Documented

Coding Instruction: Indicate if the pressure in the inferior vena cava, pre-implant was not documented.

Target Value: N/A

Technical Specification

Code: 112000001525
Code System: ACC NCDR
Short Name: IVCPreND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Valid Range: Data Source: User

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava







Section: TTVP Pre-Implant Parent: TTVP

Element: 13827 Preimplant Right Atrial Pressure

Coding Instruction: Indicate the mean right atrial pressure, pre-implant.

Target Value: The value between start of procedure and prior to the intervention

Technical Specification

Code: 276755008
Code System: SNOMED CT
Short Name: RAPPre
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm/[Hg]
Default Value: Null
Usual Pange: 1 - 10 mm

Usual Range: 1 - 10 mm[Hg]
Valid Range: 0 - 35 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14290 Preimplant Right Atrial Pressure

Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14290 Preimplant Right Atrial Pressure Not Documented

Coding Instruction: Indicate if the mean right atrial pressure is not documented.

Target Value: N/A

Technical Specification

Code: 276755008
Code System: SNOMED CT
Short Name: RAPPreND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Element: NO
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: TTVP Pre-Implant Parent: TTVP

Element: 14281 Preimplant Right Ventricular Systolic Pressure

Coding Instruction: Indicate the right ventricular systolic pressure, preimplant .

Target Value: The value between start of procedure and prior to the intervention

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSPPre
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 10 - 80 mm[Hg] Valid Range: 1 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13831 Preimplant Right Ventricular

Systolic Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND ------ Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13831 Preimplant Right Ventricular Systolic Pressure Not Documented

Coding Instruction: Indicate if the right ventricular systolic pressure, pre-implant was not documented.

Target Value: N/A

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSPPreND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Valid Range: Data Source: User

Operator: Equal







Section: TTVP Pre-Implant Parent: TTVP

Element: 13834 Preimplant Tricuspid Valve Diastolic Gradient

Coding Instruction: Indicate the tricuspid valve diastolic gradient, pre-implant.

Target Value: The value between start of procedure and prior to the intervention

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: TVDGradPre
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 1-15 mm

Usual Range: 1 - 15 mm[Hg] Valid Range: 1 - 50 mm[Hg] Data Source: User

Parent/Child Validation

Element: 13836 Preimplant Tricuspid Valve

Diastolic Gradient Not Documented

Operator: Equal

Value: No (or Not Answered)

AND -----

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13836 Preimplant Tricuspid Valve Diastolic Gradient Not Documented

Coding Instruction: Indicate if the tricuspid valve diastolic gradient was not documented pre implant.

Target Value: N/A

Technical Specification

Code: 112000001512 Code System: ACC NCDR Short Name: TVDGradPreND

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Element: No Data Type: BL Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: TTVP Post-Implant Parent: TTVP

Element: 13821 Post Implant Superior Vena Cava Pressure

Coding Instruction: Indicate the pressure in the superior vena cava post-implant.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 112000001524
Code System: ACC NCDR
Short Name: SVDPost
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 1 - 10 mm

Usual Range: 1 - 10 mm[Hg] Valid Range: 0 - 35 mm[Hg] Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava

AND

Element: 13822 Post Implant Superior Vena Cava

Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13822 Post Implant Superior Vena Cava Pressure Not Documented

Coding Instruction: Indicate if the pressure in the superior vena cava post-implant was not documented.

Target Value: N/A

Technical Specification

Code: 112000001524
Code System: ACC NCDR
Short Name: SVDPostND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava







Section: TTVP Post-Implant Parent: TTVP

Element: 13824 Post Implant Inferior Vena Cava Pressure

Coding Instruction: Indicate the pressure in the inferior vena cava post-implant.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 112000001525
Code System: ACC NCDR
Short Name: IVCPost
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 1 - 10 mm[Hg]

Usual Range: 1 - 10 mm[Hg]
Valid Range: 0 - 35 mm[Hg]
Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava

AND

Element: 13826 Post Implant Inferior Vena Cava

Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13826 Post Implant Inferior Vena Cava Pressure Not Documented

Coding Instruction: Indicate the pressure in the inferior vena cava post-implant was not documented.

Target Value: N/A

Technical Specification

Code: 112000001525
Code System: ACC NCDR
Short Name: IVCPostND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

it of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior Vena Cava

Element: 13816 Tricuspid Valve Replacement

Location

Operator: Equal

Value: Inferior and Superior Vena Cava







Section: TTVP Post-Implant Parent: TTVP

Element: 13828 Post Implant Right Atrial Pressure

Coding Instruction: Indicate the mean right atrial pressure, post implant.

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 276755008
Code System: SNOMED CT
Short Name: RAPPost
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 1 - 10 mm

Usual Range: 1 - 10 mm[Hg]
Valid Range: 0 - 35 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13830 Post Implant Right Atrial Pressure

Not Documented

Operator: Equal

Value: No (or Not Answered)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13830 Post Implant Right Atrial Pressure Not Documented

Coding Instruction: Indicate if the mean right atrial pressure, post-implant, was not documented.

Target Value: N/A

Technical Specification

Code: 276755008
Code System: SNOMED CT
Short Name: RAPPostND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Element:

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: TTVP Post-Implant Parent: TTVP

Element: 13832 Post Implant Right Ventricular Systolic Pressure

Coding Instruction: Indicate the right ventricular systolic pressure, post-implant .

Target Value: The last value between the implant and the end of current procedure

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSPPost
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]

Default Value: Null
Usual Range: 10 - 80 mm[Hg]
Valid Range: 1 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13833 Post Implant Right Ventricular

Systolic Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

AND -----

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13833 Post Implant Right Ventricular Systolic Pressure Not Documented

Coding Instruction: Indicate if the right ventricular systolic pressure, post-implant was not documented.

Target Value: N/A

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: RVSPPostND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal







Section: TTVP Post-Implant Parent: TTVP

Element: 13835 Post Implant Tricuspid Valve Diastolic Gradient

 $\begin{tabular}{ll} \textbf{Coding Instruction:} & \textbf{Indicate the tricuspid valve diastolic gradient, post-implant.} \\ \end{tabular}$

Target Value: The last value between the implant and the end of current procedure

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: TVDGradPost
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 1 - 15 mm

Usual Range: 1 - 15 mm[Hg]
Valid Range: 1 - 50 mm[Hg]
Data Source: User

Parent/Child Validation

Element: 13837 Post Implant Tricuspid Valve

Diastolic Gradient Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13837 Post Implant Tricuspid Valve Diastolic Gradient Not Documented

Coding Instruction: Indicate if the tricuspid valve diastolic gradient was not documented post implant.

Target Value: N/A

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: TVDGradPostND

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: TTVP Devices Parent: TTVP

Element: 13531 Tricuspid Valve Device Counter

Coding Instruction: This is a software-assigned value. The counter will start at one and be incremented by one for each device or system used.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.851

Code System: ACC NCDR Short Name: TVDevCounter

Missing Data: Illegal Harvested: Yes (TTVP) Is Identifier: No

Is Base Element: Yes Is Followup No Data Type: CTR Precision: 3 Selection Type: Single Unit of Measure: Default Value: Null

> Valid Range: 1 - 999 Data Source: Automatic

Usual Range:

Parent/Child Validation

Element: 13505 Procedure Aborted

Operator: Equal Value: No AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14483 Transcatheter Tricuspid Valve Device ID

Coding Instruction: Indicate the device ID of the tricuspid valve.

Target Value: The value on current procedure

Technical Specification

Code: 703201004 Code System: SNOMED CT Short Name: TTVDeviceID Missing Data: Illegal Harvested: Yes (TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD

Selection Type: Single (Dynamic List)

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Precision:

Parent/Child Validation

Element: 13531 Tricuspid Valve Device Counter

Operator:

Value: Any Value







Section: TTVP Devices Parent: TTVP

Element: 14520 Tricuspid Valve Device Diameter

Coding Instruction: Indicate the tricuspid valve device diameter (in mm).

Target Value: The value on current procedure

Technical Specification Code: 703201004

Code System: SNOMED CT Short Name: TTVDeviceDia Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null
Usual Range: 16 - 36 mm
Valid Range: 5 - 100 mm

Data Source: User

Parent/Child Validation

Element: 14483 Transcatheter Tricuspid Valve

Device ID

Value: Any Value

Operator:

Element: 13842 Tricuspid Valve Device Serial Number

Coding Instruction: Indicate the serial number of the tricuspid valve device implanted during the procedure.

Target Value: The value on current procedure

Technical Specification

Code: 703201004
Code System: SNOMED CT
Short Name: TVDeviceSN
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Followup
Element:
Data Type: ST
Precision: 30
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 13537 Tricuspid Valve Device Implanted

Successfully

Operator: Equal Value: Yes







Section: TTVP Devices Parent: TTVP

Element: 14571 Transcatheter Tricuspid Valve Unique Device ID

Coding Instruction: Indicate the full unique device identifier (UDI) for the implanted device

Target Value: The value on current procedure

Supporting Definition: Unique Device Identifier (UDI)

An identifier that is the main (primary) lookup for a medical device product and meets the requirements to uniquely identify a device through its distribution and use. This value is

supplied to the FDA by the manufacturer.

Source: US FDA

Technical Specification

Code: 2.16.840.1.113883.3.3719

Code System: ACC NCDR Short Name: TTV_UDI Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: ST
Precision: 150
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Parent/Child Validation

Element: 13537 Tricuspid Valve Device Implanted

Successfully

Valid Range: Data Source: User

Operator: Equal Value: Yes

Element: 13537 Tricuspid Valve Device Implanted Successfully

Coding Instruction: Indicate if the device was implanted successfully.

Target Value: The value on current procedure

Technical Specification

Code: 703201004 Code System: SNOMED CT

Short Name: TVDeviceImplantSuccessful

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: Bl

Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13531 Tricuspid Valve Device Counter

Operator:

Value: Any Value







Section: TTVP Devices Parent: TTVP

Element: 13540 Reason Tricuspid Valve Device Not Implanted Successfully

Coding Instruction: Indicate the reason the device was not implanted successfully.

Target Value: The value on current procedure

Technical Specification

Code: 112000002014
Code System: ACC NCDR
Short Name: TV_Unsuccessful

Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 13537 Tricuspid Valve Device Implanted

Successfully

Data Source: User

Operator: Equal Value: No

Reason Tricupsid Valve Device Not Implanted Successfully - 1.3.6.1.4.1.19376.1.4.1.6.5.569

Selection	Definition	Source	Code	Code System
Adverse Event			112000001505	ACC NCDR
Anchor Pull Through			112000001530	ACC NCDR
Device Embolization			112000001324	ACC NCDR
Device Malfunction			112000001504	ACC NCDR
Improper Device Positioning			112000001325	ACC NCDR
Improper Device Sizing			112000001326	ACC NCDR
Inability to Deliver Device Anchor			112000001533	ACC NCDR
Inability to Deploy the Stent			112000001532	ACC NCDR
Inability to Deploy the Valve			112000001531	ACC NCDR
Inability to Grasp Leaflets			112000001501	ACC NCDR
Inability to Reduce Annular Dimension			112000001534	ACC NCDR
Inability to Reduce Tricuspid Regurgitation			112000001535	ACC NCDR
Inferior Vena Cava Too Large	•		112000001536	ACC NCDR
Leaflet Detachment			112000001537	ACC NCDR
Single Leaflet Device Attachment			112000001538	ACC NCDR
Tricuspid Valve Injury			112000001539	ACC NCDR
Tricuspid Valve Stenosis			49915006	SNOMED CT
Other			100000351	ACC NCDR







	- Stelle	- Intra or Post-Procedure Events	Parent: Lab Visit			
Element: 12153		Intra or Post Procedure Events			al Specific	ation
Coding Instru	ction:	Indicate if there were any Intra or Post Prod	cedure Events.	Code: Code System:	1000142478 ACC NCDR	
Target '	Value:	e: Any occurrence between start of procedure and until next procedure or discharge		Short Name:		
Vendor Instruction:		When an Intra or Post Procedure Events (1: Events Occurred (9002) must not be Null	2153) are selected then Intra/Post-Procedure	Missing Data: Harvested:	Report Yes (BDS, TATMVrpr, TTV)	
		(14275) - may only be entered/selected one	vents (12153), Occurred (9002) and Event Date ce	Is Identifier: Is Base Element: Is Followup Element: Data Type: Precision: Selection Type: Unit of Measure: Default Value: Usual Range: Valid Range: Data Source:	No Yes No CD Single (Dynam	
Intra or Post Procedure Selection	Events	s - 1.3.6.1.4.1.19376.1.4.1.6.5.706	Source		Code	Code Syster
Annular Rupture	Annula umbrel related ventric transca to the a classifi supra-a	ar rupture (or 'annulus rupture') is an alla term covering different procedural- linjuries of the aortic root and the left ular outflow tract (LVOT) during atheter aortic valve replacement. According anatomical location of the injury, it can be led into 4 types: intra-annular, subannular, annular, and combined rupture an also be called an 'aortic root rupture' and e of the device landing zone.'	Pasic, M, Unbehaun, A, et al. Annular Rupture During Transcatheter Aortic Valve Replacement. JACC Cardiovascular Interventions, Vol 8 (2015), #1, 1-9.	1120	00001835	ACC NCD
Aortic Dissection	Include aortic of percuta classifi dependinvolve relation dissector repair of treated reserve. Type A aortic a aorta. I type III aorta of ascendin reserve. Type E left sub of the a	e only Stanford classification type A or B dissections, requiring surgical or aneous intervention. The Stanford cation is divided into type A and B ding on whether the ascending aorta is ed. The Stanford classification is in close aship to clinical practice, as type A tions generally require primary surgical whereas type B dissections generally are I medically as initial treatment with surgery ed for any complications. A Involves the ascending aorta and/or arch, and possibly the descending aorta. ar can originate in the ascending aorta, the arch, or, more rarely, in the descending ti includes DeBakey type I, II and retrograde (dissection originating in the descending or aortic arch but extending into the ding aorta). B Involves the descending aorta (distal to be clavian artery origin), without involvement ascending aorta or aortic arch. It includes ey type III without retrograde extension into	Poonyagariyagorn H, Hook M, Bhatt DL. Cardiovascular emergencies. In: Cleveland Clinic: Current Clinical Medicine 2009. 1st ed. Philadelphia, Pa: Saunders Elsevier; 2008: chap 14; Ankel F. Aortic dissection. In: Marx JA, ed. Rosen's Emergency Medicine: Concepts and Clinical Practice. 7th ed. Philadelphia, Pa: Mosby Elsevier; 2009: chap 83.	3	08546005	SNOMED C
ASD Defect Closure due o Transseptal	A proc defect	cending aorta. edure was required to close an atrial-septal as a result of the transseptal erization procedure.		1120	00001885	ACC NCD
Atrial Fibrillation	Atrial fi prolong initiatio to addi proced (cardio	brillation procedure. It is in the state of			49436004	SNOMED C

1000142440

Bleeding - Access Site

Indicate if the patient experienced a confirmed bleeding event at the access site observed and

ACC NCDR







Section: Post-Prod	edure - Intra or Post-Procedure Events	Parent: Lab Visit	
	documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells;		
	3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a GI bleed).		
Bleeding - Gastrointestina	I The patient experienced a confirmed gastrointestinal bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a Gl bleed).	7447400:	3 SNOMED C
Bleeding - Genitourinary	Indicate if the patient experienced a confirmed genitourinary bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding.	41794100:	3 SNOMED C
Bleeding - Hematoma at Access Site	Indicate if the patient experienced a confirmed hematoma at the access site observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a Gl bleed).	38549400	B SNOMED C
Bleeding - Other	The patient experienced bleeding from a site not otherwise specified, such as pulmonary bleeding or a subdural hematoma (not a hemorrhagic stroke). To qualify, the bleeding should be associated with any of the following documented in the medical record: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site or balloon angioplasty to seal an arterial tear).		1 ACC NCD
Bleeding - Retroperitoneal	Indicate if the patient experienced a confirmed retroperitoneal bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear).	9554900	1 SNOMED C
Cardiac Arrest	Cardiac arrest is defined as acute cardiac event documented by one of the following: ventricular fibrillation, rapid ventricular tachycardia or bradycardia rhythms with hemodynamic compromise causing loss of consciousness, pulseless rhythms (PEA), or asystole requiring cardiopulmonary resuscitation (two or more chest compressions or open chest massage, emergency temporary pacing, pericardiocentesis,	Data Governance Subcommittee of the NCDR's \$410429000 SQOC	O SNOMED C







	prosthetic valve involvement.			
Endocarditis	Confirmed diagnosis of endocarditis by blood culture and/or vegetation on or around a heart valve. This may include native tissue, ring or	Society of Thoracic Surgeons (STS)	56819008	SNOMED C
Dialysis (New Requirement)	Acute or worsening renal failure necessitating a new requirement for renal dialysis (renal dialysis includes hemodialysis and peritoneal dialysis). If a patient is on receiving continuous veno-venous hemofiltration (CVVH) as a result of renal failure (and not as treatment to remove fluid for heart failure), code yes.		100014076	ACC NCDI
Device Thrombosis	Any thrombus attached to or near the valve that was implanted during the procedure that occludes part of the blood flow path, interferes with valve function, or is sufficiently large to warrant treatment.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001839	ACC NCDI
Device Related Event - Other	Indicate if an otherwise unspecified device-related event requiring unanticipated treatment occurred. This includes any delivery system related event.		112000001828	ACC NCDI
Owigo Deleted Event	valve orientation within the aortic outflow track resulting in a new echocardiographic confirmed flow disturbance (pre- and post- filmed documentation). Note: Code device embolization if the device is outside of the outflow tract.		442000004929	ACC NCD
Device Migration	Device migration of the prosthetic valve is x-ray confirmed movement of the valve from its initial implantation site such that there is a change in		370512004	SNOMED C
Device Embolization	The device became displaced from its initial implantation site so that it is no longer in its original position.		112000001324	ACC NCD
Delivery System Component Embolization	A component of the delivery system became detached and embolized into the heart or vascular system of the patient.		112000001841	ACC NCD
	performed at your facility if it is accompanied by appropriate documentation in the medical record by the provider. Code no if documentation ONLY included antibody testing (IgG).			
	Notes: It is acceptable to code the diagnosis of COVID-19 based on testing that was NOT			
COVID-19 Positive	occurring during or after the procedure. The patient had a diagnosis of COVID-19, confirmed with a laboratory performed polymerase chain reaction (PCR) test.		112000001982	ACC NCD
Compression	new, partial or complete obstruction of a coronary ostium, either by the valve prosthesis itself, the native leaflets, calcifications, or dissection,		11200001037	ACC NOD
Complete Leaflet Clip Detachment Coronary Artery	A complete detachment of the leaflet clip from the mitral valve leaflets occurred. Angiographic or echocardiographic evidence of a		112000001840	ACC NCD
Intervention Other Unplanned	surgery or a cath lab intervention that was unplanned. This does not include an intervention or procedure already identified as an adverse event in the TVT Registry (e.g. AV reintervention, other vascular surgery or intervention, pacemaker or ICD implant).			
Cardiac Surgery or	aorta, with or without tamponade associated with the perforation. If tamponade occurs there would be fluid in the pericardial space compromising cardiac filling, and requiring intervention such as pericardiocentesis or returning to the operating room. This should be documented by either: 1. Echo showing pericardial fluid and signs of tamponade such as right heart compromise, or 2. Systemic hypotension due to pericardial fluid compromising cardiac function. The patient subsequently underwent cardiac		112000001892	ACC NCDI
Cardiac Perforation	institution of ECMO, or defibrillation) and without these measures death would have almost certainly resulted. A perforation of the myocardium, aortic annulus or		36191001:123005000=302509004	SNOMED C







Section. Post-Proc	edure - Intra or Post-Procedure Events	Parent: Lab Visit		
	requiring insertion of an implantable cardioverter/defibrillator.			
Left Ventricular Outflow Fract Obstruction	Left ventricular outflow tract obstruction (pressure gradient assessed by with echo-Doppler velocities or by catheter-based pressure measurement) was documented in the medical record.		253546004	SNOMED C
Aitral Leaflet or Subvalvular Injury	A mitral leaflet or subvalvular injury was detected during surgery or ascertained by echocardiogram.		112000001886	ACC NCD
Myocardial Infarction	documented and clinically significant myocardial necrosis. The MI can be periprocedural (<72 hours after the procedure) or spontaneous (>72 hours after the index procedure). 1. Peri-procedural MI (<72 h after the index procedure) (a) New ischemic symptoms (e.g. chest pain or	Transcatheter Aortic Valve Implantation Clinical Trials (JACC, 2012, vol 60, No 15)	22298006	SNOMED C
	shortness of breath), or new ischemic signs (e.g. ventricular arrhythmias, new or worsening heart failure, new ST-segment changes, hemodynamic instability, new pathological Q waves in at least two contiguous leads, imaging evidence of new loss of viable myocardium or new wall motion abnormality) AND			
	(b) Elevated cardiac biomarkers (preferable CK-MB) within 72 h after the indexprocedure, consisting of at least one sample post-procedure with a peak value exceeding 15 x as the upper reference limit for troponin or 5 x forCK-MB.* If cardiac biomarkers are increased at baseline (>99th percentile), a further increase in at least 50% post-procedure is required AND the peak value must exceed the previously stated limit.			
	2. Spontaneous MI (_72 h after the index procedure) any one of the following criteria:			
	(a) Detection of rise and/or fall of cardiac biomarkers (preferably troponin) with at least one value above the 99th percentile URL, together with the evidence of myocardial ischemia with at least one of the following: -Symptoms of ischemia -ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] -New pathological Q-waves in at least two contiguous leads -Imaging evidence of a new loss of viable myocardium or new wall motion abnormality			
	(b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood.			
	(c) Pathological findings of an acute myocardial infarction.			
Pacemaker Lead Dislodgement or Dysfunction	Pacemaker lead dislodgement or pacemaker dysfunction was documented in the medical record		112000001884	ACC NCD
Percutaneous Coronary Intervention		National Cardiovascular Data Registry (NCDR)	415070008	SNOMED C
ermanent Pacemaker	The patient developed a new dysrhythmia requiring insertion of a permanent pacemaker.		449397007	SNOMED C







Section: Post-Proc	edure - Intra or Post-Procedure Events	Parent: Lab Visit		
Pulmonary Embolism	Intravascular migration of a venous thrombus to the pulmonary arterial circulation. A 'Proved Pulmonary Embolism' is proved by a positive pulmonary angiogram, an unequivocally positive helical CT scan, a high-probability ventilation-perfusion scan, or autopsy.	Banovac, F., et al. Reporting Standards for Endovascular Treatment of Pulmonary Embolism. Journal of Vascular Interventional Radiology 2010; 21:44–53	59282003	SNOMED C
Reintervention - Aortic Valve	The patient returned to the operating room or cath lab for any aortic valve re-intervention.		112000001827	ACC NCDI
	Note: Please complete adjudication worksheet for every documented aortic valve reintervention, regardless of type of reintervention.			
Reintervention - Mitral Valve	The patient returned to the operating room or cath lab for any mitral valve re-intervention.		112000001893	ACC NCDF
	Note: Please complete adjudication worksheet for every documented mitral valve reintervention, regardless of type of reintervention.			
Reintervention - Tricuspid Valve	The patient returned to the operating room or cath lab for any tricuspid valve re-intervention.		112000001820	ACC NCDF
	Note: Please complete adjudication worksheet for every documented tricuspid valve reintervention, regardless of type of reintervention.			
Single Leaflet Device Attachment	Single leaflet device attachment was documented in the medical record.		112000001538	ACC NCDF
Stroke - Hemorrhagic	An acute episode of focal or global cerebral or spinal dysfunction caused by intraparenchymal, intraventricular or subarachnoid hemorrhage.	Hicks, K., Tcheng, J. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials A Report of the ACC/AHA Task Force on Clinical Data Standards	230706003	SNOMED CT
	Note: Subdural hematomas are intracranial hemorrhagic events and not strokes.	(Writing Committee to Develop Cardiovascular Endpoints Data Standards). JACC 2015, 66 (4), p 403-469		
Stroke - Ischemic	An acute episode of focal cerebral, spinal, or retinal dysfunction caused by infarction of central nervous system tissue.	Hicks, K., Tcheng, J. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials A Report of the ACC/AHA Task Force on Clinical Data Standards	422504002	SNOMED CT
	Note: Hemorrhage may be a consequence of ischemic stroke. In this situation, the stroke is an ischemic stroke with hemorrhagic transformation and not a hemorrhagic stroke.	(Writing Committee to Develop Cardiovascular Endpoints Data Standards). JACC 2015, 66 (4), p 403-469		
Stroke - Undetermined	A stroke of undetermined origin is defined as an acute episode of focal or global neurological dysfunction caused by presumed brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction but with insufficient information to allow categorization as ischemic or hemorrhagic.	Hicks KA, Tcheng JE, Bozkurt B, et al. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards (Writing Committee to Develop Cardiovascular Endpoints Data Standards). J Am Coll Cardiol. 2015;66 (4):403-469. doi:10.1016/j.jacc.2014.12.018.	230713003	SNOMED CT
Transient Ischemic Attack (TIA)	A transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.	Society for Thoracic Surgeons (STS)	266257000	SNOMED CT
Transseptal Complication	The patient experienced an adverse event as a result of the transseptal access.		112000001833	ACC NCDF
Vascular Complication - Major	Major vascular complications include any of the following: 1. Any aortic dissection, aortic rupture, annulus rupture, left ventricle perforation, or new apical aneurysm/pseudo-aneurysm; 2. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneurysm, hematoma, irreversible nerve injury, compartment syndrome, percutaneous closure device failure) leading to death, life threatening or major bleeding*, visceral ischemia or neurological impairment; 3. Distal embolization (non-cerebral) from a vascular source requiring surgery or resulting in amputation or irreversible end-organ damage; 4. The use of unplanned endovascular or surgical intervention associated with death, major bleeding		11200000460	ACC NCDF







Section: Post-Procedure - Intra or Post-Procedure Events Parent: Lab Visit 5. Any new ischemia documented by patient symptoms, physical exam, and/or decreased or absent blood flow on lower extremity angiogram; 6. Surgery for access site-related nerve injury; 7. Permanent access site-related nerve injury. *Refers to VARC bleeding definitions Note: "ipsilateral lower extremity" was removed from #5 to have the ability to account for ischemia from any access site. ACC NCDR Vascular Complication -Minor vascular complications include any of the Updated Standardized Endpoint Definitions for 112000001823 Transcatheter Aortic Valve Implantation (JACC, following: Minor 1. Access site or access-related vascular injury 2012, Vol 60, No 15) (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneuysms, hematomas, percutaneous closure device failure) not leading to death, life-threatening or major bleeding*, visceral ischemia or neurological impairment; 2. Distal embolization treated with embolectomy and/or thrombectomy and not resulting in amputation or irreversible end-organ damage: 3. Any unplanned endovascular stenting or unplanned surgical intervention not meeting the criteria for a major vascular complication; 4. Vascular repair or the need for vascular repair (via surgery, ultrasound-guided compression, transcatheter embolization, or stent-graft). *Refers to VARC bleeding definitions Vascular Surgery or The patient required unplanned vascular surgery 112000000467 ACC NCDR Intervention - Unplanned or intervention to correct a bleeding complication or vascular related complication. Note: If a balloon angioplasty of the access site or access related sites is performed as a routine procedure to ensure adequate hemostasis of the site, then this would not qualify as an Unplanned Vascular Surgery or Intervention. However, if a

Element: 9002 Intra/Post-Procedure Events Occurred

captured.

Coding Instruction: Indicate if the specific intra or post procedure event(s) occurred.

balloon angioplasty is performed in an attempt to treat a bleeding or vascular access complication (i.e. bleeding at access site, dissection, stenosis, narrowing of vessel, etc.), then Unplanned Vascular Surgery or Intervention should be

Target Value: Any occurrence between start of procedure and until next procedure or discharge

Technical Specification

Code: 1000142479

Code System: ACC NCDR

Short Name: PostProcOccurred

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 12153 Intra or Post Procedure Events

Operator:

Value: Any Value







Section: Post-Procedure - Intra or Post-Procedure Events Parent: Lab Visit

Element: 14275 Intra and Post Procedure Event Date

Coding Instruction: Indicate the date the event occurred.

Target Value: Any occurrence between start of procedure and until next procedure or discharge

Vendor Instruction: Intra and Post Procedure Event Date (14275) must be Greater than or Equal to Procedure Start

Date and Time (7000)

Technical Specification

Code: 10001424780 Code System: ACC NCDR

Short Name: IntraPostProcEventDate

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: DT

Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:

Valid Range:

Data Source: User

Parent/Child Validation

Element: 9002 Intra/Post-Procedure Events

Occurred

Operator: Equal Value: Yes







Section: In-Hospita	al Event Information	Parent: Lab Visit		
Element: 14312	Adjudication Event		Technical Specifi	cation
Coding Instru	action: Indicate the event being adjudicated.		Code: 1120000018 Code System: ACC NCDR Short Name: AJ AdjudEv	
_	Undetermined, Transient Ischemic Attack (e: N/A a: When Adjudication Event (14312) is Equal to (Stroke - Hemorrhagic,Stroke - Ischemic,Stroke - Undetermined,Transient Ischemic Attack (TIA)) then Transcatheter Valve Therapy Procedure Type (14273) must be Equal to (TAVR,TMVr,TMVR)		rent TAVR, TMVR, /P)
once		2) and Date (14313) - may only be entered/selected udication Event Code (14312) must match with Intra ara or Post Procedure Event Code (12153)	Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single (Dyna Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User	amic List)
Selection	Definition	Source	Code	Code Systen
Annular Rupture	Annular rupture (or 'annulus rupture') is an umbrella term covering different procedural-related injuries of the aortic root and the left ventricular outflow tract (LVOT) during transcatheter aortic valve replacement. According to the anatomical location of the injury, it can be classified into 4 types: intra-annular, subannular, supra-annular, and combined rupture	Pasic, M, Unbehaun, A, et al. Annular Rupture During Transcatheter Aortic Valve Replacement. JACC Cardiovascular Interventions, Vol 8 (2015), #1, 1-9.	112000001835	ACC NCDF
	This can also be called an 'aortic root rupture' and 'rupture of the device landing zone.'			
Aortic Dissection	Include only Stanford classification type A or B aortic dissections, requiring surgical or percutaneous intervention. The Stanford classification is divided into type A and B depending on whether the ascending aorta is involved. The Stanford classification is in close relationship to clinical practice, as type A dissections generally require primary surgical repair whereas type B dissections generally are treated medically as initial treatment with surgery reserved for any complications.	Poonyagariyagorn H, Hook M, Bhatt DL. Cardiovascular emergencies. In: Cleveland Clinic: Current Clinical Medicine 2009. 1st ed. Philadelphia, Pa: Saunders Elsevier; 2008: chap 14; Ankel F. Aortic dissection. In: Marx JA, ed. Rosen's Emergency Medicine: Concepts and Clinical Practice. 7th ed. Philadelphia, Pa: Mosby Elsevier; 2009: chap 83.	308546005	SNOMED C
	Type A - Involves the ascending aorta and/or aortic arch, and possibly the descending aorta. The tear can originate in the ascending aorta, the aortic arch, or, more rarely, in the descending aorta. It includes DeBakey type I, II and retrograde type III (dissection originating in the descending aorta or aortic arch but extending into the ascending aorta).			
	Type B - Involves the descending aorta (distal to left subclavian artery origin), without involvement of the ascending aorta or aortic arch. It includes DeBakey type III without retrograde extension into the ascending aorta.			
ASD Defect Closure due to Transseptal Catheterization	A procedure was required to close an atrial-septa defect as a result of the transseptal catheterization procedure.	al	112000001885	ACC NCD
Atrial Fibrillation	Atrial fibrillation or flutter requiring treatment or prolonged hospitalization. Treatment includes initiation of a NEW/DIFFERENT medication therapy to address the arrhythmia; or a procedure/intervention to address the arrhythmia (cardioversion, permanent pacemaker/defibrillator ablation, etc.).	,	49436004	SNOMED C







Section: In-Hospita	al Event Information	Parent: Lab Visit	
	documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells;		
	3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a GI bleed).		
Bleeding - Gastrointestina	The patient experienced a confirmed gastrointestinal bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a Gl bleed).	74474003	SNOMED C
Bleeding - Genitourinary	Indicate if the patient experienced a confirmed genitourinary bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding.	417941003	SNOMED C
Bleeding - Hematoma at Access Site	Indicate if the patient experienced a confirmed hematoma at the access site observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear, endoscopy with cautery of a Gl bleed).	385494008	SNOMED C
Bleeding - Other	The patient experienced bleeding from a site not otherwise specified, such as pulmonary bleeding or a subdural hematoma (not a hemorrhagic stroke). To qualify, the bleeding should be associated with any of the following documented in the medical record: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site or balloon angioplasty to seal an arterial tear).		ACC NCDI
Retroperitoneal	Indicate if the patient experienced a confirmed retroperitoneal bleeding event observed and documented in the medical record that was associated with any of the following: 1. Hemoglobin drop of >=3 g/dL; 2. Transfusion of whole blood or packed red blood cells; 3. Procedural intervention/surgery at the bleeding site to reverse/stop or correct the bleeding (such as surgical closures/exploration of the arteriotomy site, balloon angioplasty to seal an arterial tear).	95549001	SNOMED C
Cardiac Arrest	Cardiac arrest is defined as acute cardiac event documented by one of the following: ventricular fibrillation, rapid ventricular tachycardia or bradycardia rhythms with hemodynamic compromise causing loss of consciousness, pulseless rhythms (PEA), or asystole requiring cardiopulmonary resuscitation (two or more chest compressions or open chest massage, emergency temporary pacing, pericardiocentesis,	Data Governance Subcommittee of the NCDR's 410429000 SQOC	SNOMED C







Section: In-Hospit	al Event Information	Parent: Lab Visit		
	institution of ECMO, or defibrillation) and without these measures death would have almost certainly resulted.			
Cardiac Perforation	A perforation of the myocardium, aortic annulus or aorta, with or without tamponade associated with the perforation. If tamponade occurs there would be fluid in the pericardial space compromising cardiac filling, and requiring intervention such as pericardiocentesis or returning to the operating room. This should be documented by either: 1. Echo showing pericardial fluid and signs of tamponade such as right heart compromise, or 2. Systemic hypotension due to pericardial fluid compromising cardiac function.		36191001:123005000=302509004	SNOMED C
Cardiac Surgery or Intervention - Other Unplanned	The patient subsequently underwent cardiac surgery or a cath lab intervention that was unplanned. This does not include an intervention or procedure already identified as an adverse event in the TVT Registry (e.g. AV reintervention, other vascular surgery or intervention, pacemaker or ICD implant).		112000001892	ACC NCDF
Complete Leaflet Clip Detachment	A complete detachment of the leaflet clip from the mitral valve leaflets occurred.		112000001840	ACC NCDF
Coronary Artery Compression	Angiographic or echocardiographic evidence of a new, partial or complete obstruction of a coronary ostium, either by the valve prosthesis itself, the native leaflets, calcifications, or dissection, occurring during or after the procedure.		112000001837	ACC NCDF
COVID-19 Positive	The patient had a diagnosis of COVID-19, confirmed with a laboratory performed polymerase chain reaction (PCR) test. Notes: It is acceptable to code the diagnosis of COVID-19 based on testing that was NOT performed at your facility if it is accompanied by appropriate documentation in the medical record by the provider.		112000001982	ACC NCDR
	Code no if documentation ONLY included antibody testing (IgG).			
Delivery System Component Embolization	A component of the delivery system became detached and embolized into the heart or vascular system of the patient.		112000001841	ACC NCDF
Device Embolization	The device became displaced from its initial implantation site so that it is no longer in its original position.		112000001324	ACC NCDF
Device Migration	Device migration of the prosthetic valve is x-ray confirmed movement of the valve from its initial implantation site such that there is a change in valve orientation within the aortic outflow track resulting in a new echocardiographic confirmed flow disturbance (pre- and post- filmed documentation). Note: Code device embolization if the device is outside of the outflow tract.		370512004	SNOMED CT
Device Related Event - Other	Indicate if an otherwise unspecified device-related event requiring unanticipated treatment occurred. This includes any delivery system related event.		112000001828	ACC NCDR
Device Thrombosis	Any thrombus attached to or near the valve that was implanted during the procedure that occludes part of the blood flow path, interferes with valve function, or is sufficiently large to warrant treatment.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001839	ACC NCDR
Dialysis (New Requirement)	Acute or worsening renal failure necessitating a new requirement for renal dialysis (renal dialysis includes hemodialysis and peritoneal dialysis). If a patient is on receiving continuous veno-venous hemofiltration (CVVH) as a result of renal failure (and not as treatment to remove fluid for heart failure), code yes.		100014076	ACC NCDR
Endocarditis	Confirmed diagnosis of endocarditis by blood culture and/or vegetation on or around a heart valve. This may include native tissue, ring or	Society of Thoracic Surgeons (STS)	56819008	SNOMED CT







Section: In-Hospita	al Event Information	Parent: Lab Visit		
	requiring insertion of an implantable cardioverter/defibrillator.			
eft Ventricular Outflow Fract Obstruction	Left ventricular outflow tract obstruction (pressure gradient assessed by with echo-Doppler velocities or by catheter-based pressure measurement) was documented in the medical record.		253546004	SNOMED C
Aitral Leaflet or Subvalvular Injury	A mitral leaflet or subvalvular injury was detected during surgery or ascertained by echocardiogram.		112000001886	ACC NCD
Vyocardial Infarction		Transcatheter Aortic Valve Implantation Clinical Trials (JACC, 2012, vol 60, No 15)	22298006	SNOMED C
	shortness of breath), or new ischemic signs (e.g. ventricular arrhythmias, new or worsening heart failure, new ST-segment changes, hemodynamic instability, new pathological Q waves in at least two contiguous leads, imaging evidence of new loss of viable myocardium or new wall motion abnormality) AND			
	(b) Elevated cardiac biomarkers (preferable CK-MB) within 72 h after the indexprocedure, consisting of at least one sample post-procedure with a peak value exceeding 15 x as the upper reference limit for troponin or 5 x forCK-MB.* If cardiac biomarkers are increased at baseline (>99th percentile), a further increase in at least 50% post-procedure is required AND the peak value must exceed the previously stated limit.			
	2. Spontaneous MI (_72 h after the index procedure) any one of the following criteria:			
	(a) Detection of rise and/or fall of cardiac biomarkers (preferably troponin) with at least one value above the 99th percentile URL, together with the evidence of myocardial ischemia with at least one of the following: -Symptoms of ischemia -ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] -New pathological Q-waves in at least two contiguous leads -Imaging evidence of a new loss of viable myocardium or new wall motion abnormality			
	(b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood.			
	(c) Pathological findings of an acute myocardial infarction.			
Pacemaker Lead Dislodgement or Dysfunction	Pacemaker lead dislodgement or pacemaker dysfunction was documented in the medical record		112000001884	ACC NCD
Percutaneous Coronary ntervention		National Cardiovascular Data Registry (NCDR)	415070008	SNOMED C
Permanent Pacemaker	The patient developed a new dysrhythmia requiring insertion of a permanent pacemaker.		449397007	SNOMED C







Section: In-Hospita	al Event Information	Parent: Lab Visit		
Pulmonary Embolism	Intravascular migration of a venous thrombus to the pulmonary arterial circulation. A 'Proved Pulmonary Embolism' is proved by a positive pulmonary angiogram, an unequivocally positive helical CT scan, a high-probability ventilation-perfusion scan, or autopsy.	Banovac, F., et al. Reporting Standards for Endovascular Treatment of Pulmonary Embolism. Journal of Vascular Interventional Radiology 2010; 21:44–53	59282003	SNOMED C
Reintervention - Aortic Valve	The patient returned to the operating room or cath lab for any aortic valve re-intervention.		112000001827	ACC NCDF
	Note: Please complete adjudication worksheet for every documented aortic valve reintervention, regardless of type of reintervention.			
Reintervention - Mitral √alve	The patient returned to the operating room or cath lab for any mitral valve re-intervention.		112000001893	ACC NCDF
	Note: Please complete adjudication worksheet for every documented mitral valve reintervention, regardless of type of reintervention.			
Reintervention - Tricuspid Valve	The patient returned to the operating room or cath lab for any tricuspid valve re-intervention.		112000001820	ACC NCDF
	Note: Please complete adjudication worksheet for every documented tricuspid valve reintervention, regardless of type of reintervention.			
Single Leaflet Device Attachment	Single leaflet device attachment was documented in the medical record.		112000001538	ACC NCDF
Stroke - Hemorrhagic	An acute episode of focal or global cerebral or spinal dysfunction caused by intraparenchymal, intraventricular or subarachnoid hemorrhage.	Hicks, K., Tcheng, J. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials A Report of the ACC/AHA Task Force on Clinical Data Standards	230706003	SNOMED CT
	Note: Subdural hematomas are intracranial hemorrhagic events and not strokes.	(Writing Committee to Develop Cardiovascular Endpoints Data Standards). JACC 2015, 66 (4), p 403-469		
Stroke - Ischemic	An acute episode of focal cerebral, spinal, or retinal dysfunction caused by infarction of central nervous system tissue.	Hicks, K., Tcheng, J. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials A Report of the ACC/AHA Task Force on Clinical Data Standards	422504002	SNOMED CT
	Note: Hemorrhage may be a consequence of ischemic stroke. In this situation, the stroke is an ischemic stroke with hemorrhagic transformation and not a hemorrhagic stroke.	(Writing Committee to Develop Cardiovascular Endpoints Data Standards). JACC 2015, 66 (4), p 403-469		
Stroke - Undetermined	A stroke of undetermined origin is defined as an acute episode of focal or global neurological dysfunction caused by presumed brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction but with insufficient information to allow categorization as ischemic or hemorrhagic.	Hicks KA, Tcheng JE, Bozkurt B, et al. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards (Writing Committee to Develop Cardiovascular Endpoints Data Standards). J Am Coll Cardiol. 2015;66 (4):403-469. doi:10.1016/j.jacc.2014.12.018.	230713003	SNOMED CT
Transient Ischemic Attack (TIA)	A transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.	Society for Thoracic Surgeons (STS)	266257000	SNOMED CT
Transseptal Complication	The patient experienced an adverse event as a result of the transseptal access.		112000001833	ACC NCDR
Vascular Complication - Major	Major vascular complications include any of the following: 1. Any aortic dissection, aortic rupture, annulus rupture, left ventricle perforation, or new apical aneurysm/pseudo-aneurysm; 2. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneurysm, hematoma, irreversible nerve injury, compartment syndrome, percutaneous closure device failure) leading to death, life threatening or major bleeding*, visceral ischemia or neurological impairment; 3. Distal embolization (non-cerebral) from a vascular source requiring surgery or resulting in amputation or irreversible end-organ damage; 4. The use of unplanned endovascular or surgical intervention associated with death, major bleeding	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	11200000460	ACC NCDR







Section: In-Hospit	al Event Information	Parent: Lab Visit		
	5. Any new ischemia documented by patient symptoms, physical exam, and/or decreased or absent blood flow on lower extremity angiogram; 6. Surgery for access site-related nerve injury; 7. Permanent access site-related nerve injury. *Refers to VARC bleeding definitions Note: "ipsilateral lower extremity" was removed from #5 to have the ability to account for ischemia from any access site.			
Vascular Complication - Minor	Minor vascular complications include any of the following: 1. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneuysms, hematomas, percutaneous closure device failure) not leading to death, life-threatening or major bleeding*, visceral ischemia or neurological impairment; 2. Distal embolization treated with embolectomy and/or thrombectomy and not resulting in amputation or irreversible end-organ damage; 3. Any unplanned endovascular stenting or unplanned surgical intervention not meeting the criteria for a major vascular complication; 4. Vascular repair or the need for vascular repair (via surgery, ultrasound-guided compression, transcatheter embolization, or stent-graft). *Refers to VARC bleeding definitions	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001823	ACC NCDR
Vascular Surgery or Intervention - Unplanned	The patient required unplanned vascular surgery or intervention to correct a bleeding complication or vascular related complication. Note: If a balloon angioplasty of the access site or access related sites is performed as a routine procedure to ensure adequate hemostasis of the site, then this would not qualify as an Unplanned Vascular Surgery or Intervention. However, if a balloon angioplasty is performed in an attempt to treat a bleeding or vascular access complication (i.e. bleeding at access site, dissection, stenosis, narrowing of vessel, etc.), then Unplanned Vascular Surgery or Intervention should be captured.		11200000467	ACC NCDR







Section: In-Hospital Event Information

Parent: Lab Visit

Element: 14313 Adjudication Event Date

Coding Instruction: Indicate the date the clinical event being adjudicated occurred.

Target Value: N/A

Vendor Instruction: The Adjudication Event Date (14313) / Adjudication Event Code (14312) must match with Intra

or Post-Procedure Event Date (14275) / Intra or Post Procedure Event Code (12153)

Technical Specification

Code: 112000001816
Code System: ACC NCDR
Short Name: AJ_EventDate
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined
Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)
Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Tricuspid Valve

Element: 14314 Adjudication Status

Coding Instruction: Indicate whether the patient was alive or deceased on the date the adjudication was

performed.

Target Value: N/A

Vendor Instruction: Adjudication Status (14314) as 'Deceased' must be answered only once in the episode.

Technical Specification

Code: 112000001817
Code System: ACC NCDR
Short Name: AJ_Status
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
No

Element: NO
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Adjudication Life Status - 1.3.6.1.4.1.19376.1.4.1.6.5.726

Selection	Definition	Source	Code	Code System
Alive			438949009	SNOMED CT
Deceased			20	HL7 Discharge disposition



Element: 14315



Full Specifications **Data Dictionary v3.0**

Visit



Section: In-Hospital Event Information	Parent: Lab

Adjudication Date of Death

Coding Instruction: Indicate the date the patient was declared dead.

Target Value: N/A

Vendor Instruction: Adjudication Date of Death (14315) must be Greater than or Equal to Adjudication Event Date

14313

Technical Specification

Code: 399753006
Code System: SNOMED CT
Short Name: AJ_DeathDate
Missing Data: Report

Is Identifier: No

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Element: 14314 Adjudication Status

Parent/Child Validation

Operator: Equal Value: Deceased

Element: 14462 In Hospital Clinical Comments

Coding Instruction: Provide information and details that may assist in assessing the event(s) being adjudicated.

Target Value: N/A

Technical Specification

Code: 423016009
Code System: SNOMED CT
Short Name: AJ_CommentsInHosp

Missing Data: Report

Is Identifier: No

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: ST
Precision: 1000
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User







Section: Stroke Or TIA

Parent: In-Hospital Event Information

Element: 14316 Symptom Onset Date

Coding Instruction: Indicate the date of symptom onset of the neurologic deficit.

Target Value: N/A

Technical Specification

Code: 112000000125
Code System: ACC NCDR
Short Name: AJ_SxOnset
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined
Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr







Section: Stroke Or TIA

Parent: In-Hospital Event Information

Element: 14317 Neurologic Deficit with Rapid Onset

Coding Instruction: Indicate if the patient had a sudden onset of a focal or global neurologic deficit (regardless of the duration of symptoms) with at least one of the following present: change in level of

consciousness, hemiplegia, hemiparesis, numbness or sensory loss affecting one side of the body, dysphasia or aphasia, hemianopia, amaurosis fugax, other neurological signs or

symptoms consistent with a stroke.

Target Value: N/A

Technical Specification

Code: 264552009
Code System: SNOMED CT
Short Name: AJ_NeuroDef
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined

Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr







Section: Stroke Or TIA Parent: In-Hospital Event Information

Element: 14318 Neurologic Deficit Clinical Presentation

Coding Instruction: Indicate the clinical presentation of the neurologic deficit.

Target Value: N/A

Technical Specification

Code: 264552009
Code System: SNOMED CT
Short Name: AJ_NeuroClinPresent

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14317 Neurologic Deficit with Rapid

Onset
Operator: Equal

Value: Yes

Neurologic Deficit Clinical Presentation - 1.3.6.1.4.1.19376.1.4.1.6.5.716

Selection	Definition	Source	Code	Code System
TIA or Stroke (CVA)			100014109	ACC NCDR
Non Stroke Neurologic Def	icit		112000001860	ACC NCDR

Element: 14319 Neurologic Symptom Duration Greater Than or Equal to 24 hours

Coding Instruction: Indicate if the duration of the neurologic symptoms lasted >= 24 hours.

Target Value: N/A

Technical Specification

Code: 308921004 Code System: SNOMED CT

Short Name: AJ_NeuroSymptDuration

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14318 Neurologic Deficit Clinical

Presentation

Operator: Equal

Value: TIA or Stroke (CVA)







Section: Stroke Or TIA Parent: In-Hospital Event Information

Element: 14320 Brain Imaging Performed

Coding Instruction: Indicate if neuroimaging was performed.

Target Value: N/A

Technical Specification

Code: 441986001 Code System: SNOMED CT Short Name: AJ_BrainImag Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14318 Neurologic Deficit Clinical

Presentation

Operator: Equal

Value: TIA or Stroke (CVA)

Element: 14349 Brain Imaging Type

Coding Instruction: Indicate the type of neuroimaging performed.

Target Value: N/A

Technical Specification

Code: 441986001
Code System: SNOMED CT
Short Name: AJ_BrainImageType

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation
Element: 14320 Brain Imaging Performed

Operator: Equal Value: Yes

Imaging Type - 1.3.6.1.4.1.19376.1.4.1.6.5.417

Selection	Definition	Source	Code	Code System
Computed Tomography			77477000	SNOMED CT
Computed Tomography with Contrast			112000001861	ACC NCDR
Magnetic Resonance Imaging			113091000	SNOMED CT
Magnetic Resonance Imaging with Contrast			51619007	SNOMED CT
Other Imaging			112000001862	ACC NCDR



Element: 14350



Full Specifications Data Dictionary v3.0



Section: Stroke Or TIA Parent: In-Hospital Event Information

Brain Imaging Findings Coding Instruction: Indicate the type of deficit found as a result of the neuroimaging study.

Target Value: N/A

Technical Specification Code: 112000001979 Code System: ACC NCDR Short Name: BI_Find

Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14320 Brain Imaging Performed

Operator: Equal Value: Yes

Brain Imaging Finding - 1.3.6.1.4.1.19376.1.4.1.6.5.717

Selection	Definition	Source	Code	Code System
Infarct	Neuroimaging evidence of CNS infarction in the corresponding vascular territory (brain, spinal cord, or retinal cell death), with or without hemorrhage.	Adapted from: Lansky, A.J., et al. Proposed Standardized Neurological Endpoints for Cardiovascular Clinical Trials (An Academic Research Consortium Initiative) JACC 2017, 69 (6): 679-690	55641003	SNOMED CT
Hemorrhage	Neuroimaging evidence of central nervous system (CNS) hemorrhage within the brain parenchyma, subarachnoid space, ventricular system, spinal cord, or retina that is not caused by trauma.	Adapted from: Lansky, A.J., et al. Proposed Standardized Neurological Endpoints for Cardiovascular Clinical Trials (An Academic Research Consortium Initiative) JACC 2017, 69 (6): 679-690	50960005	SNOMED CT
No Deficit			100001231	ACC NCDR







Section: Stroke Or TIA

Parent: In-Hospital Event Information

Element: 14351 Event Related Sequelae

Coding Instruction: Indicate the sequelae related to the stroke or TIA.

Target Value: N/A

Technical Specification

Code: 362977000
Code System: SNOMED CT
Short Name: Adj_ERS
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Multiple

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14318 Neurologic Deficit Clinical

Presentation
Operator: Equal

Value: TIA or Stroke (CVA)

Event Related Sequelae - 1.3.6.1.4.1.19376.1.4.1.6.5.737

Selection	Definition	Source	Code	Code System
Death			419620001	SNOMED CT
Permanent Vegetativ	ve State		723151005	SNOMED CT
Altered Consciousne	ess		3006004	SNOMED CT
Blindness			193699007	SNOMED CT
Aphasia			87486003	SNOMED CT
Loss of Motor Functi	ion		112000001936	ACC NCDR
Loss of Sensory Fur	nction		33653009	SNOMED CT
Facial Paralysis			280816001	SNOMED CT
Prolonged Length of	Stay		112000001937	ACC NCDR
Other			100000351	ACC NCDR



Element: 14352



Full Specifications Data Dictionary v3.0



Section: Stroke Or TIA Parent: In-Hospital Event Information

Discharge Location After Event Coding Instruction: Indicate the discharge location after the stroke or TIA.

Target Value: N/A

Technical Specification

Code: 75528-0 Code System: LOINC Short Name: AJ_DLAE Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: Yes Is Followup No Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

Is Identifier: No

Valid Range: Data Source: User

Parent/Child Validation

Element: 14314 Adjudication Status Operator: Equal Value: Alive

----- AND -----

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event Operator: Equal

Value: Stroke - Undetermined Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

----- AND -----Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Discharge Location - 1.3.6.1.4.1.19376.1.4.1.6.5.41

Selection	Definition	Source	Code	Code System
Home			01	HL7 Discharge disposition
Skilled Nursing Facility	Skilled nursing facilities (SNF) are typically sub programs used for longer anticipated length of		03	HL7 Discharge disposition
	Note: Sometimes SNFs may have acute rehabileds within their facility. If the patient is dischara SNF for acute rehab (requiring a higher level code "extended care/TCU/rehab".	arged to		
Extended Care/TCU/Rehab	An extended care unit, transitional care unit or unit typically provides a high level of intensive as well as specialized nursing and physician c discharge setting may also be called subacute long term acute care (LTACH).	therapy are. This	62	HL7 Discharge disposition
Other Acute Care Hospital			02	HL7 Discharge disposition
Left Against Medical Advice (AMA)	The patient was discharged or eloped against advice.	medical	07	HL7 Discharge disposition
Other Discharge Location			100001249	ACC NCDR







Section: Stroke Or TIA

Parent: In-Hospital Event Information

Element: 14421 Patient Discharged to Prior Place of Living

Coding Instruction: Indicate if the patient was discharged to their prior place of living.

Target Value: N/A

Technical Specification

Code: 112000001882 Code System: ACC NCDR Short Name: AJ_PriorLiving Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14314 Adjudication Status

Operator: Equal Value: Alive AND

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

AND Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Stroke Or TIA

Parent: In-Hospital Event Information

Element: 14353 Stroke Diagnosed During Autopsy

Coding Instruction: Indicate if the stroke was diagnosed during autopsy.

Target Value: N/A

Technical Specification

Code: 5605004
Code System: SNOMED CT
Short Name: AJ_AutDxStroke

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14314 Adjudication Status

Operator: Equal Value: Deceased

AND ----

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14312 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined
Element: 14312 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Boolean with Information Not Available - 1.3.6.1.4.1.19376.1.4.1.6.5.718

Dociedii With Information Not Available - 1.5.0.1.4.1.13570.1.4.1.0.5.710					
Selection	Definition	Source	Code	Code System	
No			100013073	ACC NCDR	
Yes			100013072	ACC NCDR	
Information Not Avai	ilable		112000001866	ACC NCDR	







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14354 Aortic Valve Reintervention Type

Coding Instruction: Indicate the type of aortic valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: AJ_ReIntType
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacemen	t		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replac	cement		112000001875	ACC NCDR
Balloon Valvuloplasty	/		112000001469	ACC NCDR
Leaflet Clip Procedure	e		112000001778	ACC NCDR
Paravalvular Leak Clo	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14355 Aortic Valve Reintervention Primary Indication

Coding Instruction: Indicate the primary indication for the reintervention. If more than one indication is present,

code the indication the operator feels has the highest significance.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: AJ_PrimaryInd
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation			40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14356 Aortic Valve Regurgitation

Coding Instruction: Indicate the highest level of aortic regurgitation prior to the aortic valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001869 Code System: ACC NCDR Short Name: AJ_AlSev

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14355 Aortic Valve Reintervention

Primary Indication

Operator: Equal

Value: Regurgitation

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14357 Paravalvular Aortic Regurgitation

Coding Instruction: Indicate the highest severity of paravalvular regurgitation prior to the aortic valve

reintervention

Note: If trace/trivial is documented, code "none".

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: AJ_PVSev
Missing Data: Report

Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null Usual Range: Valid Range:

Data Source: User

Parent/Child Validation

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 14356 Aortic Valve Regurgitation

Operator: Equal
Value: Moderate

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14358 Central Aortic Regurgitation

Coding Instruction: Indicate the highest severity of central regurgitation prior to the aortic valve reintervention.

Note: If trace/trivial is documented, code "none".

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: AJ_CenSev
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 14356 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR

Element: 14359 Aortic Valve Area

Coding Instruction: Indicate the smallest aortic valve area (in cm squared).

Target Value: N/A

Technical Specification

Code: 112000001280
Code System: ACC NCDR
Short Name: AJ_AVA
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 0.20 - 4.00 cm2 **Valid Range:** 0.05 - 5.00 cm2

Data Source: User

Parent/Child Validation

Element: 14355 Aortic Valve Reintervention

Primary Indication

Operator: Equal Value: Stenosis







Section: AV Re-Intervention

Parent: In-Hospital Event Information

Element: 14282 Aortic Valve Mean Gradient

Coding Instruction: Indicate the aortic valve mean gradient in mm Hg.

Target Value: N/A

Technical Specification

 Code:
 112000001398

 Code System:
 ACC NCDR

 Short Name:
 AJ_AVG

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR)

Valid Range: 5 - 50 mm[Hg]
Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14355 Aortic Valve Reintervention

Primary Indication

Operator: Equal Value: Stenosis







Section: MV Re-Intervention

Parent: In-Hospital Event Information

Element: 14360 Mitral Valve Reintervention Type

Coding Instruction: Indicate the type of mitral valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: MVReinType

Missing Data: Report
Harvested: Yes (BDS, TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacemen	t		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replac	cement		112000001875	ACC NCDR
Balloon Valvuloplasty	/		112000001469	ACC NCDR
Leaflet Clip Procedur	e		112000001778	ACC NCDR
Paravalvular Leak Cl	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: MV Re-Intervention

Parent: In-Hospital Event Information

Element: 14361 Mitral Valve Reintervention Indication

Coding Instruction: Indicate the primary indication for the reintervention. If more than one indication is present,

code the indication the operator feels has the highest significance.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: MVReintInd

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVr

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation			40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: Tricuspid Valve Re-Intervention

Parent: In-Hospital Event Information

Element: 14322 Tricuspid Valve Reintervention Type

Coding Instruction: Indicate the type of tricuspid valve re-intervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: AJ_TVReIn

Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD

Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Tricuspid Valve

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacemen	t		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replace	cement		112000001875	ACC NCDR
Balloon Valvuloplasty	/		112000001469	ACC NCDR
Leaflet Clip Procedur	e		112000001778	ACC NCDR
Paravalvular Leak Cl	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: Tricuspid Valve Re-Intervention

Parent: In-Hospital Event Information

Element: 14347 Tricuspid Valve Reintervention Primary Indication

Coding Instruction: Indicate the primary indication for the tricuspid valve re-intervention.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: AJ_TVInd
Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14312 Adjudication Event

Operator: Equal

Value: Reintervention - Tricuspid Valve
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation			40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: Tricuspid Valve Re-Intervention

Parent: In-Hospital Event Information

Element: 14383 Tricuspid Valve Regurgitation

Coding Instruction: Indicate the severity of tricuspid valve regurgitation.

Target Value: N/A

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: AJ_TR
Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14347 Tricuspid Valve Reintervention

Primary Indication

Operator: Equal

Value: Regurgitation

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR



Element: 13763



Full Specifications Data Dictionary v3.0



Section: Post-Procedure Hemoglobin

Parent: Post-Procedure Clinical Data

Hemoglobin Coding Instruction: Indicate the hemoglobin (Hgb) value in g/dL.

Target Value: The lowest value between end of current procedure and discharge

Supporting Definition: Hemoglobin

Hemoglobin (Hb or Hgb) is the iron-containing oxygen-transport metalloprotein in the red blood cells. It carries oxygen from the lungs to the rest of the body (i.e. the tissues) where it releases the oxygen to burn nutrients and provide energy. Hemoglobin concentration measurement is among the most commonly performed blood tests, usually as part of a complete blood count. If the concentration is below normal, this is called anemia. Anemias are classified by the size of red blood cells: "microcytic" if red cells are small, "macrocytic" if they are large, and "normocytic" if otherwise. Dehydration or hyperhydration can greatly influence

measured hemoglobin levels.

Source: http://s.details.loinc.org/LOINC/718-7.html?sections=Simple

Technical Specification

Code: 718-7 Code System: LOINC Short Name: PostProcHgb1 Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR.

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: PQ Precision: 4,2 Selection Type: Single Unit of Measure: g/dL Default Value: Null

> Usual Range: 5.00 - 20.00 g/dL Valid Range: 1.00 - 50.00 g/dL

Data Source: User

Parent/Child Validation Element: 14243 Hemoglobin Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 14243 Hemoglobin Not Drawn

Coding Instruction: Indicate if a post-procedure hemoglobin was not collected.

Target Value: N/A

Technical Specification

Code: 718-7 Code System: LOINC Short Name: PProcHgbND Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User







Section: Post-Procedure 12 Lead

Parent: Post-Procedure Clinical Data

Technical Specification Element: 13616 12 Lead Electrocardiogram Performed Code: 164847006 Coding Instruction: Indicate if post procedure 12 lead ECG was performed. Code System: SNOMED CT Short Name: POpEKG Target Value: Any occurrence between end of current procedure and discharge Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null

Element: 13765 12 Lead Electrocardiogram Findings

Coding Instruction: Indicate the post procedure 12 lead ECG findings. If more than one ECG is performed,

document the findings from any ECG.

Target Value: Any occurrence between end of current procedure and discharge

Vendor Instruction: Cannot select option No Significant Changes with any other option: Pathological Q Wave,

Cardiac Arrhythmia, New Left Bundle Branch Block, Pathological Q Wave, Cardiac Arrhythmia

or New Left Bundle Branch Block

Technical Specification

Code: 112000001362
Code System: ACC NCDR
Short Name: PoP_EKGChange

Missing Data: Report

Usual Range: Valid Range: Data Source: User

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13616 12 Lead Electrocardiogram

Performed

Operator: Equal Value: Yes

12 Lead Electrocardiogram Findings - 1.3.6.1.4.1.19376.1.4.1.6.5.535

Selection	Definition	Source	Code	Code System
Cardiac Arrhythmia	The patient has a new onset of an atrial or ventricula arrhythmia requiring medication or other therapy. Th includes brady or tachy arrhythmias.		698247007	SNOMED CT
No Significant Changes			112000001391	ACC NCDR
Pathological Q Wave			164918000	SNOMED CT
New Left Bundle Branch Block			100014019	ACC NCDR



Element: 10060



Full Specifications **Data Dictionary v3.0**



Section: Post-Procedure Creatinine

Parent: Post-Procedure Clinical Data

Coding Instruction: Indicate the creatinine (Cr) level mg/dL.

Target Value: The last value on discharge

Creatinine

Supporting Definition: Creatinine

Creatinine or creatine anhydride, is a breakdown product of creatine phosphate in muscle. The loss of water molecule from creatine results in the formation of creatinine. It is transferred to the kidneys by blood plasma, whereupon it is eliminated by glomerular filtration and partial tubular excretion. Creatinine is usually produced at a fairly constant rate and measuring its serum level is a simple test. A rise in blood creatinine levels is observed only with marked damage to functioning nephrons; therefore this test is not suitable for detecting early kidney disease. Creatine and creatinine are metabolized in the kidneys, muscle, liver and pancreas.

Source: http://s.details.loinc.org/LOINC/2160-0.html?sections=Simple

Technical Specification

Code: 2160-0
Code System: LOINC
Short Name: DCCreatinine
Missing Data: Report

Harvested: Yes (TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: mg/dL
Default Value: Null

Usual Range: 0.10 - 5.00 mg/dL **Valid Range:** 0.10 - 30.00 mg/dL

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----

Element: 10061 Creatinine Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 10061 Creatinine Not Drawn

Coding Instruction: Indicate if a discharge creatinine level was not drawn.

Target Value: The last value on discharge

Technical Specification

Code: 2160-0 Code System: LOINC

Short Name: DCCreatinineND

Missing Data: Report

Harvested: Yes (TAVR, TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy Procedure Type

Valid Range: Data Source: User

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Post-Procedure Highest Creatinine

Parent: Post-Procedure Clinical Data

Element: 13764 Creatinine

Coding Instruction: Indicate the post-procedure creatinine level in mg/dL. If more than one level is available, code

the peak level.

Target Value: The highest value between end of current procedure and discharge

Supporting Definition: Creatinine

Creatinine or creatine anhydride, is a breakdown product of creatine phosphate in muscle. The loss of water molecule from creatine results in the formation of creatinine. It is transferred to the kidneys by blood plasma, whereupon it is eliminated by glomerular filtration and partial tubular excretion. Creatinine is usually produced at a fairly constant rate and measuring its serum level is a simple test. A rise in blood creatinine levels is observed only with marked damage to functioning nephrons; therefore this test is not suitable for detecting early kidney disease. Creatine and creatinine are metabolized in the kidneys, muscle, liver and pancreas.

Source: http://s.details.loinc.org/LOINC/2160-0.html?sections=Simple

Technical Specification

Code: 2160-0
Code System: LOINC
Short Name: PoProc_Creat
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Usual Range: 0.10 - 9.00 mg/dL **Valid Range:** 0.10 - 30.00 mg/dL

Data Source: User

Parent/Child Validation

Element: 14293 Highest Creatinine Not Drawn

Operator: Equal

Value: No (or Not Answered)

Element: 14293 Highest Creatinine Not Drawn

Coding Instruction: Indicate if the highest creatinine level was not drawn.

Target Value: N/A

Technical Specification

Code: 2160-0
Code System: LOINC
Short Name: HighCrea_ND
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13592 Echocardiogram Performed

Coding Instruction: Indicate the type of echo performed prior to discharge.

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 40701008
Code System: SNOMED CT
Short Name: POpTTEch
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13645 Echocardiogram Not Performed

Operator: Equal

Value: No (or Not Answered)

Echocardiogram Type - 1.3.6.1.4.1.19376.1.4.1.6.5.526

 Selection
 Definition
 Source
 Code System

 Transesophageal Echocardiogram (TEE)
 105376000
 SNOMED CT

 Transthoracic Echo (TTE)
 433236007
 SNOMED CT

Element: 13645 Echocardiogram Not Performed

 $\begin{tabular}{ll} \textbf{Coding Instruction:} & Indicate if an echocardiogram was not performed. \\ \end{tabular}$

Target Value: N/A

Technical Specification

Code: 40701008

Code System: SNOMED CT
Short Name: EchoND

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13493 Echocardiogram Date

Coding Instruction: Indicate the date the echocardiogram was performed.

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 40701008
Code System: SNOMED CT
Short Name: POpTTEchDate
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13495 Aortic Valve Area

Coding Instruction: Indicate the smallest aortic valve area (in cm2).

Target Value: The lowest value between end of current procedure and discharge

Technical Specification Code: 112000001280

Code System: ACC NCDR
Short Name: PP_AVArea
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No

Data Type: PQ
Precision: 3,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 0.20 - 4.00 cm2 **Valid Range:** 0.05 - 5.00 cm2

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13675 Aortic Valve Mean Gradient

Coding Instruction: Indicate the mean gradient (in mm Hg) across the aortic valve.

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 112000001398
Code System: ACC NCDR
Short Name: PP_AVMeanGradient

Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Lisual Range: 5 - 50 mm[

Usual Range: 5 - 50 mm[Hg]
Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13526 Aortic Valve Regurgitation

Coding Instruction: Indicate the severity of aortic valve regurgitation.

If mild-moderate is documented, code as mild.

If moderate-severe is documented, code as moderate.

Target Value: The last value between end of current procedure and next procedure or discharge

Technical Specification

Code: 60234000
Code System: SNOMED CT
Short Name: PP_AR
Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND ----

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13494 Mitral Regurgitation

Coding Instruction: Indicate the severity of mitral valve regurgitation.

If mild-moderate is documented, code as mild.

Target Value: The last value between end of current procedure and next procedure or discharge

Technical Specification

Code: 48724000
Code System: SNOMED CT
Short Name: PP_MR
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.728

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Moderate-Severe			1000142345	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13677 Tricuspid Valve Regurgitation

Coding Instruction: Indicate the severity of tricuspid valve regurgitation.

If mild-moderate is documented, code as mild.

If moderate-severe is documented, code as moderate.

Target Value: The last value between end of current procedure and next procedure or discharge

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: PP_TR
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13779 Effective Regurgitant Orifice Area

Coding Instruction: Indicate the effective regurgitant orifice area (EROA), in cm2.

Target Value: The highest value between end of current procedure and next procedure or discharge

Technical Specification

Code: 112000001437
Code System: ACC NCDR
Short Name: PP_MV_EOA
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 2,1
Selection Type: Single
Unit of Measure: cm2
Default Value: Null
Usual Range: 0.1 - 5.0 cm2
Valid Range: 0.1 - 5.0 cm2
Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND ------

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal
Value: TMVr

Element: 13769 Effective Regurgitant Orifice Area Method of Assessment

Coding Instruction: Indicate the method used to assess the effective regurgitant orifice area. If multiple methods

are available, code the 3D planimetry method first, then PISA.

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 112000001437
Code System: ACC NCDR
Short Name: PP_MV_EOA_MOA

Missing Data: Report

Is Identifier: No

Harvested: Yes (TMVR, TMVrpr)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13779 Effective Regurgitant Orifice Area

Operator:

Value: Any Value

Effective Requiritant Orifice Area Method of Assessment - 1 3 6 1 4 1 19376 1 4 1 6 5 547

Selection	Definition	Source	Code	Code System	
3D Planimetry			112000001438	ACC NCDR	
Proximal Isovelocity Surface Area			112000001439	ACC NCDR	
Quantitative Doppler			112000001440	ACC NCDR	
Other			100000351	ACC NCDR	







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13770 Mitral Valve Mean Gradient

Coding Instruction: Indicate the mean gradient (in mm Hg) across the mitral valve.

Target Value: The highest value between end of current procedure and discharge

Supporting Definition: Mitral Valve Mean Gradient

The average gradient across the mitral valve occurring during the entire systole.

Source: Baumgartner, H. et. al (2009). Echocardiographic assessment of valve stenosis:

EAE/ASE

recommendations for clinical practice.

Technical Specification

Code: 112000001191
Code System: ACC NCDR
Short Name: PP_MVMeanGradient

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Identifier: No

Usual Range: 5 - 50 mm[Hg]
Valid Range: 0 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

value: INVR

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal
Value: TMVr

Element: 13771 Mitral Valve Area

Coding Instruction: Indicate the smallest mitral valve area in centimeters squared.

Target Value: The lowest value between end of current procedure and discharge

Supporting Definition: Mitral Valve Area

Measurement of mitral valve area.

Source:

Technical Specification

Code: 251012002
Code System: SNOMED CT
Short Name: PP_MVArea
Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 3.00 - 6.00 cm2 **Valid Range:** 0.05 - 12.00 cm2

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 13772 Left Ventricular Outflow Tract Peak Velocity

Coding Instruction: Indicate the left ventricular outflow tract peak velocity in m/sec.

Target Value: The highest value between end of current procedure and discharge

Technical Specification

 Code:
 112000002047

 Code System:
 ACC NCDR

 Short Name:
 PP_LVOT

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: m/sec
Default Value: Null

Usual Range: 0.5 - 5.0 m/sec Valid Range: 0.1 - 10.0 m/sec

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

Element: 13774 Systolic Anterior Motion Present

 $\begin{tabular}{ll} \textbf{Coding Instruction:} & Indicate if systolic anterior motion of the mitral valve was present. \\ \end{tabular}$

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 112000001481
Code System: ACC NCDR
Short Name: PP_SAM
Missing Data: Report
Harvested: Yes (TMVR)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Data Source: User

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 14507 Tricuspid Valve Diastolic Gradient

Coding Instruction: Indicate the post-procedure tricuspid valve diastolic gradient in mm Hg. This can also be called

the TV inflow gradient.

Target Value: The highest value between end of current procedure and next procedure or discharge

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: PP_TVDGrad
Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 1 - 15 mm[Hg]
Valid Range: 1 - 50 mm[Hg]
Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND

Element: 14508 Tricuspid Valve Diastolic Gradient

Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14508 Tricuspid Valve Diastolic Gradient Not Documented

Coding Instruction: Indicate if the tricuspid valve diastolic gradient was not documented post-procedure.

Target Value: N/A

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: PP_TVDGradND

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: Yes

Is Followup
Element:
Data Type: BL
Precision:

Selection Type: Single Unit of Measure: Default Value: Null

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND ----

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 14294 Tricuspid Valve Annulus Size

Coding Instruction: Indicate the tricuspid valve annulus size in mm. Documentation using end-diastolic, 4 chamber view is preferred.

Target Value: The lowest value between end of current procedure and next procedure or discharge

Technical Specification

Code: 112000001513 Code System: ACC NCDR Short Name: PP_TVAnnulus

Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No Is Base Element: Yes

Is Followup No Data Type: PQ Precision: 2,0

Selection Type: Single Unit of Measure: mm Default Value: Null Usual Range: 15 - 60 mm Valid Range: 1 - 80 mm Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND

Element: 14495 Tricuspid Valve Annulus Size Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14495 Tricuspid Valve Annulus Size Not Documented

Coding Instruction: Indicate if the tricuspid valve annulus size was not documented.

Target Value: N/A

Technical Specification

Code: 112000001513 Code System: ACC NCDR Short Name: PP_TVAnnulusND

Missing Data: Report Harvested: Yes (TTVP) Is Identifier: No

Is Base Element: Yes Is Followup No Data Type: BL Precision:

Selection Type: Single Unit of Measure: Default Value: Null

> **Usual Range:** Valid Range: Data Source: User

> > Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----

Element: 14273 Transcatheter Valve Therapy

Procedure Type Operator: Equal







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 14295 End Diastolic Mid Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic mid right ventricular (RV) diameter, using the 4 chamber view (in cm).

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 112000001514
Code System: ACC NCDR
Short Name: PP_MidRVDia
Missing Data: Report

Selection Type: Single
Unit of Measure: cm
Default Value: Null
Usual Range: 1.0 - 7.0 cm
Valid Range: 0.1 - 9.9 cm
Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----

Element: 14496 End Diastolic Mid Right Ventricle

Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14496 End Diastolic Mid Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the end-diastolic mid right ventricular (RV) diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001514
Code System: ACC NCDR
Short Name: PP_MidRVDiaND

Missing Data: Report Harvested: Yes (TTVP) Is Identifier: No

Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:

Usual Range: Valid Range: Data Source: User

Default Value: Null

Parent/Child Validation

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 14296 End Diastolic Basal Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic basal right ventricular (RV) diameter, using the 4 chamber view (in

cm).

Target Value: Any occurrence between end of current procedure and discharge

Technical Specification

Code: 112000001515
Code System: ACC NCDR

Short Name: PP_BasalRVDia Missing Data: Report

Harvested: Yes (TTVP)
Is Identifier: No

 Is Base Element:
 Yes

 Is Followup Element:
 No

 Data Type:
 PQ

 Precision:
 2,1

 Selection Type:
 Single

Unit of Measure: cm
Default Value: Null
Usual Range: 1.0 - 7.0 cm
Valid Range: 0.1 - 9.9 cm
Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal
Value: Tricuspid Valve Procedure

----- AND -----

Element: 14497 End Diastolic Basal Right

Ventricle Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14497 End Diastolic Basal Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the end diastolic basal right ventricular (RV) diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001515
Code System: ACC NCDR
Short Name: PP BasalRVDiaND

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type
Operator: Equal







Section: Post-Procedure Echocardiogram Findings

Parent: Post-Procedure

Element: 14297 Right Ventricular Systolic Pressure

Coding Instruction: Indicate the right ventricular systolic pressure in mm Hg recorded post procedure. Note: If

more than one RVSP documented, code the highest value.

Target Value: The highest value between end of current procedure and next procedure or discharge

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: PP_RVSP
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: PQ
Precision: 3,0

Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 15 - 30 mm[Hg] Valid Range: 1 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13592 Echocardiogram Performed **Operator:** Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14498 Right Ventricular Systolic

Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14498 Right Ventricular Systolic Pressure Not Documented

Coding Instruction: Indicate if the right ventricular systolic pressure was not documented.

Target Value: N/A

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: PP_RVSYSND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Base Element: Yes

Parent/Child Validation

Element: 13592 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13592 Echocardiogram Performed

Operator: Equal

Element: 14273 Transcatheter Valve Therapy Procedure Type

Operator: Equal







Section: Post-Procedure AV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14503 Paravalvular Aortic Regurgitation

Coding Instruction: Indicate the severity of paravalvular aortic valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: PP_ParaAR
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13526 Aortic Valve Regurgitation

Operator: Equal
Value: Trace/Trivial
----- AND

Element: 14524 Paravalvular Aortic Regurgitation

Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure AV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14524 Paravalvular Aortic Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular aortic valve regurgitation was not documented post-

procedure

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: PP_ParaARND
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13526 Aortic Valve Regurgitation

Operator: Equal
Value: Trace/Trivial
----- AND

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: Post-Procedure AV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14499 Central Aortic Regurgitation

Coding Instruction: Indicate the severity of central aortic valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: PP_CentralAR
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13526 Aortic Valve Regurgitation

Operator: Equal
Value: Trace/Trivial

AND ----
Element: 14487 Central Aortic Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure AV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14487 Central Aortic Regurgitation Not Documented

Coding Instruction: Indicate if central aortic valve regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: PP_CentralARND

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13526 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13526 Aortic Valve Regurgitation

Operator: Equal

Value: Trace/Trivial

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TAVR







Section: Post-Procedure MV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 13766 Paravalvular Mitral Regurgitation

Coding Instruction: Indicate the severity of paravalvular mitral valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 112000001428 Code System: ACC NCDR Short Name: PP_ParaMR Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No Is Base Element: Yes Is Followup Element: No Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Moderate

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

----- AND -----

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

----- AND -----Element: 14525 Paravalvular Mitral Regurgitation

Not Documented

Operator: Equal

Value: No (or Not Answered)

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Taire regulgitation coverity 4 holomatinostics				
Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure MV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14525 Paravalvular Mitral Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular mitral regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: PP_ParaMRND
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Operator: Equal Value: TMVR

Element: 13494 Mitral Regurgitation

Element. 15494 William Reguly

Operator: Equal Value: Mild

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Post-Procedure MV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 13767 Central Mitral Regurgitation

Coding Instruction: Indicate the severity of central mitral valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

 Code:
 112000001433

 Code System:
 ACC NCDR

 Short Name:
 PP_CentralMR

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR)

Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13494 Mitral Regurgitation

Operator: Equal
Value: Moderate

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal Value: TMVR

AND -----

Element: 14488 Central Mitral Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Turio regularita de la companya de l				
Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure MV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14488 Central Mitral Regurgitation Not Documented

Coding Instruction: Indicate if central mitral regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: PP_CentralMRND

Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Data Source: User

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Mild

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13494 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13494 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Post-Procedure TV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14505 Paravalvular Tricupsid Regurgitation

Coding Instruction: Indicate the severity of paravalvular tricuspid valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: PP_ParaTR
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13677 Tricuspid Valve Regurgitation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND ----

Element: 14526 Paravalvular Tricupsid

Regurgitation Not Documented

Operator: Equal

Value: No (or Not Answered)

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure TV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14526 Paravalvular Tricupsid Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular tricuspid regurgitation was not documented post-

procedure

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: PP_ParaTRND
Missing Data: Report

Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal

Value: Moderate

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Trace/Trivial







Section: Post-Procedure TV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14501 Central Tricupsid Regurgitation

Coding Instruction: Indicate the severity of central tricuspid valve regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value between end of current procedure and discharge

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: PP_CentraITR
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13677 Tricuspid Valve Regurgitation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----Element: 14489 Central Tricupsid Regurgitation

Not Documented

Operator: Equal

Value: No (or Not Answered)

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Post-Procedure TV Regurgitation

Parent: Post-Procedure Echocardiogram Findings

Element: 14489 Central Tricupsid Regurgitation Not Documented

Coding Instruction: Indicate if central tricuspid valve regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: PP_CentralTRND

Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14273 Transcatheter Valve Therapy

Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal

Value: Mild

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13677 Tricuspid Valve Regurgitation

Operator: Equal Value: Trace/Trivial







Data Source: User

Data Source: User

Section: Discharge	Parent: Root		
Element: 10100	Discharge Date	Technic	al Specification
Coding Instruction:	Indicate the date on which the patient was discharged from your facility.	Code: Code System:	1000142457 ACC NCDR
_	The value on discharge	Short Name: Missing Data:	
Vendor Instruction:	Discharge Date (10100) must be Greater than or Equal to 01/01/2021 Discharge Date (10100) and Arrival Date and Time (3001) must not overlap on multiple episodes		Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
		Is Identifier:	No
		Is Base Element: Is Followup Element:	
		Data Type: Precision:	DT
		Selection Type: Unit of Measure:	Single
		Default Value: Usual Range: Valid Range:	Null

Element: 10070	Discharge Provider's Last Name	Technical Specification
Coding Instruction:	Indicate the last name of the discharge provider.	Code: 1000142453 Code System: ACC NCDR
	Note(s): If the name exceeds 50 characters, enter the first 50 characters only.	Short Name: DCLName Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr,
	The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical record.	TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: Data Type: LN
Target Value:	The value on discharge	Precision: 50 Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Element: 10071	Discharge Provider's First Name	Technical Specification
		Code: 1000142453
Coding Instruction:	Indicate the first name of the discharge provider.	Code System: ACC NCDR
	Note(s):	Short Name: DCFName
	If the name exceeds 50 characters, enter the first 50 characters only.	Missing Data: Report
		Harvested: Yes (TAVR, TMVR, TMVrpr,
	The completion of this data element is voluntary and at the discretion of your facility. NCDR will	TTVP)
	use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical record.	Is Identifier: No
		Is Base Element: Yes
		Is Followup No
		Element: NO
	record.	Data Type: FN
Target Value:	The value on discharge	Precision: 50
		Selection Type: Single
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:







Data Source: User

Technical Specification

Section: Discharge	Parent: Root	
Element: 10072	Discharge Provider's Middle Name	Technical Specification
Coding Instruction:	Indicate the middle name of the discharge provider.	Code: 1000142453 Code System: ACC NCDR
	Note(s): It is acceptable to specify the middle initial.	Short Name: DCMName Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr,
	If there is no middle name given, leave field blank.	TTVP) Is Identifier: No
	If there are multiple middle names, enter all of the middle names sequentially. If the name exceeds 50 characters, enter the first 50 letters only.	Is Base Element: Yes Is Followup Element: No
	The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality	Data Type: MN Precision: 50 Selection Type: Single
	improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical record.	Unit of Measure: Default Value: Null Usual Range:
Target Value:	The value on discharge	Valid Range:

Element: 10073	Discharge Provider's NPI	Technical Specification
	Indicate the National Provider Identifier (NPI) of the provider that discharged the patient. NPI's, assigned by the Centers for Medicare and Medicaid Services (CMS), are used to uniquely identify physicians for Medicare billing purposes. Note(s): The completion of this data element is voluntary and at the discretion of your facility. NCDR will use the data to provide reporting at the physician level, which may assist physicians with demonstrating value based care as well as support your facility's engagement in quality improvement efforts. If completed, NCDR will defer to your facility's determination of assigning Admitting, Attending, and Discharging Provider roles, as supported by the patient medical	Code: 1000142453 Code System: ACC NCDR Short Name: DCNPI Missing Data: Report Harvested: Yes (TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element:
Target Value:	record. The value on discharge	Data Type: NUM Precision: 10 Selection Type: Single
		Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Discharge Status	rediffical opeomodition
•	Code: 75527-2
Indicate whether the patient was alive or deceased at discharge.	Code System: LOINC
The value on discharge	Short Name: DCStatus
	Missing Data: Illegal
	Harvested: Yes (BDS, TAVR, TMVR,
	TMVrpr, TTVP)
	Is Identifier: No
	Is Base Element: Yes
	Is Followup No
	Element: NO
	Data Type: CD
	Precision:
	Selection Type: Single
	Unit of Measure:
	Default Value: Null
	Usual Range:
	Valid Range:
	Data Source: User
	Indicate whether the patient was alive or deceased at discharge. The value on discharge

Discharge Life Status -	1.3.6.1.4.1.19376.1.4.1.6.5.42
Discharge Life Status -	1.3.0.1.4.1.13370.1.4.1.0.3.42

Discharge Status

Selection	Definition	Source	Code	Code System
Alive			438949009	SNOMED CT
Deceased			20	HL7 Discharge disposition

Element: 10105







Section: Discharge Parent: Root

Element: 10116 Cardiac Rehabilitation Referral

Coding Instruction: Indicate if the patient has been referred to an outpatient cardiac rehab program with face to

discharge. The referral may be to a traditional outpatient cardiac rehab program with face-toface interactions and training sessions or may include other novel delivery options.

lace interactions and training sessions of may include other novel delivery options

Target Value: The value on discharge
Supporting Definition: Cardiac Rehabilitation Referral

1. Documented communication between the healthcare provider and the patient to recommend

an outpatient CR program

AND

2A. Official referral order is sent to outpatient CR program

OF

2B. Documentation of patient refusal to justify why patient information was not sent to the CR

program

Source: Source: Thomas, R.J., et al. 2018 ACC/AHA Clinical Performance and Quality Measures for Cardiac Rehabilitation: A Report of the American College of Cardiology/American

Heart Association Task Force on Performance Measures. Journal of the American College of Cardiology, Vol 71, Issue 16, April 2018, pages 1814-1837

Technical Specification

Code: 100014067
Code System: ACC NCDR
Short Name: DC_CardRehab

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Data Type: CD
Precision:

Selection Type: Single Unit of Measure: Default Value: Null

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 10105 Discharge Status

Operator: Equal Value: Alive

Cardiac Rehab - 1.3.6.1.4.1.19376.1.4.1.6.5.334

Selection	Definition	Source	Code	Code System
No - Reason Not Documented			100014064	ACC NCDR
No - Medical Reason Documented	Patient deemed by a medical provider to have a medically unstable, life-threatening condition or has other cognitive or physical impairments that preclude CR participation.	Thomas, R.J., et al. 2018 ACC/AHA Clinical Performance and Quality Measures for Cardiac Rehabilitation: A Report of the American College of Cardiology/American Heart Association Task Force on Performance Measures. Journal of the American College of Cardiology, Vol 71, Issue 16, April 2018, pages 1814-1837	100014066	ACC NCDR
No - Health Care System Reason Documented	Patient is discharged to a nursing care or long-term care facility, or patient lacks medical coverage for CR.	Thomas, R.J., et al. 2018 ACC/AHA Clinical Performance and Quality Measures for Cardiac Rehabilitation: A Report of the American College of Cardiology/American Heart Association Task Force on Performance Measures. Journal of the American College of Cardiology, Vol 71, Issue 16, April 2018, pages 1814-1837	100014065	ACC NCDR
No - Patient - Oriented Reason	No traditional CR program available to the patient, within 60 min [travel time] from the patient's home, or patient does not have access to an alternative model of CR delivery that meets all criteria for a CR program.	n Thomas, R.J., et al. 2018 ACC/AHA Clinical Performance and Quality Measures for Cardiac Rehabilitation: A Report of the American College of Cardiology/American Heart Association Task Force on Performance Measures. Journal of the American College of Cardiology, Vol 71, Issue 16, April 2018, pages 1814-1837	112000000520	ACC NCDR
Yes			100013072	ACC NCDR







Section: Discharge Parent: Root

Element: 10110 Discharge Location

Coding Instruction: Indicate the location to which the patient was discharged.

Target Value: The value on discharge

Technical Specification

Code: 75528-0
Code System: LOINC
Short Name: DCLocation
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 10105 Discharge Status

Operator: Equal Value: Alive

Discharge Location - 1.3.6.1.4.1.19376.1.4.1.6.5.41

Selection	Definition	Source	Code	Code System
Home			01	HL7 Discharge disposition
Skilled Nursing Facility	Skilled nursing facilities (SNF) are typically sub-acute programs used for longer anticipated length of stay.		03	HL7 Discharge disposition
	Note: Sometimes SNFs may have acute rehabilitation beds within their facility. If the patient is discharged to a SNF for acute rehab (requiring a higher level of carcode "extended care/TCU/rehab".	0		
Extended Care/TCU/Rehab	An extended care unit, transitional care unit or rehab unit typically provides a high level of intensive therap as well as specialized nursing and physician care. The discharge setting may also be called subacute care clong term acute care (LTACH).	/ nis	62	HL7 Discharge disposition
Other Acute Care Hospital			02	HL7 Discharge disposition
Left Against Medical Advice (AMA)	The patient was discharged or eloped against medica advice.	ıl	07	HL7 Discharge disposition
Other Discharge Location			100001249	ACC NCDR

Element: 10115 Hospice Care

Coding Instruction: Indicate if the patient was discharged to hospice care.

Target Value: The value on discharge

Technical Specification

Code: 385763009
Code System: SNOMED CT
Short Name: DCHospice
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element: No
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Is Identifier: No

Parent/Child Validation 10105 Discharge Status

Element: 10105 Operator: Equal Value: Alive







Section: Discharge Parent: Root

Element: 10120 Death During the Procedure

Coding Instruction: Indicate if the patient expired during the procedure.

Note(s): Make sure to only capture 'death during the procedure' in the procedure appropriate

registry.

For example, if the patient had a CathPCI procedure and a TVT procedure in the same episode of care (hospitalization) but different cath lab visits and the death occurred during the TVT procedure, code 'Yes' only in the TVT Registry and not the CathPCI Registry. If the CathPCI procedure and TVT procedure occurred during the same cath lab visit then code 'Yes' in both

registries.

Target Value: Any occurrence on discharge

Technical Specification

Code: 100000923
Code System: ACC NCDR
Short Name: DeathProcedure

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single

Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 10105 Discharge Status

Operator: Equal Value: Deceased



Element: 10125



Full Specifications **Data Dictionary v3.0**



Section: Discharge Parent: Root

Coding Instruction: Indicate the primary cause of death, i.e. the first significant abnormal event which ultimately led

to death

Cause of Death

Target Value: The value on time of death

Technical Specification

Code: 184305005
Code System: SNOMED CT
Short Name: DeathCause
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: OD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 10105 Discharge Status

Operator: Equal Value: Deceased

Cause of Death Clinical Finding - 1.3.6.1.4.1.19376.1.4.1.6.5.88

Selection	Definition	Source	Code	Code System
Acute myocardial infarction	Death by any cardiovascular mechanism (e.g., arrhythmia, sudden death, heart failure, stroke, pulmonary embolus, peripheral arterial disease) within 30 days after an acute myocardial infarction, related the immediate consequences of the MI, such as progressive HF or recalcitrant arrhythmia. There may be other assessable (attributable) mechanisms of cardiovascular death during this time period, but for simplicity, if the cardiovascular death occurs <=30 days of an acute myocardial infarction, it will be		100000960	ACC NCDR
	considered a death due to myocardial infarction.			
Sudden cardiac death	Death that occurs unexpectedly, and not within 30 days of an acute MI.		100000978	ACC NCDR
Heart failure	Death associated with clinically worsening symptoms and/or signs of heart failure.		100000964	ACC NCDR
Stroke	Death after a stroke that is either a direct consequence of the stroke or a complication of the stroke.	е	100000977	ACC NCDR
Cardiovascular procedure	Death caused by the immediate complication(s) of a cardiovascular procedure.		100000962	ACC NCDR
Cardiovascular hemorrhage	Death related to hemorrhage such as a non-stroke intracranial hemorrhage, non-procedural or non-traumatic vascular rupture (e.g., aortic aneurysm), or hemorrhage causing cardiac tamponade.		100000961	ACC NCDR
Other cardiovascular reason	Cardiovascular death not included in the above categories but with a specific, known cause (e.g., pulmonary embolism, peripheral arterial disease).		100000972	ACC NCDR
Pulmonary	Non-cardiovascular death attributable to disease of th lungs (excludes malignancy).	е	100000975	ACC NCDR
Renal	Non-cardiovascular death attributable to renal failure.		100000976	ACC NCDR
Gastrointestinal	Non-cardiovascular death attributable to disease of th esophagus, stomach, or intestines (excludes malignancy).	е	100000963	ACC NCDR
Hepatobiliary	Non-cardiovascular death attributable to disease of th liver, gall bladder, or biliary ducts (exclude malignancy		100000966	ACC NCDR
Pancreatic	Non-cardiovascular death attributable to disease of th pancreas (excludes malignancy).	е	100000974	ACC NCDR
Infection	Non-cardiovascular death attributable to an infectious disease.		100000967	ACC NCDR
Inflammatory/Immunologic	Non-cardiovascular death attributable to an inflammatory or immunologic disease process.		100000968	ACC NCDR
Hemorrhage	Non-cardiovascular death attributable to bleeding that not considered cardiovascular hemorrhage or stroke per this classification.	is	100000965	ACC NCDR
Non-cardiovascular procedur or surgery	e Death caused by the immediate complication(s) of a non-cardiovascular procedure or surgery.		100000971	ACC NCDR







Section: Discharge	Parent: F	Root	
Trauma	Non-cardiovascular death attributable to trauma.	100000980	ACC NCDR
Suicide	Non-cardiovascular death attributable to suicide.	100000979	ACC NCDR
Neurological	Non-cardiovascular death attributable to disease of the nervous system (excludes malignancy).	100000970	ACC NCDR
Malignancy	Non-cardiovascular death attributable to malignancy.	100000969	ACC NCDR
Other non-cardiovascular reason	Non-cardiovascular death attributable to a cause other than those listed in this classification (specify organ system).	100000973	ACC NCDR

Technical Specification Element: 9275 Packed Red Blood Cell Transfusion Code: 71493000 Coding Instruction: Indicate if there was a transfusion(s) of packed red blood cells. Code System: SNOMED CT Short Name: PostTransfusion Target Value: Any occurrence between start of procedure and until next procedure or discharge Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range:

Element: 13670 Packed Red Blood Cell Units Transfused

Coding Instruction: Indicate the total number of units transfused of packed red blood cells.

Target Value: The total value between start of first procedure until discharge

Technical Specification

Code: 100014031
Code System: ACC NCDR
Short Name: DC_RBCUnit
Missing Data: Report

Valid Range: Data Source: User

 $\textbf{Harvested:} \ \ \text{Yes (BDS, TAVR, TMVR,} \\$

TMVrpr, TTVP) **Is Identifier:** No

Is Base Element: Yes
Is Followup
Element:
Data Type: NUM
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 9275 Packed Red Blood Cell Transfusion

Operator: Equal Value: Yes



Element: 10200



Full Specifications Data Dictionary v3.0



Section: Discharge Medications

Parent: Discharge

Discharge Medication Code

Coding Instruction: Indicate the assigned identification number associated with the medications the patient was

prescribed upon discharge.

Note(s):

Discharge medications not required for patients who expired, discharged to "Other acute care

hospital", "Left against medical advice (AMA)" or are receiving Hospice Care.

The medication(s) collected in this field are controlled by the Medication Master file. This file is maintained by the NCDR and will be made available on the internet for downloading and importing/updating into your application. Each medication in the Medication Master file is assigned to a value set. The value set is used to separate procedural medications from medications prescribed at discharge. The separation of these medications is depicted on the

data collection form.

Target Value: N/A

Technical Specification

Code: 100013057 Code System: ACC NCDR Short Name: DC_MedID Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup No Element: Data Type: CD Precision:

Selection Type: Single (Dynamic List)

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 10110 Discharge Location

Operator: Equal Value: Home

Element: 10110 Discharge Location

Operator: Equal

Value: Extended Care/TCU/Rehab Element: 10110 Discharge Location Operator: Equal

Value: Other Discharge Location Element: 10110 Discharge Location

Operator: Equal

Value: Skilled Nursing Facility AND

Element: 10115 Hospice Care

Operator: Equal Value: No

Element: 10105 Discharge Status

----- AND -----

Operator: Equal Value: Alive

Discharge Medication - 1.3.6.1.4.1.19376.1.4.1.6.5.165

Selection	Definition	Source	Code	Code System
Angiotensin Converting			41549009	SNOMED CT
Enzyme Inhibitor				
Aldosterone Antagonist			372603003	SNOMED CT
Direct thrombin inhibitor			414010005	SNOMED CT
Warfarin			11289	RxNorm
Aspirin			1191	RxNorm
Angiotensin II Receptor Blo	ocker		372913009	SNOMED CT
Beta Blocker			33252009	SNOMED CT
Diuretics Not Otherwise			112000001417	ACC NCDR
Specified				
Loop Diuretics			29051009	SNOMED CT
Thiazides			372747003	SNOMED CT
Direct Factor Xa Inhibitor			112000000696	ACC NCDR
P2Y12 Antagonist			112000001003	ACC NCDR







Section: Discharge Medications

Parent: Discharge

Element: 10205 Discharge Medication Prescribed

Coding Instruction: Indicate if the medication was prescribed, not prescribed, or was not prescribed for either a

medical or patient reason.

Note(s):

Discharge medications do not need to be recorded for patients who were discharged to "Other

acute care hospital", "Left against medical advice (AMA)" or are receiving Hospice Care is

'Yes'

Target Value: The value on discharge

Vendor Instruction: When Discharge Medication Code (10200) is selected Discharge Medications Prescribed

(10205) cannot be Null

Technical Specification

Code: 432102000
Code System: SNOMED CT
Short Name: DC_MedAdmin
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: Yes
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Is Identifier: No

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 10200 Discharge Medication Code

Operator:

Value: Any Value

Discharge Medication Administration - 1.3.6.1.4.1.19376.1.4.1.6.5.86

Selection	Definition	Source	Code	Code System
Yes - Prescribed			100001247	ACC NCDR
Not Prescribed - No Reason	n		100001048	ACC NCDR
Not Prescribed - Medical Reason			100001034	ACC NCDR
Not Prescribed - Patient			100001071	ACC NCDR

Element: 14576 Loop Diuretic Dose

Coding Instruction: Specify the total daily dose of the loop diuretic that was prescribed to the patient at discharge.

Target Value: The value on discharge

Technical Specification

Code: 112000001975 Code System: ACC NCDR

Short Name: DischMed_LoopDiureticDose

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: No
Precision: 3,0
Selection Type: Single
Unit of Measure: mg
Default Value: Null
Usual Range: 1 - 40 mg
Valid Range: 1 - 300 mg

Data Source: User

Parent/Child Validation

Element: 10200 Discharge Medication Code

Operator: Equal

Value: Loop Diuretics

AND -----
Element: 10205 Discharge Medication Prescribed

Operator: Equal

Value: Yes - Prescribed







Section: Follow Up	Parent: Root		
lement: 11000	Follow-Up Assessment Date	Technic	al Specification
0 - 11 1 1 1 1	In directs the data of the fallowing accessment was a softeness d	Code:	1000142364
Coding instruction:	Indicate the date of the follow-up assessment was performed.	Code System:	
Target Value:	The value on Follow-up		F_AssessmentDate
Vendor Instruction:	Follow-Up Assessment Date (11000) must be Greater than or Equal to 01/01/2021	Missing Data:	•
	1	Harvested:	Yes (BDS, TAVR, TMVR
	Follow-Up Assessment Date (11000) must be Greater than or Equal to Follow-Up Reference	Is Identifier:	TMVrpr, TTVP)
	Episode Arrival Date and Time (11002)	Is Base Element:	
	A Follow-up Assessment Date may only be entered/selected once	Is Followup	
		Element:	Yes
	Follow-Up Assessment Date (11000) must be Greater than Follow-Up Reference Procedure	Data Type:	DT
	Start Date and Time (11001)	Precision:	
		Selection Type:	Single
		Unit of Measure:	
		Default Value:	Null
		Usual Range:	
		Valid Range:	
		Data Source:	User
		Tankais	al Cassification
lement: 10999	Follow-Up Unique Key		al Specification
Coding Instruction:	Indicate the unique key associated with each patient follow-up record as assigned by the	Code System:	
	EMR/EHR or your software application.	1	FollowUpKey
Target Value:	N/A	Missing Data:	
· 3 · · · · · · · · ·		_	Yes (BDS, TAVR, TMVR)
			TMVrpr, TTVP)
		Is Identifier:	Yes
		Is Base Element:	
		Is Followup	Yes
		Element:	165
		Data Type:	
		Precision:	
		Selection Type:	Single
		Unit of Measure:	NI. II
		Default Value:	Null
		Usual Range:	
		Valid Range: Data Source:	Automatia
		Data Source:	Automatic
Element: 11001	Follow-Up Reference Procedure Start Date and Time	Technic	al Specification
	·		1000142372
Cadina Instruction.	Indicate the reference procedure start date and time on the follow-up assessment date.	1	
Coding instruction:	indicate the reference procedure start date and time on the follow-up assessment date.	Code System:	ACC NCDR

		.000.120.2
Indicate the reference procedure start date and time on the follow-up assessment date.	Code System:	ACC NCDR
The value on Follow-up	Short Name:	RefProcStartDateTime
	Missing Data:	Illegal
	Harvested:	Yes (BDS, TAVR, TMVR,
		TMVrpr, TTVP)
	Is Identifier:	No
	Is Followup	Yes
	Element:	103
	Data Type:	TS
	Precision:	
	Selection Type:	Single
	Unit of Measure:	
	Default Value:	Null
	Usual Range:	
	Valid Range:	
	Data Source:	User
	Indicate the reference procedure start date and time on the follow-up assessment date. The value on Follow-up	The value on Follow-up Short Name: Missing Data:







Valid Range: Data Source: User

Data Source: User

Technical Specification
reclinical opecinication
e reference Code: 1000142436 Code System: ACC NCDR Short Name: RefArrivalDateTime
Missing Data: Illegal Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
Is Identifier: No Is Base Element: No
Is Followup Element: Pata Type: TS
Precision: Selection Type: Single
Unit of Measure: Default Value: Null Usual Range:
•

Element: 13705	Transcatheter Valve Therapy Reference Procedure Type	Technical Specification
		Code: 112000001167
Coding Instruction:	Indicate the procedure type performed at the reference procedure start date/time.	Code System: ACC NCDR
Target Value:	The value on Follow-up	Short Name: F_RefProType
Vandar Instruction	When Transcatheter Valve Therapy Reference Procedure Type (13705) is Equal to	Missing Data: Illegal
vendor mstruction.	(TMVr,TMVR,Tricuspid Valve Procedure) then Follow-Up Medications Code (11990) must be Equal to (Aldosterone Antagonist,Angiotensin Converting Enzyme Inhibitor,Angiotensin II	Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
	Receptor Blocker, Beta Blocker, Diuretics Not Otherwise Specified, Loop Diuretics, Thiazides)	Is Identifier: No
		Is Base Element: No
		Is Followup Yes
		Data Type: CD
		Precision:
		Selection Type: Multiple
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:

Transcatheter Valve Therapy Procedure - 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selection	Definition	Source	Code	Code System
TAVR	Transcatheter aortic valve replacement		41873006	SNOMED CT
TMVr	Transcatheter mitral repair procedure		112000001801	ACC NCDR
TMVR	Transcatheter mitral valve replacement		112000001458	ACC NCDR
Tricuspid Valve Procedure	Transcatheter tricuspid valve procedures include eithe a transcatheter tricuspid valve replacement or transcatheter tricuspid valve repair.	er	112000001977	ACC NCDR



Element: 11004



Full Specifications Data Dictionary v3.0



Section: Follow Up Parent: Root

Follow-Up Status Coding Instruction: Indicate whether the patient was alive or deceased at the date the follow-up was performed.

Target Value: The value on Follow-up

Technical Specification Code: 308273005 Code System: SNOMED CT Short Name: F_Status

Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: No Is Followup Element: Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:**

> Valid Range: Data Source: User

Follow-Up Status - 1.3.6.1.4.1.19376.1.4.1.6.5.372

Selection Definition Source Code **Code System** Alive 438949009 SNOMED CT Deceased 20 HL7 Discharge disposition 399307001 SNOMED CT Lost to follow-up

Element: 14338 Follow-Up Reference Discharge Date

Coding Instruction: Indicate the date of discharge for the episode of care that included the reference procedure.

Target Value: The value on Follow-up

Vendor Instruction: Follow-Up Reference Discharge Date (14338) must not be Null

Technical Specification

Code: 112000001859 Code System: ACC NCDR

Short Name: FU_RefDischargeDate

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No Is Followup Yes Element: Data Type: DT Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range:

Valid Range: Data Source: User







Section: Follow Up	Parent: Root	
Element: 11006	Follow-Up Date of Death	Technical Specification
Coding Instruction:	Indicate the date the patient was declared dead.	Code: 1000142373 Code System: ACC NCDR
Target Value:	The value on Follow-up	Short Name: F_DeathDate
Vendor Instruction:	Follow-Up Date of Death (11006) must be Greater than Follow-Up Reference Procedure Start Date and Time (11001)	Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP)
	Follow-Up Date of Death (11006) must be Greater than or Equal to Follow-Up Reference Discharge Date (14338)	Is Identifier: No Is Base Element: No
	Follow-Up Date of Death (11006) must be Less than or Equal to Follow-Up Assessment Date	Is Followup Yes Element:
	(11000)	Data Type: DT Precision:
		Selection Type: Single Unit of Measure: Default Value: Null Usual Range:
		Valid Range: Data Source: User
		Parent/Child Validation Element: 11004 Follow-Up Status Operator: Equal

Element: 11003 Method to Determine Follow-Up Status

Coding Instruction: Indicate the method to determine follow-up status.

Target Value: The value on Follow-up

Technical Specification

Code: 100014059
Code System: ACC NCDR
Short Name: F_Method
Missing Data: Report

Value: Deceased

Harvested: Yes (TAVR, TMVR, TMVrpr,

TVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Multiple
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Method to Determine Follow-up status - 1.3.6.1.4.1.19376.1.4.1.6.5.370

Selection	Definition	Source	Code	Code System
Office Visit			183654001	SNOMED CT
Medical Records			100014060	ACC NCDR
Letter from Medical Prov	vider		100014061	ACC NCDR
Phone Call			100014062	ACC NCDR
Social Security Death M	laster (1000142362	ACC NCDR
Hospitalized			1000142363	ACC NCDR
Obituary List			112000001406	ACC NCDR
Centers for Medicare ar Medicaid Services Linke			112000001407	ACC NCDR
Other			100000351	ACC NCDR



Element: 11007



Full Specifications **Data Dictionary v3.0**



Section: Follow Up Parent: Root

Coding Instruction: Indicate the primary cause of death, i.e. the first significant abnormal event which ultimately led

to death

Cause of Death

Target Value: The value on Follow-up

Technical Specification

Code: 184305005
Code System: SNOMED CT
Short Name: F_DeathCause
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 11004 Follow-Up Status

Operator: Equal Value: Deceased

Cause of Death Clinical Finding - 1.3.6.1.4.1.19376.1.4.1.6.5.88

Selection	Definition	Source	Code	Code System
Acute myocardial infarction	Death by any cardiovascular mechanism (e.g., arrhythmia, sudden death, heart failure, stroke, pulmonary embolus, peripheral arterial disease) withi 30 days after an acute myocardial infarction, related the immediate consequences of the MI, such as progressive HF or recalcitrant arrhythmia. There may be other assessable (attributable) mechanisms of cardiovascular death during this time period, but for simplicity, if the cardiovascular death occurs <=30 days of an acute myocardial infarction, it will be considered a death due to myocardial infarction.	to	100000960	ACC NCDR
Sudden cardiac death	Death that occurs unexpectedly, and not within 30 days of an acute MI.		100000978	ACC NCDR
Heart failure	Death associated with clinically worsening symptoms and/or signs of heart failure.	S	100000964	ACC NCDR
Stroke	Death after a stroke that is either a direct consequen of the stroke or a complication of the stroke.	ce	100000977	ACC NCDR
Cardiovascular procedure	Death caused by the immediate complication(s) of a cardiovascular procedure.		100000962	ACC NCDR
Cardiovascular hemorrhage	Death related to hemorrhage such as a non-stroke intracranial hemorrhage, non-procedural or non-traumatic vascular rupture (e.g., aortic aneurysm), or hemorrhage causing cardiac tamponade.		100000961	ACC NCDR
Other cardiovascular reason	Cardiovascular death not included in the above categories but with a specific, known cause (e.g., pulmonary embolism, peripheral arterial disease).		100000972	ACC NCDR
Pulmonary	Non-cardiovascular death attributable to disease of the lungs (excludes malignancy).	ne	100000975	ACC NCDR
Renal	Non-cardiovascular death attributable to renal failure.		100000976	ACC NCDR
Gastrointestinal	Non-cardiovascular death attributable to disease of the esophagus, stomach, or intestines (excludes malignancy).	ne	100000963	ACC NCDR
Hepatobiliary	Non-cardiovascular death attributable to disease of the liver, gall bladder, or biliary ducts (exclude malignance)		100000966	ACC NCDR
Pancreatic	Non-cardiovascular death attributable to disease of the pancreas (excludes malignancy).	ne	100000974	ACC NCDR
Infection	Non-cardiovascular death attributable to an infectious disease.	3	100000967	ACC NCDR
Inflammatory/Immunologic	Non-cardiovascular death attributable to an inflammatory or immunologic disease process.		100000968	ACC NCDR
Hemorrhage	Non-cardiovascular death attributable to bleeding tha not considered cardiovascular hemorrhage or stroke per this classification.		100000965	ACC NCDR
Non-cardiovascular procedure or surgery	Death caused by the immediate complication(s) of a non-cardiovascular procedure or surgery.		100000971	ACC NCDR







Section: Follow Up		Parent: Root	
Trauma	Non-cardiovascular death attributable to trauma.	100000980	ACC NCDR
Suicide	Non-cardiovascular death attributable to suicide.	100000979	ACC NCDR
Neurological	Non-cardiovascular death attributable to disease of the nervous system (excludes malignancy).	100000970	ACC NCDR
Malignancy	Non-cardiovascular death attributable to malignancy.	100000969	ACC NCDR
Other non-cardiovascular reason	Non-cardiovascular death attributable to a cause other than those listed in this classification (specify organ system).	100000973	ACC NCDR

Element: 13805 Residence

Coding Instruction: Indicate the primary residence of the patient at the time of follow-up.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001506
Code System: ACC NCDR
Short Name: F_Residence
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)
Is Identifier: No

Is Base Element: No
Is Followup Yes
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation
Element: 11004 Follow-Up Status

Operator: Equal Value: Alive

Element: 14511 Residence Not Documented

Operator: Equal

Value: No (or Not Answered)

Residence - 1.3.6.1.4.1.19376.1.4.1.6.5.562

Selection	Definition	Source	Code	Code System
Home with No Health Aid	The patient lives at home with no health-aid (this includes living in senior living facilities with no assistance).		112000001507	ACC NCDR
Home with Health Aid	The patient lives at home with health-aid (this include living in senior living facilities with assistance).	S	112000001508	ACC NCDR
Long Term Care	The patient lives in a long-term care facility that provides the person's health or personal care needs during a short or long period of time.	National Institute of Aging at the National Institutes of Health	42665001	SNOMED CT
Other			100000351	ACC NCDR







Section: Follow Up Parent: Root

Element: 14511 Residence Not Documented

Coding Instruction: Indicate if the primary residence of the patient was not documented during follow-up.

Target Value: N/A

Technical Specification

Code: 112000001506
Code System: ACC NCDR
Short Name: F_ResidenceND

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 11004 Follow-Up Status

Operator: Equal Value: Alive



Element: 13775



Full Specifications Data Dictionary v3.0



Section: Follow-Up Clinical Assessment Parent: Follow Up

Hemoglobin

Coding Instruction: Indicate the hemoglobin (Hgb) value in g/dL.

Note(s):

This may include POC (Point of Care) testing results or results obtained prior to arrival at this

facility.

Target Value: The last value between discharge (or previous follow-up) and current follow-up assessment

Supporting Definition: Hemoglobin

Hemoglobin (Hb or Hgb) is the iron-containing oxygen-transport metalloprotein in the red blood cells. It carries oxygen from the lungs to the rest of the body (i.e. the tissues) where it releases the oxygen to burn nutrients and provide energy. Hemoglobin concentration measurement is among the most commonly performed blood tests, usually as part of a complete blood count. If the concentration is below normal, this is called anemia. Anemias are classified by the size of red blood cells: "microcytic" if red cells are small, "macrocytic" if they are large, and "normocytic" if otherwise. Dehydration or hyperhydration can greatly influence

measured hemoglobin levels.

Source: http://s.details.loinc.org/LOINC/718-7.html?sections=Simple

Technical Specification

Code: 718-7 Code System: LOINC **Short Name:** FU_ProcHgb1 Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: No Is Followup Element: Data Type: PQ Precision: 4,2 Selection Type: Single Unit of Measure: g/dL Default Value: Null

> Usual Range: 5.00 - 20.00 g/dL Valid Range: 1.00 - 50.00 g/dL

Data Source: User

Parent/Child Validation Element: 14326 Hemoglobin Not Drawn

Operator: Equal

Value: No (or Not Answered)

Technical Specification Element: 14326 Hemoglobin Not Drawn

Coding Instruction: Indicate if a follow-up hemoglobin was not collected.

Target Value: N/A

Code: 718-7 Code System: LOINC Short Name: FUHgbND Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: No Is Followup Element: Data Type: BL Precision: Selection Type: Single Unit of Measure:

Default Value: Null Usual Range: Valid Range: Data Source: User







Section: Follow-Up Clinical Assessment

Parent: Follow Up

Element: 13310 Creatinine

Coding Instruction: Indicate the creatinine value.

Target Value: The last value between discharge (or previous follow-up) and current follow-up assessment

Supporting Definition: Creatinine

Creatinine or creatine anhydride, is a breakdown product of creatine phosphate in muscle. The loss of water molecule from creatine results in the formation of creatinine. It is transferred to the kidneys by blood plasma, whereupon it is eliminated by glomerular filtration and partial tubular excretion. Creatinine is usually produced at a fairly constant rate and measuring its serum level is a simple test. A rise in blood creatinine levels is observed only with marked damage to functioning nephrons; therefore this test is not suitable for detecting early kidney disease. Creatine and creatinine are metabolized in the kidneys, muscle, liver and pancreas.

Source: http://s.details.loinc.org/LOINC/2160-0.html?sections=Simple

Technical Specification

Code: 2160-0
Code System: LOINC
Short Name: Follow_Creat
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: mg/dL
Default Value: Null

Usual Range: 0.10 - 9.00 mg/dL Valid Range: 0.10 - 30.00 mg/dL

Data Source: User

Parent/Child Validation

Operator: Equal

Value: No (or Not Answered)

Element: 13311 Creatinine Not Drawn

Element: 13311 Creatinine Not Drawn

Coding Instruction: Indicate if a follow-up creatinine level was not collected.

Target Value: N/A

Technical Specification

Code: 2160-0 Code System: LOINC

Short Name: FollowCreatinineNotDrawn

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Is Identifier: No







Section: Follow-Up Clinical Assessment

Parent: Follow Up

Element: 13688 New York Heart Association Classification

Coding Instruction: Indicate the patient's latest dyspnea or functional class, coded as the New York Heart

Association (NYHA) classification.

Target Value: The value on Follow-up

Supporting Definition: NYHA

The NYHA classes focus on exercise capacity and the symptomatic status of the disease.

Source: 2013 ACCF/AHA Guideline for the Management of Heart Failure; J Am Coll Cardiol.

2013;62(16):e147-e239. doi:10.1016/j.jacc.2013.05.019

Technical Specification

Code: 420816009
Code System: SNOMED CT
Short Name: F_NYHA
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14333 New York Heart Association

Classification Not Documented

Operator: Equal

Value: No (or Not Answered)

NYHA Functional Classification - 1.3.6.1.4.1.19376.1.4.1.6.5.8

Selection	Definition	Source	Code	Code System
Class I	Patients with cardiac disease but without resulting limitations of physical activity. Ordinary physical activit does not cause undue fatigue, palpitation, or dyspnea.	The Criteria Committee of the New York Heart y Association. Nomenclature and Criteria for Diagnosis of Diseases of the Heart and Great Vessels. 9th ed. Boston, Mass: Little, Brown & Co; 1994:253-256.	420300004	SNOMED CT
Class II	Patients with cardiac disease resulting in slight limitation of physical activity. They are comfortable at rest. Ordinary physical activity results in fatigue, palpitation or dyspnea.		421704003	SNOMED CT
Class III	Patients with cardiac disease resulting in marked limitation of physical activity. They are comfortable at rest. Less than ordinary activity causes fatigue, palpitation, or dyspnea.		420913000	SNOMED CT
Class IV	Patients with cardiac disease resulting in inability to carry on any physical activity without discomfort. Symptoms are present even at rest or minimal exertion If any physical activity is undertaken, discomfort is increased.		422293003	SNOMED CT

	Element: 14333	New York Heart Association Classification Not Documented	recnnical Specification
	Coding Instruction:	: Indicate if NYHA was not documented during the follow-up assessment period.	Code: 420816009
			Code System: SNOMED CT

Target Value: The value on Follow-up

Short Name: F_NYHAND

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Base Element: No







Section: Follow-Up Clinical Assessment

Parent: Follow Up

Element: 13689 12 Lead Electrocardiogram Performed

Coding Instruction: Indicate if a 12 lead ECG was performed in the follow-up assessment period.

Target Value: The value on Follow-up

Technical Specification

Code: 164847006 Code System: SNOMED CT Short Name: F_12LeadEKG Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:
Data Source: User

Element: 13621 12 Lead Electrocardiogram Findings

Coding Instruction: Indicate the 12 lead ECG findings during follow-up. If more than one ECG is performed,

document the findings from any ECG.

Target Value: The value on Follow-up

Vendor Instruction: Cannot select option No Significant Changes with any other option: Pathological Q Wave,

Cardiac Arrhythmia, New Left Bundle Branch Block, Pathological Q Wave, Cardiac Arrhythmia

or New Left Bundle Branch Block

Technical Specification

Code: 112000001362
Code System: ACC NCDR
Short Name: F_EKGChange
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Init of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13689 12 Lead Electrocardiogram

Performed

Operator: Equal Value: Yes

12 Lead Electrocardiogram Findings - 1.3.6.1.4.1.19376.1.4.1.6.5.535

Selection	Definition	Source	Code	Code System
Cardiac Arrhythmia	ardiac Arrhythmia The patient has a new onset of an atrial or ventricular arrhythmia requiring medication or other therapy. This includes brady or tachy arrhythmias.		698247007	SNOMED CT
No Significant Changes			112000001391	ACC NCDR
Pathological Q Wave			164918000	SNOMED CT
New Left Bundle Branch Block			100014019	ACC NCDR







Section: Follow-Up Imaging

Parent: Follow-Up Echocardiogram

Element: 13492 Echocardiogram Performed

Coding Instruction: Indicate whether an echo (and the type of echo) was performed in the follow-up assessment

period

Target Value: Any occurrence on follow-up

Technical Specification

Code: 40701008
Code System: SNOMED CT
Short Name: F_POpTTEch
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14512 Echocardiogram Not Performed

Operator: Equal

Value: No (or Not Answered)

Echocardiogram Type - 1.3.6.1.4.1.19376.1.4.1.6.5.526

 Selection
 Definition
 Source
 Code System

 Transesophageal Echocardiogram (TEE)
 105376000
 SNOMED CT

 Transthoracic Echo (TTE)
 433236007
 SNOMED CT

Element: 14512 Echocardiogram Not Performed

Coding Instruction: Indicate if an echocardiogram was not performed during follow-up.

Target Value: N/A

Technical Specification

Code: 40701008

Code System: SNOMED CT
Short Name: F_EchoND
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User







Section: Follow-Up Imaging

Parent: Follow-Up Echocardiogram

Element: 13593 Echocardiogram Date

Coding Instruction: Indicate the date the echocardiogram was performed.

Target Value: Any occurrence on follow-up

Technical Specification

Code: 40701008

Code System: SNOMED CT
Short Name: F_POpTTEchDate

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13690 Left Ventricular Ejection Fraction

Coding Instruction: Indicate the left ventricular ejection fraction.

Target Value: The value on Follow-up Supporting Definition: Most Recent LVEF %

The left ventricular ejection fraction is the percentage of blood emptied from the left ventricle at

the end of contraction.

Source: ACC Clinical Data Standards, Society for Thoracic Surgeons Adult Cardiac Surgery

Database (STS)

Technical Specification

Code: 10230-1
Code System: LOINC
Short Name: F_LVEF
Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: %
Default Value: Null
Usual Range: 5 - 90 %
Valid Range: 1 - 99 %
Data Source: User

Parent/Child Validation
Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND

Element: 13691 Left Ventricular Ejection Fraction

Not Assessed

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up Imaging

Parent: Follow-Up Echocardiogram

Element: 13691 Left Ventricular Ejection Fraction Not Assessed

Coding Instruction: Indicate whether the left ventricular ejection fraction was not assessed.

Target Value: The value on Follow-up

Technical Specification

Code: 100001027
Code System: ACC NCDR
Short Name: F_LVEFNA
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)







Section: Follow-Up Aortic Valve

Parent: Follow-Up Echocardiogram

Element: 13676 Aortic Valve Mean Gradient

Coding Instruction: Indicate the highest aortic valve mean gradient in mm Hg.

Target Value: The highest value on follow up

Technical Specification

Code: 112000001398
Code System: ACC NCDR
Short Name: F_AVMeanGradient

Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)

| Is Identifier: No |
| Is Base Element: No |
| Is Followup Element: PQ |
| Precision: 3,0 |
| Selection Type: Single |
| Unit of Measure: mm[Hg] |
| Default Value: Null |
| Usual Range: 5-50 mm[Hg]

Usual Range: 5 - 50 mm[Hg]
Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13669 Aortic Valve Area

Coding Instruction: Indicate the smallest aortic valve area, in cm2.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001280
Code System: ACC NCDR
Short Name: F_AVArea
Missing Data: Report
Harvested: Yes (TAVR)
Is Identifier: No
Is Base Element: No

Is Followup
Element:
Data Type: PQ
Precision: 3,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 0.20 - 4.00 cm2 **Valid Range:** 0.05 - 5.00 cm2

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR







Section: Follow-Up AV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 13527 Aortic Valve Regurgitation

Coding Instruction: Indicate the severity of aortic valve regurgitation.

If mild-moderate is documented, code as mild.

If moderate-severe is documented, code as moderate.

Target Value: The value on Follow-up

Technical Specification

Code: 60234000
Code System: SNOMED CT
Short Name: F_AR
Missing Data: Report

Harvested: Yes (BDS, TAVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up AV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14504 Paravalvular Aortic Regurgitation

Coding Instruction: Indicate the severity of paravalvular aortic regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001428 Code System: ACC NCDR Short Name: F_ParaAR Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: CD Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial ----- AND -----

Element: 14527 Paravalvular Aortic Regurgitation

Not Documented

Operator: Equal

Value: No (or Not Answered)

----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: TAVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

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Selection	Definition	Source	Code	Code System		
None			112000001910	ACC NCDR		
Mild			112000000380	ACC NCDR		
Moderate			112000000381	ACC NCDR		
Severe			112000000382	ACC NCDR		







Section: Follow-Up AV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14527 Paravalvular Aortic Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular aortic regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: F_ParaARND
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: No
Is Followup Element: Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Operator: Equal Value: Mild

Element: 13527 Aortic Valve Regurgitation

Operator: Equal
Value: Moderate

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial







Section: Follow-Up AV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14500 Central Aortic Regurgitation

Coding Instruction: Indicate the severity of central aortic regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_CentAR
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13527 Aortic Valve Regurgitation

Operator: Equal
Value: Trace/Trivial
----- AND

Element: 14490 Central Aortic Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

······································				
Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up AV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14490 Central Aortic Regurgitation Not Documented

Coding Instruction: Indicate if central aortic regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_CentARND
Missing Data: Report
Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: No
Is Followup Element: Percision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 13527 Aortic Valve Regurgitation

Operator: Equal
Value: Moderate

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 13527 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13778 Mitral Valve Mean Gradient

Coding Instruction: Indicate the highest mitral valve mean gradient, in mm Hg.

Target Value: The highest value on follow up

Supporting Definition: Mitral Valve Mean Gradient

The average gradient across the mitral valve occurring during the entire systole.

Source: Baumgartner, H. et. al (2009). Echocardiographic assessment of valve stenosis:

EAE/ASE

recommendations for clinical practice.

Technical Specification

Code: 112000001191
Code System: ACC NCDR
Short Name: F_MeanMVGrad

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Usual Range: 5 - 50 mm[Hg]
Valid Range: 0 - 150 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13768 Effective Regurgitant Orifice Area

Coding Instruction: Indicate the effective regurgitant orifice area (EROA), in cm2.

Target Value: The highest value on follow up

Technical Specification

Code: 112000001437
Code System: ACC NCDR
Short Name: F_MV_EOA
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: PQ
Precision: 2,1
Selection Type: Single
Unit of Measure: cm2
Default Value: Null
Usual Range: 0.1 - 5.0 cm2
Valid Range: 0.1 - 5.0 cm2
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Element: 13780 Effective Regurgitant Orifice Area Method of Assessment

Coding Instruction: Indicate the method used to assess the effective orifice area. If multiple methods are available,

code the 3D planimetry method first, then PISA.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001437
Code System: ACC NCDR
Short Name: F_MV_EOA_MOA

Missing Data: Report

Is Identifier: No

Harvested: Yes (TMVR, TMVrpr)

Is Base Element: No
Is Followup Yes
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13768 Effective Regurgitant Orifice Area

Operator:

Value: Any Value

Effective Regurgitant Orifice Area Method of Assessment - 1.3.6.1.4.1.19376.1.4.1.6.5.547

Selection	Definition	Source	Code	Code System
3D Planimetry			112000001438	ACC NCDR
Proximal Isovelocity Surface			112000001439	ACC NCDR
Area				
Quantitative Doppler			112000001440	ACC NCDR
Other			100000351	ACC NCDR







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13781 Mitral Valve Area

Coding Instruction: Indicate the smallest mitral valve area in centimeters squared.

Target Value: The value on Follow-up

Supporting Definition: Mitral Valve Area

Measurement of mitral valve area.

Source:

Technical Specification

Code: 251012002
Code System: SNOMED CT
Short Name: F_MVA
Missing Data: Report
Harvested: Yes (TMVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: PQ
Precision: 4,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Usual Range: 3.00 - 6.00 cm2 **Valid Range:** 0.05 - 12.00 cm2

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Reference Procedure Type

rator: Equal

Operator: Equal Value: TMVR

Element: 13773 Left Ventricular Outflow Tract Peak Velocity

Coding Instruction: Indicate the left ventricular outflow tract peak velocity in m/sec.

Target Value: The highest value on follow up

Technical Specification

 Code:
 112000002047

 Code System:
 ACC NCDR

 Short Name:
 F_LVOT

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR)

Is Base Element: No
Is Followup
Element: Yes
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: m/sec
Default Value: Null

Is Identifier: No

Usual Range: 0.5 - 5.0 m/sec Valid Range: 0.1 - 10.0 m/sec Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13782 Systolic Anterior Motion Present

Coding Instruction: Indicate if systolic anterior motion of the mitral valve was present.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001481 Code System: ACC NCDR Short Name: F_SAM Missing Data: Report Harvested: Yes (TMVR)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: BL Precision: Selection Type: Single Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) AND -----

Element: 13705 Transcatheter Valve Therapy Reference Procedure Type

Operator: Equal Value: TMVR







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13783 Left Ventricular Internal Systolic Dimension

Coding Instruction: Indicate the left ventricular internal systolic dimension in cm.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001424 Code System: ACC NCDR Short Name: F_LVIDs Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: PQ Precision: 2,1 Selection Type: Single Unit of Measure: cm Default Value: Null Usual Range: 2.5 - 4.5 cm Valid Range: 1.0 - 9.0 cm Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

----- AND -----Element: 14536 Left Ventricular Internal Systolic

Dimension Not Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14536 Left Ventricular Internal Systolic Dimension Not Measured

Coding Instruction: Indicate if the left ventricular internal systolic dimension was not measured.

Target Value: N/A

Technical Specification

Code: 112000001424
Code System: ACC NCDR
Short Name: F_LVIDs_NM
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Pecsion:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

value: INVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13784 Left Ventricular Internal Diastolic Dimension

Coding Instruction: Indicate the left ventricular internal diastolic dimension in cm.

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001425

 Code System:
 ACC NCDR

 Short Name:
 F_LVIDd

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Yes
Element:
Data Type: PQ
Precision: 3,1
Selection Type: Single
Unit of Measure: cm
Default Value: Null
Usual Range: 3.5 - 5.5 cm
Valid Range: 1.0 - 10.0 cm
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Dimension Not Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14537 Left Ventricular Internal Diastolic Dimension Not Measured

Coding Instruction: Indicate if the left ventricular internal diastolic dimension was not measured.

Target Value: N/A

Technical Specification

 Code:
 112000001425

 Code System:
 ACC NCDR

 Short Name:
 F_LVIDd_NM

 Missing Data:
 Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

value: TIVIVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13786 Left Ventricular End Systolic Volume

Coding Instruction: Indicate the left ventricular end systolic volume in ml.

Target Value: The value on Follow-up

Technical Specification

Code: 250931004 Code System: SNOMED CT Short Name: F_LVESV Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mL Default Value: Null Usual Range: 10 - 150 mL Valid Range: 1 - 300 mL Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr ----- AND -----

Element: 14539 Left Ventricular End Systolic

Volume Not Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14539 Left Ventricular End Systolic Volume Not Measured

Coding Instruction: Indicate if the left ventricular end systolic volume was not measured.

Target Value: N/A

Technical Specification

Code: 250931004
Code System: SNOMED CT
Short Name: F_LVESV_NM
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Element: Percision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13785 Left Ventricular End Diastolic Volume

Coding Instruction: Indicate the left ventricular end diastolic volume in ml.

Target Value: The value on Follow-up

Technical Specification

Code: 250932006 Code System: SNOMED CT Short Name: F_LVEDV Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mL Default Value: Null Usual Range: 40 - 250 mL Valid Range: 1 - 400 mL Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr ----- AND -----

Element: 14538 Left Ventricular End Diastolic

Volume Not Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14538 Left Ventricular End Diastolic Volume Not Measured

Coding Instruction: Indicate if the left ventricular end diastolic volume was not measured.

Target Value: N/A

Technical Specification

Code: 250932006
Code System: SNOMED CT
Short Name: F_LVEDV_NM
Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Element: Percision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

value: HVIVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13787 Left Atrial Volume

Coding Instruction: Indicate the left atrial volume in ml.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001426 Code System: ACC NCDR Short Name: F_LAVol Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: mL Default Value: Null Usual Range: 10 - 90 mL Valid Range: 1 - 500 mL Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr ----- AND -----

Element: 14540 Left Atrial Volume Not Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14540 Left Atrial Volume Not Measured

Coding Instruction: Indicate if the left atrial volume was not measured.

Target Value: N/A

Technical Specification

 Code:
 112000001426

 Code System:
 ACC NCDR

 Short Name:
 F_LAVol_NM

 Missing Data:
 Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND ------

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

value: HVIVK

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 13788 Left Atrial Volume Index

Coding Instruction: Indicate the left atrial volume index in mL/m2.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001427 Code System: ACC NCDR Short Name: F_LAVolIndex Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No Is Base Element: No Is Followup Element: Yes Data Type: PQ Precision: 3,0 Selection Type: Single Unit of Measure: ml/m2 Default Value: Null Usual Range: 10 - 90 ml/m2 Valid Range: 1 - 250 ml/m2

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Data Source: User

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr ----- AND -----

Element: 14582 Left Atrial Volume Index Not

Measured

Operator: Equal

Value: No (or Not Answered)







Section: Follow-Up MV Imaging

Parent: Follow-Up Echocardiogram

Element: 14582 Left Atrial Volume Index Not Measured

Coding Instruction: Indicate if the left atrial volume index was not measured.

Target Value: N/A

Technical Specification

Code: 112000001427
Code System: ACC NCDR
Short Name: F_LAVolIndex_NM

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR







Section: Follow-Up MV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 13673 Mitral Regurgitation

Coding Instruction: Indicate highest level of mitral regurgitation.

If mild-moderate is documented, code as mild.

Target Value: The value on Follow-up

Technical Specification

Code: 48724000
Code System: SNOMED CT
Short Name: F_MR
Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr,

TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND ------

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.728

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Moderate-Severe			1000142345	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up MV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 13776 Paravalvular Mitral Regurgitation

Coding Instruction: Indicate the severity of paravalvular mitral regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: F_ParaMR
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13673 Mitral Regurgitation

Operator: Equal
Value: Moderate

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up MV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14528 Paravalvular Mitral Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular mitral regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: F_ParaMRND
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: No
Is Followup Element: Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Follow-Up MV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 13777 Central Mitral Regurgitation

Coding Instruction: Indicate the severity of central mitral regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_CentralMR
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: No
Is Followup Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Mild

Element: 13673 Mitral Regurgitation

Operator: Equal
Value: Moderate

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe

AND -----Element: 14491 Central Mitral Regurgitation Not

Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

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Selection	Definition	Source	Code	Code System	
None			112000001910	ACC NCDR	
Mild			112000000380	ACC NCDR	
Moderate			112000000381	ACC NCDR	
Severe			112000000382	ACC NCDR	







Section: Follow-Up MV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14491 Central Mitral Regurgitation Not Documented

Coding Instruction: Indicate if central mitral regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_CentralMRND
Missing Data: Report

Harvested: Yes (BDS, TMVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

----- AND -----

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13673 Mitral Regurgitation

Operator: Equal

Operator: Equal Value: Mild

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Severe

Element: 13673 Mitral Regurgitation

Operator: Equal Value: Trace/Trivial

Element: 13673 Mitral Regurgitation

Operator: Equal

Value: Moderate-Severe







Section: Follow-Up TV Imaging

Parent: Follow-Up Echocardiogram

Element: 14545 Tricuspid Valve Diastolic Gradient

Coding Instruction: Indicate the tricuspid valve diastolic gradient in mm Hg. This can also be called the TV inflow

gradient

Target Value: The highest value on follow up

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: F_TVDGrad
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: PQ
Precision: 2,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 1 - 15 mm[Hg]
Valid Range: 1 - 50 mm[Hg]
Data Source: User

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Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14546 Tricuspid Valve Diastolic Gradient
Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14546 Tricuspid Valve Diastolic Gradient Not Documented

Coding Instruction: Indicate if the tricuspid valve diastolic gradient was not documented.

Target Value: N/A

Technical Specification

Code: 112000001512
Code System: ACC NCDR
Short Name: F_TVDGradND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Is Base Element: No

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Data Source: User

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
----- AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Imaging

Parent: Follow-Up Echocardiogram

Element: 14547 Tricuspid Valve Annulus Size

Coding Instruction: Indicate the tricuspid valve annulus size in mm. Document the size using end-diastolic, 4

chamber view is preferred (in mm).

Target Value: The value on Follow-up

Technical Specification

Code: 112000001513

Code System: ACC NCDR Short Name: F_TVAnnulus

Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No

Is Base Element: No
Is Followup
Element: Yes

Element: Yes

Data Type: PQ

Precision: 2,0
Selection Type: Single
Unit of Measure: mm
Default Value: Null

Usual Range: 15 - 60 mm Valid Range: 1 - 80 mm Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

----- AND -----

Element: 14548 Tricuspid Valve Annulus Size Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14548 Tricuspid Valve Annulus Size Not Documented

Coding Instruction: Indicate if the tricuspid valve annulus size was not documented.

Target Value: N/A

Technical Specification

Code: 112000001513
Code System: ACC NCDR
Short Name: F_TVAnnulusND

Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes

Data Type: BL
Precision:
Selection Type: Single

Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Imaging

Parent: Follow-Up Echocardiogram

Element: 14549 End Diastolic Mid Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic mid right ventricular (RV) diameter, using the 4 chamber view (in cm).

Target Value: The value on Follow-up

Technical Specification

Code: 112000001514 Code System: ACC NCDR Short Name: F_MidRVDia

Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No Is Base Element: No

Is Followup Element: Data Type: PQ Precision: 2,1 Selection Type: Single Unit of Measure: cm

Default Value: Null Usual Range: 1.0 - 7.0 cm Valid Range: 0.1 - 9.9 cm Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

AND Element: 14550 End Diastolic Mid Right Ventricle

Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14550 End Diastolic Mid Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the end-diastolic mid right ventricular diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001514 Code System: ACC NCDR Short Name: F_MidRVDiaND Missing Data: Report Harvested: Yes (TTVP)

Is Base Element: No Is Followup Element: Yes Data Type: BL Precision: Selection Type: Single

Is Identifier: No

Default Value: Null **Usual Range:** Valid Range: Data Source: User

Unit of Measure:

Parent/Child Validation

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE) ----- AND -----

Element: 13705 Transcatheter Valve Therapy Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Imaging

Parent: Follow-Up Echocardiogram

Element: 14551 End Diastolic Basal Right Ventricle Diameter

Coding Instruction: Indicate the end-diastolic basal right ventricular (RV) diameter, using the 4 chamber view (in

cm)

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001515

 Code System:
 ACC NCDR

 Short Name:
 F_BasalRVDia

 Missing Data:
 Report

Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: No

Is Followup
Element:

Pata Type: PQ
Precision: 2,1

Selection Type: Single
Unit of Measure: cm
Default Value: Null

Usual Range: 1.0 - 7.0 cm Valid Range: 0.1 - 9.9 cm Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14552 End Diastolic Basal Right

lement: 14552 End Diastolic Basal Right

Ventricle Diameter Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14552 End Diastolic Basal Right Ventricle Diameter Not Documented

Coding Instruction: Indicate if the basal diastolic mid right ventricular (RV) diameter was not documented.

Target Value: N/A

Technical Specification

Code: 112000001515
Code System: ACC NCDR
Short Name: F_BasalDiaND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: No

Is Followup
Element:

Data Type:
BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)
------ AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Imaging

Parent: Follow-Up Echocardiogram

Element: 14553 Right Ventricular Systolic Pressure

Coding Instruction: Indicate the right ventricular systolic pressure in mm Hg.

Target Value: The highest value on follow up

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: F_RVSP
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: PQ

Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null

Usual Range: 15 - 30 mm[Hg] Valid Range: 1 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed **Operator:** Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 14554 Right Ventricular Systolic

Pressure Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 14554 Right Ventricular Systolic Pressure Not Documented

Coding Instruction: Indicate if the right ventricular systolic pressure was not documented.

Target Value: N/A

Supporting Definition: RV Systolic Pressure

The maximum pressure exerted into the systemic arterial circulation during the contraction of

the right ventricle of the heart

Source: NCI EVS

Technical Specification

Code: 276772001
Code System: SNOMED CT
Short Name: F_RVSPND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No
Is Base Element: No
Is Followup

Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

AND

Element: 13705 Transcatheter Valve Therapy Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 13678 Tricuspid Valve Regurgitation

Coding Instruction: Indicate the severity of tricuspid regurgitation.

If mild-moderate is documented, code as mild.

If moderate-severe is documented, code as moderate.

Target Value: The value on Follow-up

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: F_Post_TR
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transthoracic Echo (TTE)

Element: 13492 Echocardiogram Performed

Operator: Equal

Value: Transesophageal Echocardiogram (TEE)

AND

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up TV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14506 Paravalvular Tricuspid Regurgitation

Coding Instruction: Indicate the severity of paravalvular tricuspid regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: F_ParaTR
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal

Value: Moderate

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal
Value: Trace/Trivial
----- AND

Element: 14529 Paravalvular Tricuspid

Regurgitation Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up TV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14529 Paravalvular Tricuspid Regurgitation Not Documented

Coding Instruction: Indicate if the severity of paravalvular tricuspid regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 112000001428
Code System: ACC NCDR
Short Name: F_ParaTRND
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal

Value: Moderate

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Trace/Trivial

AND -----
Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal







Section: Follow-Up TV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14502 Central Tricuspid Regurgitation

Coding Instruction: Indicate the severity of central tricuspid regurgitation.

Note: If trace/trivial is documented, code "none".

Target Value: The highest value on follow up

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_CenTR
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element:

Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal

Value: Moderate

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13678 Tricuspid Valve Regurgitation

Element: 14492 Central Tricuspid Regurgitation

Not Documented

Operator: Equal

Value: No (or Not Answered)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up TV Regurgitation

Parent: Follow-Up Echocardiogram

Element: 14492 Central Tricuspid Regurgitation Not Documented

Coding Instruction: Indicate if central tricuspid regurgitation was not documented.

Target Value: N/A

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: F_CenTRND
Missing Data: Report
Harvested: Yes (TTVP)
Is Identifier: No

Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Mild

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal
Value: Moderate

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Severe

Element: 13678 Tricuspid Valve Regurgitation

Operator: Equal Value: Trace/Trivial







Section: Follow-Up 4DCTA Parent: Follow Up

Element: 13692 4D Computed Tomography Performed

Coding Instruction: Indicate if a 4D CT was performed.

Target Value: The value on Follow-up

Technical Specification

Code: 241547009
Code System: SNOMED CT
Short Name: F_4DCT
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Pecision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13693 4D Computed Tomography Date

Coding Instruction: Indicate the date the 4D CT was performed.

Target Value: The value on Follow-up

Technical Specification

Code: 241547009
Code System: SNOMED CT
Short Name: F_4DCTdate
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Percision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13692 4D Computed Tomography

Performed
Operator: Equal
Value: Yes







Section: Follow-Up 4DCTA Parent: Follow Up

Element: 13694 Valve Thrombosis

 $\begin{tabular}{ll} \textbf{Coding Instruction:} & Indicate if there was findings of thrombus on the prosthetic valve. \\ \end{tabular}$

Target Value: The value on Follow-up

Code: 112000001917
Code System: ACC NCDR
Short Name: F_VThromb
Missing Data: Report

Technical Specification

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Yes
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 13692 4D Computed Tomography

Performed

Data Source: User

Operator: Equal Value: Yes

Element: 13695 Leaflet Dysfunction Noted

Coding Instruction: Indicate if leaflet dysfunction was noted. Leaflet dysfunction is evident when there is a finding

of "stuck leaflets" on the prosthetic valve.

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001409

 Code System:
 ACC NCDR

 Short Name:
 F_LeafDysFx

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR, TMVR, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element: Data Type: BL
Precision: Selection Type: Single
Unit of Measure: Default Value: Null
Usual Range: Valid Range:
Data Source: User

Parent/Child Validation

Element: 13692 4D Computed Tomography

Performed

Operator: Equal Value: Yes







Section: Follow-Up Six Minute Walk Test

Parent: Follow Up

Element: 13789 Six Minute Walk Test

Coding Instruction: Indicate whether a six minute walk test was performed.

Target Value: The value on Follow-up

Technical Specification

Code: 252478000

Code System: SNOMED CT
Short Name: F_SixMinWalkPerf

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element: Percision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Element: 14263 Six Minute Walk Test Reason Not Performed

Coding Instruction: Indicate the reason the six minute walk test was not performed.

Target Value: The value on Follow-up

Technical Specification

Code: 252478000 Code System: SNOMED CT

Short Name: F_SixMinWalkPerfReason

Missing Data: Report

Is Identifier: No

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13789 Six Minute Walk Test

Operator: Equal Value: No

Six Minute Walk Test Reason Not Performed - 1.3.6.1.4.1.19376.1.4.1.6.5.544

Selection	Definition	Source	Code	Code System
Non-Cardiac Reason			112000001418	ACC NCDR
Cardiac Reason			112000001419	ACC NCDR
Patient Not Willing to Wa	alk		112000001420	ACC NCDR
Not Performed by Site			112000001421	ACC NCDR







Section: Follow-Up Six Minute Walk Test

Parent: Follow Up

Element: 13790 Six Minute Walk Test Date

Coding Instruction: Indicate the date the six minute walk test was performed.

Target Value: The value on Follow-up

Technical Specification
Code: 252478000
Code System: SNOMED CT
Short Name: F_SixMinWalkDate

Missing Data: Report

Is Identifier: No

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13789 Six Minute Walk Test

Operator: Equal Value: Yes

Element: 14325 Six Minute Walk Test Total Distance

Coding Instruction: Indicate the total distance, in feet, the patient walked.

Target Value: The value on Follow-up

Technical Specification
Code: 112000001422
Code System: ACC NCDR
Short Name: F_SixMinWalkDist

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: PQ
Precision: 4,0
Selection Type: Single
Unit of Measure: ft
Default Value: Null
Usual Range: 1 - 3,000 ft
Valid Range: 1 - 3,000 ft
Data Source: User

Parent/Child Validation

Element: 13789 Six Minute Walk Test

Operator: Equal Value: Yes





Section: Follow-Up KCCQ Parent: Follow Up

Element: 13845 Kansas City Cardiomyopathy Questionnaire 12 Performed

Coding Instruction: Indicate if the Kansas City Cardiomyopathy Questionnaire (KCCQ-12) was performed.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001540 Code System: ACC NCDR

Short Name: F_KCCQ12_Performed

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No

Is Followup
Element:
Data Type: BL

Precision:
Selection Type: Single
Unit of Measure:

Usual Range: Valid Range: Data Source: User

Default Value: Null

Element: 13844 Kansas City Cardiomyopathy Questionnaire 12 Date

Coding Instruction: Indicate the date the Kansas City Cardiomyopathy Questionnaire (KCCQ-12) was performed.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001540
Code System: ACC NCDR
Short Name: F_KCCQ12_Date

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes







Section: Follow-Up KCCQ

Parent: Follow Up

Element: 13847 Kansas City Cardiomyopathy Questionnaire 12 Question 1a

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 1a

Heart Failure Limitation - Showering/bathing

Target Value: The value on Follow-up

Technical Specification

Code: 112000001541
Code System: ACC NCDR
Short Name: F_KCCQ12_1a
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 1a thru 1c - 1.3.6.1.4.1.19376.1.4.1.6.5.570

Selection	Definition	Source	Code	Code System
1 - Extremely Limited			100001173	ACC NCDR
2 - Quite a Bit Limited			100001171	ACC NCDR
3 - Moderately Limited	i		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Not at All Limited			100001167	ACC NCDR
6 - Limited for Other F			100014041	ACC NCDR







Section: Follow-Up KCCQ

Parent: Follow Up

Element: 13869 Kansas City Cardiomyopathy Questionnaire 12 Question 1b

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 1b

Heart Failure Limitation - Walking 1 block on level ground

Target Value: The value on Follow-up

Technical Specification

Code: 112000001542
Code System: ACC NCDR
Short Name: F_KCCQ12_1b

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 1a thru 1c - 1.3.6.1.4.1.19376.1.4.1.6.5.570

Selection	Definition	Source	Code	Code System
1 - Extremely Limited			100001173	ACC NCDR
2 - Quite a Bit Limited			100001171	ACC NCDR
3 - Moderately Limited	1		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Not at All Limited			100001167	ACC NCDR
6 - Limited for Other F			100014041	ACC NCDR
or Did Not Do These	Activities			







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13850 Kansas City Cardiomyopathy Questionnaire 12 Question 1c

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 1c

Heart Failure Limitation - Hurrying or jogging

Target Value: The value on Follow-up

Technical Specification

Code: 112000001543
Code System: ACC NCDR
Short Name: F_KCCQ12_1c

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Is Identifier: No

Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 1a thru 1c - 1.3.6.1.4.1.19376.1.4.1.6.5.570

Selection	Definition	Source	Code	Code System
1 - Extremely Limited			100001173	ACC NCDR
2 - Quite a Bit Limited			100001171	ACC NCDR
3 - Moderately Limited	I		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Not at All Limited			100001167	ACC NCDR
6 - Limited for Other R	Reasons		100014041	ACC NCDR
or Did Not Do These A	Activities			







Section: Follow-Up KCCQ

Parent: Follow Up

Element: 13852 Kansas City Cardiomyopathy Questionnaire 12 Question 2

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 2

Symptom Frequency - swelling in legs

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001544

 Code System:
 ACC NCDR

 Short Name:
 F_KCCQ12_2

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 2 - 1.3.6.1.4.1.19376.1.4.1.6.5.571

Selection	Definition	Source	Code	Code System
1 - Every Morning			112000001553	ACC NCDR
2 - Three or More Times Per Week But Not Everyday			112000001554	ACC NCDR
3 - One to Two Times Per Week			112000001555	ACC NCDR
4 - Less Than Once a Week			112000001556	ACC NCDR
5 - Never Over the Past Two Weeks			112000001557	ACC NCDR







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13854 Kansas City Cardiomyopathy Questionnaire 12 Question 3

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 3

Symptom Frequency - fatigue

Target Value: The value on Follow-up

Technical Specification

Code: 112000001545
Code System: ACC NCDR
Short Name: F_KCCQ12_3

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Is Identifier: No

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 3 and 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.572

Selection	Definition	Source	Code	Code System
1 - All the Time			112000001818	ACC NCDR
2 - Several Times Per D	ay		112000001559	ACC NCDR
3 - At Least Once Per D	Day		112000001560	ACC NCDR
4 - Three or More Times Week But Not Everyday			112000001554	ACC NCDR
5 - One to Two Times P Week	er		112000001555	ACC NCDR
6 - Less Than Once a V	Veek		112000001556	ACC NCDR
7 - Never Over the Pas Weeks	t Two		112000001557	ACC NCDR







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13856 Kansas City Cardiomyopathy Questionnaire 12 Question 4

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 4

Symptom Frequency - shortness of breath

Target Value: The value on Follow-up

Technical Specification

Code: 112000001546
Code System: ACC NCDR
Short Name: F_KCCQ12_4

Missing Data: Report
Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 3 and 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.572

Selection	Definition	Source	Code	Code System
1 - All the Time			112000001818	ACC NCDR
2 - Several Times Per Day			112000001559	ACC NCDR
3 - At Least Once Per Day			112000001560	ACC NCDR
4 - Three or More Times Pe Week But Not Everyday	er er		112000001554	ACC NCDR
5 - One to Two Times Per Week			112000001555	ACC NCDR
6 - Less Than Once a Wee	k		112000001556	ACC NCDR
7 - Never Over the Past Tv Weeks	vo		112000001557	ACC NCDR







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13858 Kansas City Cardiomyopathy Questionnaire 12 Question 5

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 5

Symptom Frequency - sleep sitting up due to shortness of breath

Target Value: The value on Follow-up

Technical Specification

Code: 112000001547
Code System: ACC NCDR
Short Name: F_KCCQ12_5
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 5 - 1.3.6.1.4.1.19376.1.4.1.6.5.704

Selection	Definition	Source	Code	Code System
1 - Every Night			112000001819	ACC NCDR
2 - Three or More Times Per Week But Not Everyday			112000001554	ACC NCDR
3 - One to Two Times Per Week			112000001555	ACC NCDR
4 - Less Than Once a Week			112000001556	ACC NCDR
5 - Never Over the Past Two Weeks			112000001557	ACC NCDR







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13860 Kansas City Cardiomyopathy Questionnaire 12 Question 6

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 6

Quality of Life - effect on enjoyment of life due to heart failure

Target Value: The value on Follow-up

Technical Specification

Code: 112000001548
Code System: ACC NCDR
Short Name: F KCC012 6

Short Name: F_KCCQ12_6 Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 6 - 1.3.6.1.4.1.19376.1.4.1.6.5.573

Selection	Definition	Source	Code	Code System
1 - It Has Extremely Li Enjoyment of Life	imited My		100014049	ACC NCDR
2 - It Has Limited My E of Life Quite a Bit	Enjoyment		100014050	ACC NCDR
3 - It Has Moderately I My Enjoyment of Life			100014051	ACC NCDR
4 - It Has Slightly Limit Enjoyment of Life	ted My		100014052	ACC NCDR
5 - It Has Not Limited I Enjoyment of Life at A			100014053	ACC NCDR







Section: Follow-Up KCCQ

Parent: Follow Up

Element: 13862 Kansas City Cardiomyopathy Questionnaire 12 Question 7

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 7

Quality of life - remaining life with heart failure

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001549

 Code System:
 ACC NCDR

 Short Name:
 F_KCCQ12_7

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 7 - 1.3.6.1.4.1.19376.1.4.1.6.5.574

Selection	Definition	Source	Code	Code System
1 - Not At All Satisfied			112000001561	ACC NCDR
2 - Mostly Dissatisfied			112000001562	ACC NCDR
3 - Somewhat Satisfied			112000001563	ACC NCDR
4 - Mostly Satisfied			112000001564	ACC NCDR
5 - Completely Satisfied			112000001565	ACC NCDR

Element: 13864 Kansas City Cardiomyopathy Questionnaire 12 Question 8a

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 8a

Social limitation - hobbies, recreational activities

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001550

 Code System:
 ACC NCDR

 Short Name:
 F_KCCQ12_8a

 Missing Data:
 Report

Is Identifier: No

Data Source: User

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

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Selection	Definition	Source	Code	Code System
1 - Severely Limited		,	112000001566	ACC NCDR
2 - Limited Quite a Bit			112000001567	ACC NCDR
3 - Moderately Limited	i		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Did Not Limit at All			112000001569	ACC NCDR
6 - Does Not Apply or Do for Other Reasons			112000001570	ACC NCDR







Section: Follow-Up KCCQ Parent: Follow Up

Element: 13866 Kansas City Cardiomyopathy Questionnaire 12 Question 8b

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 8b

Social limitation - working or doing household chores

Target Value: The value on Follow-up

Technical Specification

Code: 112000001551
Code System: ACC NCDR
Short Name: F_KCCQ12_8b
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

Selection	Definition	Source	Code	Code System
1 - Severely Limited			112000001566	ACC NCDR
2 - Limited Quite a Bit			112000001567	ACC NCDR
3 - Moderately Limited	I		100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Did Not Limit at All			112000001569	ACC NCDR
6 - Does Not Apply or			112000001570	ACC NCDR







Section: Follow-Up KCCQ

Parent: Follow Up

Element: 13868 Kansas City Cardiomyopathy Questionnaire 12 Question 8c

Coding Instruction: Indicate the patient's response to the Kansas City Cardiomyopathy Questionnaire (KCCQ-12)

Question 8c

Social limitation - visiting family or friends

Target Value: The value on Follow-up

Technical Specification

 Code:
 112000001552

 Code System:
 ACC NCDR

 Short Name:
 F_KCCQ12_8c

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes

Kansas City Cardiomyopathy Questionnaire 12 Answer to Question 8 - 1.3.6.1.4.1.19376.1.4.1.6.5.575

Selection	Definition	Source	Code	Code System
1 - Severely Limited			112000001566	ACC NCDR
2 - Limited Quite a Bit			112000001567	ACC NCDR
3 - Moderately Limited			100001170	ACC NCDR
4 - Slightly Limited			100014042	ACC NCDR
5 - Did Not Limit at All			112000001569	ACC NCDR
6 - Does Not Apply or Did No	ot		112000001570	ACC NCDR
Do for Other Peacons				

Element: 14535 Follow-Up KCCQ Overall Summary Score

Coding Instruction: (Auto Calculated) This field is auto-populated by your application.

Kansas City Cardiomyopathy Questionnaire (KCCQ-12) Overall Summary Score.

Note(s): The 12 patient responses are reduced into four summary scores (Physical Limitation Score, Symptom Frequency Score, Quality of Life Score, Social Limitation Score). The four summary scores are used to calculate the Overall Summary Score.

For more information, please refer to the KCCQ-12 Scoring Instructions document

provided by the STS/ACC TVT Registry.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001540
Code System: ACC NCDR
Short Name: F_KCCQ12_Overall

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: NUM
Precision: 5,2
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Data Source: Computed

Valid Range:

Parent/Child Validation

Element: 13845 Kansas City Cardiomyopathy

Questionnaire 12 Performed

Operator: Equal Value: Yes







ment: 12933	Follow-up Event Name	Technical Specification
Coding Instruction	Select from the list all of the clinical conditions, procedures, or re-admissions that occurred in	Code: 112000000795
County instruction.	the follow-up period	Code System: ACC NCDR
		Short Name: F_Condition_Event
Target Value:	N/A	Missing Data: Report
	A Follow-up - combination Name (12933), Occurred (14276) and Date (14277) - may only be entered/selected once	Harvested: Yes (BDS, TAVR, TMVF TMVrpr, TTVP)
	Chicles/Science once	Is Identifier: No
		Is Base Element: No
		Is Followup
		Element:
		Data Type: CD
		Precision:
		Selection Type: Single (Dynamic List)
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User

Selection	Definition	Source	Code	Code System
ASD Defect Closure due to Transseptal Catheterization	A procedure was required to close an atrial-septal defect as a result of the transseptal catheterization procedure.		112000001885	ACC NCDR
Atrial Fibrillation	Atrial fibrillation or flutter requiring treatment or prolonged hospitalization. Treatment includes initiation of a NEW/DIFFERENT medication therapy to address the arrhythmia; or a procedure/intervention to address the arrhythmia (cardioversion, permanent pacemaker/defibrillator, ablation, etc.).		49436004	SNOMED CT
Bleeding - Life Threatening	Life threatening or disabling bleeding is defined as: 1. Fatal bleeding OR 2. Bleeding in a critical area or organ, such as intracranial, intraspinal, intraocular, or pericardial necessitating pericardiocentesis, or intramuscular with compartment syndrome OR 3. Bleeding causing hypovolemic shock or severe hypotension requiring vasopressors or surgery OR 4. Overt source of bleeding with drop in hemoglobin of >=5 g/dl or whole blood or packed red blood cells (RBCs) transfusion >=4 U.		112000000459	ACC NCDR
Bleeding - Major	A major bleeding event, based on the 'Bleeding Academic Research Consortium' or BARC type 3a criteria is defined as: 1. Overt bleeding that is either associated with a drop in the hemoglobin level of at least 3.0 g/dL or requiring transfusion of 2 or 3 units of whole blood/RBC, or causing hospitalization or permanent injury, or requiring surgery AND 2. Does not meet VARC criteria of life-threatening or disabling bleeding.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001889	ACC NCDR
Cardiac Surgery or Intervention - Other Unplanned	The patient subsequently underwent cardiac surgery or a cath lab intervention that was unplanned. This does not include an intervention or procedure already identified as an adverse event in the TVT Registry (e.g. AV reintervention, other vascular surgery or intervention, pacemaker or ICD implant).		112000001892	ACC NCDR
COVID-19 Positive	The patient had a diagnosis of COVID-19, confirmed with a laboratory performed polymerase chain reaction (PCR) test. Notes: It is acceptable to code the diagnosis of COVID-19 based on testing that was NOT performed at your facility if it is accompanied by appropriate documentation in the medical record by the provider.		112000001982	ACC NCDR
	Code no if documentation ONLY included antibody testing (IgG).			
Deep Vein Thrombosis	Deep vein thrombosis (DVT) refers to the formation of	Office of the Surgeon General. (2008). The surgeon	128053003	SNOMED CT







Section: Follow-Up Ev	vents	Parent: Follow Up		
	· ·	general's call to action to prevent deep vein thrombosis and pulmonary embolism. Retrieved from https://www.ncbi.nlm.nih.gov/books/NBK44184/		
Device Embolization	The device became displaced from its initial implantation site so that it is no longer in its original position.	1	112000001324	ACC NCDR
Device Fracture	Partial or complete separation of any portion of the valve frame fractured into two or more parts. Do not code this event when there was a planned		112000001891	ACC NCDR
	bioprosthetic valve fracture (BVF) on a previously implanted bioprosthetic valve during the lab visit.			
Device Migration	Device migration of the prosthetic valve is x-ray confirmed movement of the valve from its initial implantation site such that there is a change in valve orientation within the aortic outflow track resulting in a new echocardiographic confirmed flow disturbance (pre- and post- filmed documentation). Note: Code device embolization if the device is outside of the outflow tract.		370512004	SNOMED CT
Device Thrombosis	Any thrombus attached to or near the valve that was implanted during the procedure that occludes part of the blood flow path, interferes with valve function, or is sufficiently large to warrant treatment.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001839	ACC NCDR
Device Related Event - Other	Indicate if an otherwise unspecified device-related event requiring unanticipated treatment occurred. This includes any delivery system related event.		112000001828	ACC NCDR
Dialysis (New Requirement)	Acute or worsening renal failure necessitating a new requirement for renal dialysis (renal dialysis includes hemodialysis and peritoneal dialysis). If a patient is on receiving continuous veno-venous hemofiltration (CVVH) as a result of renal failure (and not as treatment to remove fluid for heart failure), code yes.		100014076	ACC NCDR
Endocarditis	Confirmed diagnosis of endocarditis by blood culture and/or vegetation on or around a heart valve. This may include native tissue, ring or prosthetic valve involvement.	Society of Thoracic Surgeons (STS)	56819008	SNOMED CT
ICD	The patient developed a new dysrhythmia requiring insertion of an implantable cardioverter/defibrillator.		ACC-NCDR- ICD	ACC NCDR
Myocardial Infarction	A myocardial infarction (MI) is defined as an acute ischemic event that is associated with documented and clinically significant myocardial necrosis. The MI can be periprocedural (<72 hours after the procedure) or spontaneous (>72 hours after the index procedure). 1. Peri-procedural MI (<72 h after the index procedure) (a) New ischemic symptoms (e.g. chest pain or shortness of breath), or new ischemic signs (e.g. ventricular arrhythmias, new or worsening heart failure, new ST-segment changes, hemodynamic instability, new pathological Q waves in at least two contiguous leads, imaging evidence of new loss of viable myocardium or new wall motion abnormality) AND		22298006	SNOMED CT
	(b) Elevated cardiac biomarkers (preferable CK-MB) within 72 h after the indexprocedure, consisting of at least one sample post-procedure with a peak value exceeding 15 x as the upper reference limit for troponin or 5 x forCK-MB.* If cardiac biomarkers are increased at baseline (>99th percentile), a further increase in at least 50% post-procedure is required AND the peak value must exceed the previously stated limit.			
	Spontaneous MI (_72 h after the index procedure) any one of the following criteria:			
	(a) Detection of rise and/or fall of cardiac			

Effective for Patient Discharged January 01, 2021

biomarkers (preferably troponin) with at least one value above the 99th percentile URL, together with the evidence of myocardial ischemia with at least one of the following:

-Symptoms of ischemia







Section: Follow-Up Ev	vents	Parent: Follow Up		
	-ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] -New pathological Q-waves in at least two contiguous leads			
	-Imaging evidence of a new loss of viable myocardium or new wall motion abnormality			
	(b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood.			
	(c) Pathological findings of an acute myocardial infarction.			
PCI	A PCI is the placement of an angioplasty guide wire, balloon, or other device (e.g. stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary artery bypass graft for the purpose of mechanical coronary revascularization.		415070008	SNOMED CT
Permanent Pacemaker	The patient developed a new dysrhythmia requiring insertion of a permanent pacemaker.		449397007	SNOMED CT
Pulmonary Embolism	Intravascular migration of a venous thrombus to the pulmonary arterial circulation. A 'Proved Pulmonary Embolism' is proved by a positive pulmonary angiogram an unequivocally positive helical CT scan, a high-probability ventilation-perfusion scan, or autopsy.	Banovac, F., et al. Reporting Standards for Endovascular Treatment of Pulmonary Embolism. , Journal of Vascular Interventional Radiology 2010; 21:44–53	59282003	SNOMED CT
Readmission - (Non-Valve Related)	The patient has been readmitted to an acute care facility after discharge for a non-valve related reason.		112000001895	ACC NCDR
Readmission (Valve Related)	The patient has been readmitted to an acute care facility after discharge for a valve-related reason.		112000001894	ACC NCDR
Readmission - Cardiac (Not Heart Failure)	The patient has been readmitted to an acute care facility after discharge with a cardiac diagnosis (where the primary diagnosis is NOT heart failure).		112000001897	ACC NCDR
Readmission - Heart Failure	The patient has been readmitted to an acute care facility after discharge for the procedure with a diagnosis of heart failure.		112000001896	ACC NCDR
	The following criteria must be met for an event to be characterized as a heart failure readmission: 1. Hospitalization >=24 hours (including emergency room stay); 2. Clinical signs and/or symptoms of heart failure (including, but not limited to, new or worsening dyspnea, orthopnea, paroxysmal nocturnal dyspnea, increasing fatigue, worsening functional capacity or activity intolerance, or signs and/or symptoms of volume overload.); 3. Intravenous (e.g., diuretic or vasoactive therapy) or invasive (e.g., ultrafiltration, IABP, mechanical			
Readmission - Non-Cardiac	assistance) treatment for heart failure. The patient has been readmitted to an acute care		112000001898	ACC NCDR
Trought Troit Garage	facility after discharge for a non-cardiac related diagnosis or procedure.			,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
Reintervention - Aortic Valve	The patient returned to the operating room or cath lab for any aortic valve re-intervention.		112000001827	ACC NCDR
	Note: Please complete adjudication worksheet for every documented aortic valve reintervention, regardless of type of reintervention.			
Reintervention - Mitral Valve	The patient returned to the operating room or cath lab for any mitral valve re-intervention. Note: Please complete adjudication worksheet for every documented mitral valve reintervention,		112000001893	ACC NCDR
Reintervention - Tricuspid	regardless of type of reintervention.		112000001820	ACC NCDR
Reintervention - Tricuspid Valve	The patient returned to the operating room or cath lab for any tricuspid valve re-intervention.		112000001020	ACC NODR
	Note: Please complete adjudication worksheet for every documented tricuspid valve reintervention, regardless of type of reintervention.			
Single Leaflet Device	Single leaflet device attachment was documented in		112000001538	ACC NCDR







Section: Follow-Up Ev	ents	Parent: Follow Up		
Attachment	the medical record.			
Stroke - Ischemic	An ischemic stroke is an acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of infarction of central nervous system tissue.	Hicks KA, Tcheng JE, Bozkurt B, et al. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards (Writing Committee to Develop Cardiovascular Endpoints Data Standards). J Am Coll Cardiol. 2015;66 (4):403-469. doi:10.1016/j.jacc.2014.12.018.	422504002	SNOMED CT
Stroke - Hemorrhagic		,,	230706003	SNOMED CT
Stroke - Undetermined	A stroke of undetermined origin is defined as an acute episode of focal or global neurological dysfunction caused by presumed brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction but with insufficient information to allow categorization as ischemic or hemorrhagic.	Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart	230713003	SNOMED CT
Transient Ischemic Attack (TIA)	A transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.	Society for Thoracic Surgeons (STS)	266257000	SNOMED CT
Vascular Complication - Major	Major vascular complications include any of the following: 1. Any aortic dissection, aortic rupture, annulus rupture, left ventricle perforation, or new apical aneurysm/pseudo-aneurysm; 2. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneurysm, hematoma, irreversible nerve injury, compartment syndrome, percutaneous closure device failure) leading to death, life threatening or major bleeding*, visceral ischemia or neurological impairment; 3. Distal embolization (non-cerebral) from a vascular source requiring surgery or resulting in amputation or irreversible end-organ damage; 4. The use of unplanned endovascular or surgical intervention associated with death, major bleeding, visceral ischemia or neurological impairment; 5. Any new ischemia documented by patient symptoms, physical exam, and/or decreased or absent blood flow on lower extremity angiogram; 6. Surgery for access site-related nerve injury; 7. Permanent access site-related nerve injury. **Refers to VARC bleeding definitions Note: "ipsilateral lower extremity" was removed from #5 to have the ability to account for ischemia from any access site.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000000460	ACC NCDR
Vascular Complication - Minor	Minor vascular complications include any of the following: 1. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneuysms, hematomas, percutaneous closure device failure) not leading to death, life-threatening or major bleeding*, visceral ischemia or neurological impairment; 2. Distal embolization treated with embolectomy and/or thrombectomy and not resulting in amputation or irreversible end-organ damage; 3. Any unplanned endovascular stenting or unplanned surgical intervention not meeting the criteria for a major vascular complication; 4. Vascular repair or the need for vascular repair (via surgery, ultrasound-guided compression, transcathete embolization, or stent-graft). *Refers to VARC bleeding definitions		112000001823	ACC NCDR
Vascular Surgery or Intervention - Unplanned	The patient required unplanned vascular surgery or intervention to correct a bleeding complication or vascular related complication. Note: If a balloon angioplasty of the access site or access related sites is performed as a routine procedure to ensure adequate hemostasis of the site, then this would not qualify as an Unplanned Vascular Surgery or Intervention. However, if a balloon angioplasty is performed in an attempt to treat a		112000000467	ACC NCDR







Section: Follow-Up Events

Parent: Follow Up

bleeding or vascular access complication (i.e. bleeding at access site, dissection, stenosis, narrowing of vessel, etc.), then Unplanned Vascular Surgery or Intervention should be captured.

Element: 14276 Follow-Up Events Occurred

Coding Instruction: Indicate if the event occurred.

Target Value: Any occurrence on follow-up

Technical Specification

Code: 1000142378
Code System: ACC NCDR
Short Name: FupEvOccurred

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single

Is Identifier: No

Unit of Measure:

Default Value: Null

Usual Range:

Valid Range:

Data Source: User

Parent/Child Validation

Operator:

Value: Any Value

Element: 14277 Follow-Up Event Date

Coding Instruction: Indicate the date the event occurred.

Target Value: Any occurrence on follow-up

Technical Specification

Code: 1000142379
Code System: ACC NCDR
Short Name: FupEventDate
Missing Data: Report

Element: 12933 Follow-up Event Name

 $\textbf{Harvested:} \ \ \text{Yes (BDS, TAVR, TMVR,} \\$

TMVrpr, TTVP)

Is Base Element: No
Is Followup Yes
Element:
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 14276 Follow-Up Events Occurred

Operator: Equal Value: Yes







nent: 14385	Adjudication Event	Technical Specification
Coding Instruction:	Indicate the event being adjudicated.	Code: 112000001816 Code System: ACC NCDR
Target Value:	N/A	Short Name: F_AJ_AdjudEvent
Vendor Instruction:	An Adjudication - combination Event (14385) and Date (14386) - may only be entered/selected once	Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR TMVrpr, TTVP)
	The Adjudication Event Date (14386) / Adjudication Event Code (14385) must match with	Is Identifier: No
	Follow-Up Event Date (14277) / Follow-Up Event Code (12933)	Is Base Element: No
		Is Followup Element:
		Data Type: CD
		Precision:
		Selection Type: Single (Dynamic List)
		Unit of Measure:
		Default Value: Null
		Usual Range:
		Valid Range:
		Data Source: User

Follow Up Events - 1.3.6.1.4.1.19376.1.4.1.6.5.356

Selection	Definition	Source	Code	Code System
ASD Defect Closure due to Transseptal Catheterization	A procedure was required to close an atrial-septal defect as a result of the transseptal catheterization procedure.		112000001885	ACC NCDR
Atrial Fibrillation	Atrial fibrillation or flutter requiring treatment or prolonged hospitalization. Treatment includes initiation of a NEW/DIFFERENT medication therapy to address the arrhythmia; or a procedure/intervention to address the arrhythmia (cardioversion, permanent pacemaker/defibrillator, ablation, etc.).		49436004	SNOMED CT
Bleeding - Life Threatening	Life threatening or disabling bleeding is defined as: 1. Fatal bleeding OR 2. Bleeding in a critical area or organ, such as intracranial, intraspinal, intraocular, or pericardial necessitating pericardiocentesis, or intramuscular with compartment syndrome OR 3. Bleeding causing hypovolemic shock or severe hypotension requiring vasopressors or surgery OR 4. Overt source of bleeding with drop in hemoglobin of >=5 g/dl or whole blood or packed red blood cells (RBCs) transfusion >=4 U.		112000000459	ACC NCDR
Bleeding - Major	A major bleeding event, based on the 'Bleeding Academic Research Consortium' or BARC type 3a criteria is defined as: 1. Overt bleeding that is either associated with a drop in the hemoglobin level of at least 3.0 g/dL or requiring transfusion of 2 or 3 units of whole blood/RBC, or causing hospitalization or permanent injury, or requiring surgery AND 2. Does not meet VARC criteria of life-threatening or disabling bleeding.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001889	ACC NCDR
Cardiac Surgery or Intervention - Other Unplanned	The patient subsequently underwent cardiac surgery or a cath lab intervention that was unplanned. This does not include an intervention or procedure already identified as an adverse event in the TVT Registry (e.g. AV reintervention, other vascular surgery or intervention, pacemaker or ICD implant).		112000001892	ACC NCDR
COVID-19 Positive	The patient had a diagnosis of COVID-19, confirmed with a laboratory performed polymerase chain reaction (PCR) test. Notes: It is acceptable to code the diagnosis of COVID-19 based on testing that was NOT performed at your facility if it is accompanied by appropriate documentation in the medical record by the provider. Code no if documentation ONLY included antibody testing (ICC)		112000001982	ACC NCDR
	testing (IgG).	Office of the Surgeon General. (2008). The surgeon	128053003	SNOMED CT
Deep Vein Thrombosis	Deen vain thrombosic (DVI) refers to the termetion of			







Section: Follow-Up Ev	vent Information	Parent: Follow Up		
	· ·	general's call to action to prevent deep vein thrombosis and pulmonary embolism. Retrieved from https://www.ncbi.nlm.nih.gov/books/NBK44184/		
Device Embolization	The device became displaced from its initial implantation site so that it is no longer in its original position.	1	112000001324	ACC NCDI
Device Fracture	Partial or complete separation of any portion of the valve frame fractured into two or more parts. Do not code this event when there was a planned		112000001891	ACC NCDF
	bioprosthetic valve fracture (BVF) on a previously implanted bioprosthetic valve during the lab visit.			
Device Migration	Device migration of the prosthetic valve is x-ray confirmed movement of the valve from its initial implantation site such that there is a change in valve orientation within the aortic outflow track resulting in a new echocardiographic confirmed flow disturbance (pre- and post- filmed documentation). Note: Code device embolization if the device is outside of the outflow tract.		370512004	SNOMED C
Device Thrombosis	Any thrombus attached to or near the valve that was implanted during the procedure that occludes part of the blood flow path, interferes with valve function, or is sufficiently large to warrant treatment.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	112000001839	ACC NCDF
Device Related Event - Other	Indicate if an otherwise unspecified device-related event requiring unanticipated treatment occurred. This includes any delivery system related event.		112000001828	ACC NCDF
Dialysis (New Requirement)	Acute or worsening renal failure necessitating a new requirement for renal dialysis (renal dialysis includes hemodialysis and peritoneal dialysis). If a patient is on receiving continuous veno-venous hemofiltration (CVVH) as a result of renal failure (and not as treatment to remove fluid for heart failure), code yes.		100014076	ACC NCDF
Endocarditis	Confirmed diagnosis of endocarditis by blood culture and/or vegetation on or around a heart valve. This may include native tissue, ring or prosthetic valve involvement.	Society of Thoracic Surgeons (STS)	56819008	SNOMED CT
ICD	The patient developed a new dysrhythmia requiring insertion of an implantable cardioverter/defibrillator.		ACC-NCDR- ICD	ACC NCDF
Myocardial Infarction	A myocardial infarction (MI) is defined as an acute ischemic event that is associated with documented and clinically significant myocardial necrosis. The MI can be periprocedural (<72 hours after the procedure) or spontaneous (>72 hours after the index procedure). 1. Peri-procedural MI (<72 h after the index procedure)	•	22298006	SNOMED CT
	(a) New ischemic symptoms (e.g. chest pain or shortness of breath), or new ischemic signs (e.g. ventricular arrhythmias, new or worsening heart failure, new ST-segment changes, hemodynamic instability, new pathological Q waves in at least two contiguous leads, imaging evidence of new loss of viable myocardium or new wall motion abnormality) AND			
	(b) Elevated cardiac biomarkers (preferable CK-MB) within 72 h after the indexprocedure, consisting of at least one sample post-procedure with a peak value exceeding 15 x as the upper reference limit for troponin or 5 x forCK-MB.* If cardiac biomarkers are increased at baseline (>99th percentile), a further increase in at least 50% post-procedure is required AND the peak value must exceed the previously stated limit.			
	2. Spontaneous MI (_72 h after the index procedure) any one of the following criteria:			

- (a) Detection of rise and/or fall of cardiac biomarkers (preferably troponin) with at least one value above the 99th percentile URL, together with the evidence of myocardial ischemia with at least one of the following:
- -Symptoms of ischemia







ent Information	Parent: Follow Up		
-ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] -New pathological Q-waves in at least two contiguous leads -Imaging evidence of a new loss of viable myocardium			
or new wall motion abnormality (b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBRB, and/or evidence of			
fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood.			
(c) Pathological findings of an acute myocardial infarction.			
	National Cardiovascular Data Registry (NCDR)	415070008	SNOMED CT
The patient developed a new dysrhythmia requiring insertion of a permanent pacemaker.		449397007	SNOMED CT
Intravascular migration of a venous thrombus to the pulmonary arterial circulation. A 'Proved Pulmonary Embolism' is proved by a positive pulmonary angiogram, an unequivocally positive helical CT scan, a high-	Banovac, F., et al. Reporting Standards for Endovascular Treatment of Pulmonary Embolism. Journal of Vascular Interventional Radiology 2010; 21:44–53	59282003	SNOMED CT
The patient has been readmitted to an acute care		112000001895	ACC NCDR
The patient has been readmitted to an acute care facility after discharge for a valve-related reason.		112000001894	ACC NCDF
The patient has been readmitted to an acute care facility after discharge with a cardiac diagnosis (where the primary diagnosis is NOT heart failure).		112000001897	ACC NCDR
The patient has been readmitted to an acute care facility after discharge for the procedure with a diagnosis of heart failure.		112000001896	ACC NCDR
The following criteria must be met for an event to be characterized as a heart failure readmission: 1. Hospitalization >=24 hours (including emergency room stay); 2. Clinical signs and/or symptoms of heart failure (including, but not limited to, new or worsening dyspage, orthogoga, parroysmal nocturnal dyspage).			
increasing fatigue, worsening functional capacity or activity intolerance, or signs and/or symptoms of volume overload.); 3. Intravenous (e.g., diuretic or vasoactive therapy) or			
invasive (e.g., ultrafiltration, IABP, mechanical assistance) treatment for heart failure.			
The patient has been readmitted to an acute care facility after discharge for a non-cardiac related diagnosis or procedure.		112000001898	ACC NCDR
The patient returned to the operating room or cath lab for any aortic valve re-intervention.		112000001827	ACC NCDR
Note: Please complete adjudication worksheet for every documented aortic valve reintervention, regardless of type of reintervention.			
The patient returned to the operating room or cath lab for any mitral valve re-intervention.		112000001893	ACC NCDR
Note: Please complete adjudication worksheet for every documented mitral valve reintervention, regardless of type of reintervention.			
The patient returned to the operating room or cath lab for any tricuspid valve re-intervention.		112000001820	ACC NCDR
	-ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] -New pathological Q-waves in at least two contiguous leads -Imaging evidence of a new loss of viable myocardium or new wall motion abnormality (b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood. (c) Pathological findings of an acute myocardial infarction. A PCI is the placement of an angioplasty guide wire, balloon, or other device (e.g. stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary artery bypass graft for the purpose of mechanical coronary revascularization. The patient developed a new dysrhythmia requiring insertion of a permanent pacemaker. Intravascular migration of a venous thrombus to the pulmonary arterial circulation. A 'Proved Pulmonary Embolism' is proved by a positive pulmonary angiogram an unequivocally positive helical CT scan, a high-probability ventilation-perfusion scan, or autopsy. The patient has been readmitted to an acute care facility after discharge for a non-valve related reason. The patient has been readmitted to an acute care facility after discharge for a valve-related reason. The patient has been readmitted to an acute care facility after discharge for a valve-related reason. The patient has been readmitted to an acute care facility after discharge for searchy in a cardiac diagnosis of heart failure. The patient has been readmitted to an acute care facility after discharge for a valve-related reason. The patient has been readmited to an acute care facility after discharge for searchy in a cardiac diagnosis of heart failure. The patient returned to the operating room or cath lab for any artic valve	ECG changes indicative of new ischemia [new ST-T changes or new left bundle branch block (LBBB)] New pathological O-waves in at least two contiguous leads I-maging evidence of a new loss of viable myocardium or new wall motion abnormality (b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial ischemia, and accompanied by presumably new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples could be obtained, or at a time before the appearance of cardiac biomarkers in the blood. (c) Pathological findings of an acute myocardial infarction. A PCI is the placement of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary artery bypass graft for the purpose of mechanical coronary revascularization. The patient developed a new dysrhythmia requiring insention of a permanent pacemaker. The patient developed a new dysrhythmia requiring insention of a permanent pacemaker. The patient developed a new dysrhythmia requiring insention of a permanent pacemaker. The patient developed to a new dysrhythmia requiring insention of a permanent pacemaker. The patient developed by a positive plumonary genome and probability were interchanged for a new-lave related reason. The patient has been readmitted to an acute care facility after discharge for a non-valve related reason. The patient has been readmitted to an acute care facility after discharge for a non-valve related reason. The patient has been readmitted to an acute care facility after discharge for a non-valve related reason. The patient has been readmitted to an acute care facility after discharge for a point procedure with a diagnosis of heart failure. The patient has been readmitted to an acute care facility after discharge for a point for any and for symptoms of volume overload; 3. Intravenous (e.g., diuriter	ECG changes indicative of new ischemia (new ST-T changes or new let bundle branch block (LBBB) reverse plant logical Cwaves in a less test two configurus changes or new let bundle branch block (LBBB) reverse plant logical Cwaves in a less test two configurus changes gevidence of a new loss of viable myocardium or new wall motion abnormality (b) Sudden, unexpected cardiac death, involving cardiac arrest, often with symptoms suggestive of myocardial inchemia, and accompanies by presumbly new ST elevation, or new LBBB, and/or evidence of fresh thrombus by coronary angiography and/or at autopsy, but death occurring before blood samples of cardiac blomarkers in the blood. (c) Pathological findings of an acute myocardial inflaration. AFC is the pleasment of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy business) and the pleasment of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy business) and the pleasment of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy business) and the plant of the plant developed a new dysrhythmia requiring insertion of a permanent pacemater. Intravascular migration of a vanous thrombus to the purpose of mechanical coronary vesticularization. The patient developed a new dysrhythmia requiring insertion of a permanent pacemater. Intravascular migration of a vanous thrombus to the purpose of mechanical coronary vesticularization. 2144–53 2144–53 11200001895 11200001895 11200001895 11200001895 11200001896 11200001896 11200001896 11200001896 11200001897 11200001896 11200001897 11200001897 11200001897 11200001898 112







Section: Follow-Up Ev	ent miormation	Parent: Follow Up		
Attachment	the medical record.			
Stroke - Ischemic	An ischemic stroke is an acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of infarction of central nervous system tissue.	Hicks KA, Tcheng JE, Bozkurt B, et al. 2014 ACC/AHA Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards (Writing Committee to Develop Cardiovascular Endpoints Data Standards). J Am Coll Cardiol. 2015;66 (4):403-469. doi:10.1016/j.jacc.2014.12.018.	422504002	SNOMED CT
Stroke - Hemorrhagic		,,	230706003	SNOMED CT
Stroke - Undetermined	A stroke of undetermined origin is defined as an acute episode of focal or global neurological dysfunction caused by presumed brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction but with insufficient information to allow categorization as ischemic or hemorrhagic.	Key Data Elements and Definitions for Cardiovascular Endpoint Events in Clinical Trials: A Report of the American College of Cardiology/American Heart	230713003	SNOMED CT
Transient Ischemic Attack (TIA)	A transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.	Society for Thoracic Surgeons (STS)	266257000	SNOMED CT
Vascular Complication - Major	Major vascular complications include any of the following: 1. Any aortic dissection, aortic rupture, annulus rupture, left ventricle perforation, or new apical aneurysm/pseudo-aneurysm; 2. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneurysm, hematoma, irreversible nerve injury, compartment syndrome, percutaneous closure device failure) leading to death, life threatening or major bleeding*, visceral ischemia or neurological impairment; 3. Distal embolization (non-cerebral) from a vascular source requiring surgery or resulting in amputation or irreversible end-organ damage; 4. The use of unplanned endovascular or surgical intervention associated with death, major bleeding, visceral ischemia or neurological impairment; 5. Any new ischemia documented by patient symptoms, physical exam, and/or decreased or absent blood flow on lower extremity angiogram; 6. Surgery for access site-related nerve injury; 7. Permanent access site-related nerve injury. **Refers to VARC bleeding definitions Note: "ipsilateral lower extremity" was removed from #5 to have the ability to account for ischemia from any access site.	Updated Standardized Endpoint Definitions for Transcatheter Aortic Valve Implantation (JACC, 2012, Vol 60, No 15)	11200000460	ACC NCDR
Vascular Complication - Minor	Minor vascular complications include any of the following: 1. Access site or access-related vascular injury (dissection, stenosis, perforation, rupture, arteriovenous fistula, pseudoaneuysms, hematomas, percutaneous closure device failure) not leading to death, life-threatening or major bleeding*, visceral ischemia or neurological impairment; 2. Distal embolization treated with embolectomy and/or thrombectomy and not resulting in amputation or irreversible end-organ damage; 3. Any unplanned endovascular stenting or unplanned surgical intervention not meeting the criteria for a major vascular complication; 4. Vascular repair or the need for vascular repair (via surgery, ultrasound-guided compression, transcatheterembolization, or stent-graft). *Refers to VARC bleeding definitions		112000001823	ACC NCDR
Vascular Surgery or Intervention - Unplanned	The patient required unplanned vascular surgery or intervention to correct a bleeding complication or vascular related complication. Note: If a balloon angioplasty of the access site or access related sites is performed as a routine procedure to ensure adequate hemostasis of the site, then this would not qualify as an Unplanned Vascular Surgery or Intervention. However, if a balloon angioplasty is performed in an attempt to treat a		112000000467	ACC NCDR







Section: Follow-Up Event Information

Parent: Follow Up

bleeding or vascular access complication (i.e. bleeding at access site, dissection, stenosis, narrowing of vessel, etc.), then Unplanned Vascular Surgery or Intervention should be captured.

Element: 14386 Adjudication Event Date

Coding Instruction: Indicate the date the clinical event being adjudicated occurred.

Target Value: N/A

Vendor Instruction: The Adjudication Event Date (14386) / Adjudication Event Code (14385) must match with

Follow-Up Event Date (14277) / Follow-Up Event Code (12933)

Technical Specification

Code: 112000001816
Code System: ACC NCDR
Short Name: F_AJ_EventDate

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Base Element: No
Is Followup
Element: Yes
Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve Element: 14385 Adjudication Event

Operator: Equal

Value: Readmission - Heart Failure Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)
Element: 14385 Adjudication Event
Operator: Equal

perator: Equal

Value: Reintervention - Tricuspid Valve Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined







Section: Follow-Up Event Information

Parent: Follow Up

Element: 14387 Adjudication Status

Coding Instruction: Indicate whether the patient was alive or deceased on the date the adjudication was

performed

Target Value: N/A

Vendor Instruction: Adjudication Status (14387) as 'Deceased' must be answered only once in follow-up episode.

Technical Specification
Code: 112000001817
Code System: ACC NCDR
Short Name: F_AJ_Status
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null

Usual Range: Valid Range: Data Source: User

Adjudication Life Status - 1.3.6.1.4.1.19376.1.4.1.6.5.726

 Selection
 Definition
 Source
 Code
 Code System

 Alive
 438949009
 SNOMED CT

 Deceased
 20
 HL7 Discharge disposition

Element: 14388 Adjudication Date of Death

Coding Instruction: Indicate the date the patient was declared dead.

Target Value: N/A

Vendor Instruction: Adjudication Date of Death (14388) must be Greater than or Equal to Adjudication Event Date

(14386)

Technical Specification

Code: 399753006
Code System: SNOMED CT
Short Name: F_AJ_DeathDate
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14387 Adjudication Status

Operator: Equal Value: Deceased







Section: Follow-Up Event Information

Parent: Follow Up

Element: 14463 Follow Up Clinical Comments

Coding Instruction: Provide information and details that may assist in assessing the event(s) being adjudicated.

Target Value: N/A

Technical Specification
Code: 423016009
Code System: SNOMED CT
Short Name: AJ_CommentsFU

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element: Yes
Data Type: ST
Precision: 1000
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Data Source: User







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14389 Symptom Onset Date

Coding Instruction: Indicate the date of symptom onset of the neurologic deficit.

Target Value: N/A

Technical Specification

Code: 112000000125
Code System: ACC NCDR
Short Name: F_AJ_SxOnset

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Data Type: DT
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)
Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined

----- AND -----

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14390 Neurologic Deficit with Rapid Onset

Coding Instruction: Indicate if the patient had a sudden onset of a focal or global neurologic deficit (regardless of the duration of symptoms) with at least one of the following present: change in level of

consciousness, hemiplegia, hemiparesis, numbness or sensory loss affecting one side of the body, dysphasia or aphasia, hemianopia, amaurosis fugax, other neurological signs or

symptoms consistent with a stroke.

Target Value: N/A

Technical Specification

Code: 264552009
Code System: SNOMED CT
Short Name: F_AJ_NeuroDef
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined

Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14391 Neurologic Deficit Clinical Presentation

Coding Instruction: Indicate the clinical presentation of the neurologic deficit.

Target Value: N/A

Technical Specification

Code: 264552009 Code System: SNOMED CT

Short Name: F_AJ_NeuroClinPresent

Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

 Is Base Element:
 No

 Is Followup Element:
 Yes

 Data Type:
 CD

 Precision:
 Selection Type:
 Single

Is Identifier: No

Selection Type: Singl Unit of Measure: Default Value: Null Usual Range: Valid Range: Data Source: User

Parent/Child Validation

Element: 14390 Neurologic Deficit with Rapid

Onset
Operator: Equal

Value: Yes

Neurologic Deficit Clinical Presentation - 1.3.6.1.4.1.19376.1.4.1.6.5.716

Selection	Definition	Source	Code	Code System
TIA or Stroke (CVA)			100014109	ACC NCDR
Non Stroke Neurologic	Deficit		112000001860	ACC NCDR

Element: 14392 Neurologic Symptom Duration Greater Than or Equal to 24 hours

Coding Instruction: Indicate if the duration of the neurologic symptoms lasted >= 24 hours.

Target Value: N/A

Technical Specification

Code: 308921004 Code System: SNOMED CT

Short Name: F_AJ_NeuroSymptDuration

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No

Is Base Element: No
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14391 Neurologic Deficit Clinical

Presentation

Operator: Equal

Value: TIA or Stroke (CVA)







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14393 Brain Imaging Performed

Coding Instruction: Indicate if neuroimaging such as CT, MRI, cerebral angiography was performed.

Target Value: N/A

Technical Specification

Code: 441986001
Code System: SNOMED CT
Short Name: F_AJ_BrainImag

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14391 Neurologic Deficit Clinical

Presentation

Data Source: User

Operator: Equal

Value: TIA or Stroke (CVA)

Element: 14394 Brain Imaging Type

Coding Instruction: Indicate the type of neuroimaging performed.

Target Value: N/A

Technical Specification

Code: 441986001 Code System: SNOMED CT

Short Name: F_AJ_BrainImageType

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Yes
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation
Element: 14393 Brain Imaging Performed

Operator: Equal Value: Yes

Imaging Type - 1.3.6.1.4.1.19376.1.4.1.6.5.417

Selection	Definition	Source	Code	Code System
Computed Tomography			77477000	SNOMED CT
Computed Tomography wit Contrast	h		112000001861	ACC NCDR
Magnetic Resonance Imagi	ng		113091000	SNOMED CT
Magnetic Resonance Imagi with Contrast	ng		51619007	SNOMED CT
Other Imaging			112000001862	ACC NCDR







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14395 Brain Imaging Findings

Coding Instruction: Indicate the type of deficit found as a result of the neuroimaging study. Hemorrhage includes

intraparenchymal, intraventricular and epidural hemorrhages.

Target Value: N/A

Technical Specification
Code: 112000001979
Code System: ACC NCDR
Short Name: F_Bl_Find

Missing Data: Report
Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14393 Brain Imaging Performed

Operator: Equal Value: Yes

Brain Imaging Finding - 1.3.6.1.4.1.19376.1.4.1.6.5.717

Selection	Definition	Source	Code	Code System
Infarct	Neuroimaging evidence of CNS infarction in the corresponding vascular territory (brain, spinal cord, or retinal cell death), with or without hemorrhage.	Adapted from: Lansky, A.J., et al. Proposed Standardized Neurological Endpoints for Cardiovascular Clinical Trials (An Academic Research Consortium Initiative) JACC 2017, 69 (6): 679-690	55641003	SNOMED CT
Hemorrhage	Neuroimaging evidence of central nervous system (CNS) hemorrhage within the brain parenchyma, subarachnoid space, ventricular system, spinal cord, or retina that is not caused by trauma.	Adapted from: Lansky, A.J., et al. Proposed Standardized Neurological Endpoints for Cardiovascular Clinical Trials (An Academic Research Consortium Initiative) JACC 2017, 69 (6): 679-690	50960005	SNOMED CT
No Deficit			100001231	ACC NCDR







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14396 Event Related Sequelae

Coding Instruction: Indicate the sequelae related to the stroke or TIA.

Target Value: N/A

Technical Specification

Code: 362977000
Code System: SNOMED CT
Short Name: F_Adj_ERS
Missing Data: Report

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Multiple Unit of Measure:

Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14391 Neurologic Deficit Clinical

Presentation
Operator: Equal

Value: TIA or Stroke (CVA)

Event Related Sequelae - 1.3.6.1.4.1.19376.1.4.1.6.5.737

Selection	Definition	Source	Code	Code System
Death			419620001	SNOMED CT
Permanent Vegetativ	ve State		723151005	SNOMED CT
Altered Consciousne	ess		3006004	SNOMED CT
Blindness			193699007	SNOMED CT
Aphasia			87486003	SNOMED CT
Loss of Motor Functi	ion		112000001936	ACC NCDR
Loss of Sensory Fur	nction		33653009	SNOMED CT
Facial Paralysis			280816001	SNOMED CT
Prolonged Length of	Stay		112000001937	ACC NCDR
Other			100000351	ACC NCDR



Element: 14420



Full Specifications Data Dictionary v3.0



Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Discharge Location After Event Coding Instruction: Indicate the discharge location after the stroke or TIA.

Target Value: N/A

Technical Specification Code: 75528-0 Code System: LOINC Short Name: F_AJ_DLAE

Missing Data: Report Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr)

Is Base Element: No Is Followup Element: Data Type: CD Precision: Selection Type: Single

Is Identifier: No

Unit of Measure: Default Value: Null **Usual Range:** Valid Range: Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 14387 Adjudication Status

Operator: Equal Value: Alive

----- AND -----Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

AND

Operator: Equal Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Discharge Location - 1.3.6.1.4.1.19376.1.4.1.6.5.41

Selection	Definition	Source	Code	Code System
Home			01	HL7 Discharge disposition
Skilled Nursing Facility	Skilled nursing facilities (SNF) are typically sub-ac programs used for longer anticipated length of sta		03	HL7 Discharge disposition
	Note: Sometimes SNFs may have acute rehabilita beds within their facility. If the patient is discharg a SNF for acute rehab (requiring a higher level of code "extended care/TCU/rehab".	ed to		
Extended Care/TCU/Rehab	An extended care unit, transitional care unit or rel unit typically provides a high level of intensive the as well as specialized nursing and physician care discharge setting may also be called subacute ca- long term acute care (LTACH).	rapy This	62	HL7 Discharge disposition
Other Acute Care Hospital			02	HL7 Discharge disposition
Left Against Medical Advice (AMA)	The patient was discharged or eloped against me advice.	dical	07	HL7 Discharge disposition
Other Discharge Location			100001249	ACC NCDR







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14422 Patient Discharged to Prior Place of Living

Coding Instruction: Indicate if the patient was discharged to their prior place of living.

Target Value: N/A

Technical Specification

Code: 112000001882
Code System: ACC NCDR
Short Name: F_AJ_PriorLiving

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: BL
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined
Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Floment: 14397 Adjudication Status

Element: 14387 Adjudication Status

Operator: Equal Value: Alive

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: TAVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR







Section: Follow-Up Stroke or TIA

Parent: Follow-Up Event Information

Element: 14397 Stroke Diagnosed During Autopsy

Coding Instruction: Indicate if the stroke was diagnosed during autopsy.

Target Value: N/A

Technical Specification

Code: 5605004

Code System: SNOMED CT

Short Name: F_AJ_AutDxStroke

Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Pecsion:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Hemorrhagic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Ischemic

Element: 14385 Adjudication Event

Operator: Equal

Value: Stroke - Undetermined

Element: 14385 Adjudication Event

Operator: Equal

Value: Transient Ischemic Attack (TIA)

Element: 14387 Adjudication Status

Operator: Equal

Value: Deceased

Boolean with Information Not Available - 1.3.6.1.4.1.19376.1.4.1.6.5.718

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR
Information Not Available			112000001866	ACC NCDR







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14398 Aortic Valve Reintervention Type

Coding Instruction: Indicate the type of aortic valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: F_AJ_ReIntType

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14385 Adjudication Event

Data Source: User

Operator: Equal

Value: Reintervention - Aortic Valve

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacement	Surgical Replacement		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replacement			112000001875	ACC NCDR
Balloon Valvuloplasty	Balloon Valvuloplasty		112000001469	ACC NCDR
Leaflet Clip Procedure	e		112000001778	ACC NCDR
Paravalvular Leak Clo	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14399 Aortic Valve Reintervention Primary Indication

Coding Instruction: Indicate the primary indication for the reintervention. If more than one indication is present,

code the indication the operator feels has the highest significance.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: F_AJ_PrimaryInd

Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: No
Is Followup Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation			40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14400 Aortic Valve Regurgitation

Coding Instruction: Indicate the highest level of aortic regurgitation prior to the aortic valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001869
Code System: ACC NCDR
Short Name: F_AJ_AlSev
Missing Data: Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: No
Is Followup Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Aortic Valve

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TAVR

----- AND -----

Element: 14399 Aortic Valve Reintervention

Primary Indication

Operator: Equal

Value: Regurgitation

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

Selection	Definition	Source	Code	Code System
None	'		112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14403 Paravalvular Aortic Regurgitation

Coding Instruction: Indicate the highest severity of paravalvular aortic regurgitation prior to the aortic valve

reintervention.

Note: If trace/trivial is documented, code "none".

Target Value: N/A

Technical Specification

 Code:
 112000001428

 Code System:
 ACC NCDR

 Short Name:
 F_AJ_PVSev

 Missing Data:
 Report

Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range:

Data Source: User
Parent/Child Validation

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14401 Central Aortic Regurgitation

Coding Instruction: Indicate the highest severity of central aortic regurgitation prior to the aortic valve

reintervention

Note: If trace/trivial is documented, code "none".

Target Value: N/A

Technical Specification

Code: 112000001433
Code System: ACC NCDR
Short Name: F_AJ_CenSev

Missing Data: Report Harvested: Yes (TAVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:

Default Value: Null Usual Range: Valid Range:

Data Source: User

Parent/Child Validation

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Mild

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Moderate

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Severe

Element: 14400 Aortic Valve Regurgitation

Operator: Equal Value: Trace/Trivial

Valve Regurgitation Severity 4 - 1.3.6.1.4.1.19376.1.4.1.6.5.768

Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR

Element: 14402 Aortic Valve Area

Coding Instruction: Indicate the smallest aortic valve area (in cm squared).

Target Value: N/A

Technical Specification

Code: 112000001280
Code System: ACC NCDR
Short Name: F_AJ_AVA
Missing Data: Report
Harvested: Yes (TAVR)

Is Base Element: No
Is Followup
Element:
Data Type: PQ
Precision: 3,2
Selection Type: Single
Unit of Measure: cm2
Default Value: Null

Is Identifier: No

Usual Range: 0.20 - 4.00 cm2 **Valid Range:** 0.05 - 5.00 cm2

Data Source: User

Parent/Child Validation

Element: 14399 Aortic Valve Reintervention

Primary Indication

Operator: Equal Value: Stenosis







Section: Follow-Up AV Re-Intervention

Parent: Follow-Up Event Information

Element: 14404 Aortic Valve Mean Gradient

Coding Instruction: Indicate the aortic valve mean gradient in mm Hg.

Target Value: N/A

Technical Specification

 Code:
 112000001398

 Code System:
 ACC NCDR

 Short Name:
 F_AJ_AVG

 Missing Data:
 Report

Harvested: Yes (BDS, TAVR)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mm[Hg]
Default Value: Null
Usual Range: 5 - 50 mm[

Usual Range: 5 - 50 mm[Hg]
Valid Range: 0 - 200 mm[Hg]

Data Source: User

Parent/Child Validation

Element: 14399 Aortic Valve Reintervention

Primary Indication

Operator: Equal Value: Stenosis







Section: Follow-Up MV Re-Intervention

Parent: Follow-Up Event Information

Element: 14405 Mitral Valve Reintervention Type

Coding Instruction: Indicate the type of mitral valve reintervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: F_AJ_MVReinType

Missing Data: Report

Is Identifier: No

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Base Element: No
Is Followup Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacemen	t		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replac	cement		112000001875	ACC NCDR
Balloon Valvuloplasty	/		112000001469	ACC NCDR
Leaflet Clip Procedur	e		112000001778	ACC NCDR
Paravalvular Leak Cl	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: Follow-Up MV Re-Intervention

Parent: Follow-Up Event Information

Element: 14406 Mitral Valve Reintervention Indication

Coding Instruction: Indicate the primary indication for the reintervention. If more than one indication is present,

code the indication the operator feels has the highest significance.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: F_AJ_MVReintInd

Missing Data: Report

Harvested: Yes (BDS, TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Element: CD
Precision: Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Mitral Valve

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation	,	·	40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: Follow-up Readmission

Parent: Follow-Up Event Information

Element: 14380 Hospitalization Greater Than or Equal to 24 Hours

Coding Instruction: Indicate if the heart failure readmission required the patient to be hospitalized with treatment in

any inpatient unit for at least 24 hours, including emergency department or observation stay.

Target Value: N/A

Technical Specification

Code: 1000142363
Code System: ACC NCDR
Short Name: F_AJ_Hospital

Missing Data: Report
Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Readmission - Heart Failure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Boolean with Information Not Available - 1.3.6.1.4.1.19376.1.4.1.6.5.718

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR
Information Not Avail	able		112000001866	ACC NCDR







Section: Follow-up Readmission

Parent: Follow-Up Event Information

Element: 14381 Clinical Signs or Symptoms of Heart Failure

Coding Instruction: Indicate if the patient had clinical signs and/or symptoms of heart failure, including new or

worsening dyspnea, orthopnea, paroxysmal nocturnal dyspnea, increasing fatigue, worsening functional capacity or activity intolerance, or signs and/or symptoms of volume overload.

Target Value: N/A

Technical Specification

Code: 100014007 Code System: ACC NCDR Short Name: F_AJ_SSHF

Missing Data: Report
Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup
Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Parent/Child Validation

Element: 14385 Adjudication Event

Data Source: User

Operator: Equal

Value: Readmission - Heart Failure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Boolean with Information Not Available - 1.3.6.1.4.1.19376.1.4.1.6.5.718

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR
Information Not Available			112000001866	ACC NCDR







Section: Follow-up Readmission

Parent: Follow-Up Event Information

Element: 14382 IV or Invasive Treatment Required

Coding Instruction: Indicate if the patient had signs and symptoms of heart failure that resulted in intravenous (e.g,

diuretic or vasoactive therapy) or invasive (e.g., ultrafiltration, IABP, mechanical assistance)

treatment for heart failure.

Target Value: N/A

Technical Specification

Code: 112000001867
Code System: ACC NCDR
Short Name: F_AJ_HFTreatment

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr)

Is Identifier: No
Is Base Element: No
Is Followup Element: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Readmission - Heart Failure

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVR

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal Value: TMVr

Boolean with Information Not Available - 1.3.6.1.4.1.19376.1.4.1.6.5.718

Selection	Definition	Source	Code	Code System
No			100013073	ACC NCDR
Yes			100013072	ACC NCDR
Information Not Avail	able		112000001866	ACC NCDR







Section: Follow-Up Tricuspid Valve Re-Intervention

Parent: Follow-Up Event Information

Element: 14408 Tricuspid Valve Reintervention Type

Coding Instruction: Indicate the type of tricuspid valve re-intervention.

Target Value: N/A

Technical Specification

Code: 112000001868
Code System: ACC NCDR
Short Name: F_AJ_TVReIn

Missing Data: Report Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Tricuspid Valve
----- AND ------

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Reintervention Type - 1.3.6.1.4.1.19376.1.4.1.6.5.719

Selection	Definition	Source	Code	Code System
Surgical Replacemen	t		112000001872	ACC NCDR
Surgical Repair			112000001871	ACC NCDR
Transcatheter Replac	cement		112000001875	ACC NCDR
Balloon Valvuloplasty	/		112000001469	ACC NCDR
Leaflet Clip Procedure	e		112000001778	ACC NCDR
Paravalvular Leak Clo	osure		112000001916	ACC NCDR
Other Transcatheter			112000001873	ACC NCDR
Intervention				







Section: Follow-Up Tricuspid Valve Re-Intervention

Parent: Follow-Up Event Information

Element: 14409 Tricuspid Valve Reintervention Primary Indication

Coding Instruction: Indicate the primary indication for the tricuspid valve re-intervention.

Target Value: N/A

Technical Specification

Code: 112000001825
Code System: ACC NCDR
Short Name: F_AJ_TVInd
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup Element: Yes
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14385 Adjudication Event

Operator: Equal

Value: Reintervention - Tricuspid Valve
----- AND ------

Element: 13705 Transcatheter Valve Therapy

Reference Procedure Type

Operator: Equal

Value: Tricuspid Valve Procedure

Valve Reintervention Indication - 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selection	Definition	Source	Code	Code System
Regurgitation			40445007	SNOMED CT
Stenosis			44241007	SNOMED CT
Device Embolization			112000001324	ACC NCDR
Device Fracture			112000001891	ACC NCDR
Device Migration			370512004	SNOMED CT
Endocarditis			56819008	SNOMED CT
Paravalvular Leak			234184000	SNOMED CT
Device Thrombosis			112000001839	ACC NCDR
Valve Injury			762610001	SNOMED CT
Other			100000351	ACC NCDR







Section: Follow-Up Tricuspid Valve Re-Intervention

Parent: Follow-Up Event Information

Element: 14410 Tricuspid Valve Regurgitation

Coding Instruction: Indicate the severity of tricuspid valve regurgitation.

Target Value: N/A

Technical Specification

Code: 111287006
Code System: SNOMED CT
Short Name: F_AJ_TR
Missing Data: Report
Harvested: Yes (TTVP)

Is Identifier: No
Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Parent/Child Validation

Element: 14409 Tricuspid Valve Reintervention

Primary Indication

Operator: Equal

Value: Regurgitation

Valve Regurgitation Severity - 1.3.6.1.4.1.19376.1.4.1.6.5.767

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Selection	Definition	Source	Code	Code System
None			112000001910	ACC NCDR
Trace/Trivial			112000001911	ACC NCDR
Mild			112000000380	ACC NCDR
Moderate			112000000381	ACC NCDR
Severe			112000000382	ACC NCDR







Section: Follow-Up Medications

Parent: Follow Up

Element: 11990 Follow-Up Medications Code

Coding Instruction: Indicate the assigned identification number associated with the medications the patient was

prescribed or received.

Note(s):

The medication(s) collected in this field are controlled by the Medication Master file. This file is maintained by the NCDR and will be made available on the internet for downloading and importing/updating into your application. Each medication in the Medication Master file is assigned to a value set. The value set is used to separate procedural medications from medications prescribed at discharge. The separation of these medications is depicted on the

data collection form.

Target Value: N/A

Technical Specification

Code: 100013057
Code System: ACC NCDR
Short Name: F_MedID
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:

Is Identifier: No

Selection Type: Single (Dynamic List)

Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Follow-up Medication - 2.16.840.1.113883.3.3478.6.5.203

Selection Definition	Source	Code	Code System
Angiotensin Converting Enzyme Inhibitor		41549009	SNOMED CT
		07000000	ONOMEDOT
Aldosterone Antagonist		372603003	SNOMED CT
Direct thrombin inhibitor		414010005	SNOMED CT
Warfarin		11289	RxNorm
Aspirin		1191	RxNorm
Angiotensin II Receptor Blocker		372913009	SNOMED CT
Beta Blocker		33252009	SNOMED CT
Diuretics Not Otherwise		112000001417	ACC NCDR
Specified			
Loop Diuretics		29051009	SNOMED CT
Thiazides		372747003	SNOMED CT
Direct Factor Xa Inhibitor		112000000696	ACC NCDR
P2Y12 Antagonist		112000001003	ACC NCDR







Section: Follow-Up Medications

Parent: Follow Up

Element: 13696 Medications Prescribed

Coding Instruction: Indicated if the medication is prescribed, not prescribed or is not prescribed for either a medical

or patient reason

Target Value: The value on Follow-up

Technical Specification

Code: 432102000
Code System: SNOMED CT
Short Name: F_MedAdmin1
Missing Data: Report

Harvested: Yes (TAVR, TMVR, TMVrpr,

TTVP)

Is Base Element: No
Is Followup
Element:
Data Type: CD
Precision:
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:
Data Source: User

Is Identifier: No

Parent/Child Validation

Element: 11990 Follow-Up Medications Code

Operator:

Value: Any Value

Follow-Up Medication Administration - 1.3.6.1.4.1.19376.1.4.1.6.5.371

Selection	Definition	Source	Code	Code System
Not Prescribed - Medical			100001034	ACC NCDR
Reason				
Not Prescribed - No Reas	son		100001048	ACC NCDR
Not Prescribed - Patient			100001071	ACC NCDR
Reason				
Yes - Prescribed			100001247	ACC NCDR

Element: 14577 Loop Diuretic Dose

Coding Instruction: Specify the total daily dose of the loop diuretic that was prescribed to the patient.

Target Value: The value on Follow-up

Technical Specification

Code: 112000001975 Code System: ACC NCDR

Short Name: FUMed_LoopDiureticDose

Missing Data: Report

Harvested: Yes (TMVR, TMVrpr, TTVP) Is Identifier: No

Is Base Element: No
Is Followup Element:
Data Type: PQ
Precision: 3,0
Selection Type: Single
Unit of Measure: mg
Default Value: Null
Usual Range: 1 - 40 mg
Valid Range: 1 - 300 mg

Data Source: User

Parent/Child Validation
11990 Follow-Up Medications Code

Element: 11990 Operator: Equal

Value: Loop Diuretics

AND ----

Element: 13696 Medications Prescribed

Operator: Equal

Value: Yes - Prescribed







Usual Range:

Valid Range: 1 - 999,999

Data Source: Automatic

Data Source: Automatic

Data Source: Automatic

Section: Administration Parent: Root **Technical Specification** Element: 1000 Participant ID Code: 2.16.840.1.113883.3.3478.4.836 Coding Instruction: Indicate the participant ID of the submitting facility. Code System: ACC NCDR Short Name: PartID Target Value: N/A Missing Data: Illegal Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: Data Type: NUM Precision: 6 Selection Type: Single Unit of Measure: Default Value: Null

Technical Specification Element: 1010 Participant Name Code: 2.16.840.1.113883.3.3478.4.836 Coding Instruction: Indicate the full name of the facility where the procedure was performed. Code System: ACC NCDR Short Name: PartName Missing Data: Illegal Values should be full, official hospital names with no abbreviations or variations in spelling. Harvested: Yes (BDS, TAVR, TMVR, Target Value: N/A TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: Data Type: ST Precision: 100 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:

Technical Specification Element: 1020 Time Frame of Data Submission Code: 1.3.6.1.4.1.19376.1.4.1.6.5.45 Coding Instruction: Indicate the time frame of data included in the data submission. Format: YYYYQQ. e.g.,2016Q1 Code System: ACC NCDR Target Value: N/A Short Name: Timeframe Missing Data: Illegal Harvested: Yes (BDS, TAVR, TMVR, TMVrpr, TTVP) Is Identifier: No Is Base Element: Yes Is Followup Element: Yes Data Type: ST Precision: 6 Selection Type: Single Unit of Measure: Default Value: Null **Usual Range:** Valid Range:



Element: 1040



Full Specifications **Data Dictionary v3.0**



Section: Administration	Parent: Root
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Transmission Number

Coding Instruction: 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 submission files are

exported. The transmission number should never be repeated.

Target Value: N/A

Technical Specification

Code: 1.3.6.1.4.1.19376.1.4.1.6.5.45

Code System: ACC NCDR Short Name: Xmsnld Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element: Yes
NUM
Precision: 9
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Valid Range: 1 - 999,999,999

Data Source: Automatic

Element: 1050 Vendor Identifier

Coding Instruction: Vendor identification (agreed upon by mutual selection between the vendor and the NCDR) to

identify software vendor. This is entered into the schema automatically by vendor software. Vendors must use consistent name identification across sites. Changes to vendor name

identification must be approved by the NCDR.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.840

Code System: ACC NCDR Short Name: Vendorld Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:
Data Type: ST
Precision: 15
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:
Valid Range:

Data Source: Automatic

Element: 1060 Vendor Software Version

Coding Instruction: Vendor's software product name and version number identifying the software which created

this record (assigned by vendor). Vendor controls the value in this field. This is entered into

the schema automatically by vendor software.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.847 **Code System:** ACC NCDR

Short Name: VendorVer Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No
Is Base Element: Yes
Is Followup
Element:

Data Type: ST
Precision: 20
Selection Type: Single
Unit of Measure:
Default Value: Null
Usual Range:

Data Source: Automatic

Valid Range:



Element: 1070



Full Specifications Data Dictionary v3.0



Section: Administration Parent: Root

Registry Identifier

Coding Instruction: The NCDR registry identifier describes the data registry to which these records apply. It is

implemented in the software at the time the data is collected and records are created. This is

entered into the schema automatically by software.

Target Value: N/A

Technical Specification

Code: 2.16.840.1.113883.3.3478.4.841

Code System: ACC NCDR Short Name: Registryld Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: Data Type: ST Precision: 30 Selection Type: Single Unit of Measure:

Default Value: ACC-NCDR-TVT-3.0

Usual Range: Valid Range:

Data Source: Automatic

Element: 1071 Registry Schema Version

Coding Instruction: Schema version describes the version number of the Registry Transmission Document (RTD)

schema to which each record conforms. It is an attribute that includes a constant value indicating the version of schema file. This is entered into the schema automatically by

Target Value: N/A

Technical Specification

Code: 1000142438 Code System: ACC NCDR Short Name: SchemaVersion

Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Element: Data Type: NUM Precision: 3,1 Selection Type: Single Unit of Measure: Default Value: 1 **Usual Range:** Valid Range:

Data Source: Automatic

Element: 1085 Submission Type

Coding Instruction: Indicate if the data contained in the harvest/data file contains either standard patient episode of

care records (arrival date to discharge only) or if it contains patient follow-up records.

A transmission file with all episode of care records (from Arrival to Discharge only) is

considered a 'Base Registry Record'.

A file with patient follow-up records (any follow-up assessments performed during the quarter

selected) is considered a 'Follow-Up Record'.

Note(s):

Selecting 'Follow-Up Records Only' will transmit all patient records with Follow-up Assessment Dates (Element Ref# 11000) contained in the selected timeframe, regardless of the procedure or discharge date. For example, if a patient has a procedure on 3/30/2017, is discharged on 3/31/2017, and has a follow-up assessment on 5/6/2017, the patient's episode of care data will be transmitted in the 2017Q1 Base Registry Record file, but the Follow-up data will be

transmitted in the 2017Q2 Follow-Up File.

Target Value: N/A

Technical Specification

Code: 1000142423 Code System: ACC NCDR Short Name: SubmissionType

Missing Data: Illegal

Harvested: Yes (BDS, TAVR, TMVR,

TMVrpr, TTVP)

Is Identifier: No Is Base Element: Yes Is Followup Yes Element: Data Type: CD

Precision: Selection Type: Single Unit of Measure:

> Default Value: Null **Usual Range:** Valid Range:

Data Source: Automatic

Submission Type

Selection	Definition	Source	Code	Code System
Episode of Care Records Only			1000142424	ACC NCDR
Follow-Up Records Only			1000142425	ACC NCDR

Element: 12903

Condition History Name

Value Set Name: Condition History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.340

Selections

Atrial Fibrillation | 49436004, Atrial Flutter | 5370000, Cardiomyopathy | 85898001, Carotid Artery Stenosis | 64586002, Cerebrovascular Accident | 230690007, Cerebrovascular Disease | 62914000, Chronic Lung Disease | 413839001, COVID-19 Positive | 112000001982, Dementia - Moderate to Severe | 112000001493, Diabetes Mellitus | 73211009, Endocarditis | 56819008, Heart Failure | 84114007, Hostile Chest | 112000001489, Hypertension | 38341003, Liver Disease | 235856003, Myocardial Infarction | 22298006, Peripheral Arterial

TVT Pathway (13171) IN (TMVr)

Disease | 399957001, Porcelain Aorta | 112000001175, Transient Ischemic Attack (TIA) | 266257000

Element: 12903

Value Set Name: Condition History Name

Condition History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.340

Selections

Atrial Fibrillation | 49436004, Atrial Flutter | 5370000, Carotid Artery Stenosis | 64586002, Cerebrovascular Accident | 230690007, Cerebrovascular Disease | 62914000, Chronic Lung Disease | 413839001, Conduction Defect | 44808001, COVID-19 Positive | 112000001982, Dementia - Moderate to Severe | 112000001493, Diabetes Mellitus | 73211009, Endocarditis | 56819008, Heart Failure | 84114007, Hostile Chest | 112000001489, Hypertension | 38341003, Liver Disease | 235856003, Myocardial Infarction | 22298006, Peripheral Arterial Disease | 399957001, Porcelain Aorta | 112000001175, Transient Ischemic Attack (TIA) | 266257000

Selection Dependency

Selection Dependency

Selection Dependency

Selection Dependency

TVT Pathway (13171) IN (TAVR)

Element: 12903

Value Set Name: Condition History Name

Condition History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.340

Selections

Atrial Fibrillation | 49436004, Atrial Flutter | 5370000, Cardiomyopathy | 85898001, Carotid Artery Stenosis | 64586002, Cerebrovascular Accident | 230690007, Cerebrovascular Disease | 62914000, Chronic Lung Disease | 413839001, COVID-19 Positive | 112000001982, Dementia - Moderate to Severe | 112000001493, Diabetes Mellitus | 73211009, Endocarditis | 56819008, Heart Failure | 84114007, Hostile Chest | 112000001489, Hypertension | 38341003, Liver Disease | 235856003, Myocardial Infarction | 22298006, Peripheral Arterial Disease | 399957001, Porcelain Aorta | 112000001175, Transient Ischemic Attack (TIA) |

TVT Pathway (13171) IN (TMVR)

266257000

Element: 12903

Value Set Name: Condition History Name

Condition History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.340

Selections

Atrial Fibrillation | 49436004, Atrial Flutter | 5370000, Cardiomyopathy | 85898001, Carotid Artery Stenosis | 64586002, Cerebrovascular Accident | 230690007, Cerebrovascular Disease | 62914000, Chronic Lung Disease | 413839001, Conduction Defect | 44808001, COVID-19 Positive | 112000001982, Dementia - Moderate to Severe | 112000001493, Diabetes Mellitus | 73211009, Endocarditis | 56819008, Heart Failure | 84114007, Hostile Chest | 112000001489, Hypertension | 38341003, Liver Disease | 235856003, Myocardial Infarction | 22298006, Peripheral Arterial Disease | 399957001, Porcelain Aorta | 112000001175, Transient Ischemic Attack (TIA) | 266257000

Selection Dependency TVT Pathway (13171) IN (Tricuspid Valve Procedure)

Element: 12905

Value Set Name: Procedure History Name

Procedure History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.341

Selections

Aortic Valve Procedure | 112000001755, Aortic Valve Repair Surgery | 112816004, Aortic Valve Replacement Surgery | 725351001, Aortic Valve Replacement - Transcatheter | 41873006, Coronary Artery Bypass Graft | 232717009, Implantable Cardioverter Defibrillator |

447365002, Mitral Valve Procedure | 112000001940, Mitral Valve Annuloplasty Ring Surgery | 232744004, Mitral Valve Repair Surgery | 384641003, Mitral Valve Replacement Surgery | 53059001, Mitral Valve Transcatheter Intervention | 112000001773, PCI | 415070008, Permanent Pacemaker | 449397007, Pulmonic Valve Procedure | 112000001769, Tricuspid

Valve Procedure | 112000001941

TVT Pathway (13171) IN (TMVr)

Element: 12905

Value Set Name: Procedure History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.341

Selection Dependency TVT Pathway (13171) IN (TAVR)

Procedure History Name

Selections

Aortic Valve Procedure | 112000001755, Aortic Valve Balloon Valvuloplasty | 77166000,

Aortic Valve Repair Surgery | 112816004, Aortic Valve Replacement Surgery | 725351001, Aortic Valve Replacement - Transcatheter | 41873006, Aortic Valve Transcatheter Intervention | 112000001768, Coronary Artery Bypass Graft | 232717009, Implantable Cardioverter Defibrillator | 447365002, Mitral Valve Procedure | 112000001940, Mitral Valve Annuloplasty Ring Surgery | 232744004, Mitral Valve Repair Surgery | 384641003, Mitral

Effective for Patient Discharged January 01, 2021

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Full Specifications

Data Dictionary v3.0

STS/ACC TVT Registr

STS/ACC TVT Registry Data Dictionary v3.0 Full Specifications

Value Set Member Constraints

Valve Replacement Surgery | 53059001, Mitral Valve Transcatheter Intervention | 112000001773, PCI | 415070008, Permanent Pacemaker | 449397007, Pulmonic Valve Procedure | 112000001769, Tricuspid Valve Procedure | 112000001941

Element: 12905

Element: 12905

Element: 14241

Value Set Name: Procedure History Name

Procedure History Name

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.341

Selections

Aortic Valve Procedure | 112000001755, Aortic Valve Repair Surgery | 112816004, Aortic Valve Replacement Surgery | 725351001, Aortic Valve Replacement - Transcatheter |

41873006, Coronary Artery Bypass Graft | 232717009, Implantable Cardioverter Defibrillator | 447365002, Mitral Valve Procedure | 112000001940, Mitral Valve Annuloplasty Ring Surgery | 232744004, Mitral Valve Repair Surgery | 384641003, Mitral Valve Replacement Surgery | 53059001, Mitral Valve Transcatheter Intervention | 112000001773, PCI | 415070008, Permanent Pacemaker | 449397007, Pulmonic Valve Procedure | 112000001769, Tricuspid Valve Procedure | 112000001941

Selection Dependency TVT Pathway (13171) IN (TMVR)

TVT Pathway (13171) IN (Tricuspid Valve Procedure)

Procedure History Name

Value Set Name: Procedure History Name OID: 1.3.6.1.4.1.19376.1.4.1.6.5.341

Selections

ections Selection Dependency

Aortic Valve Procedure | 112000001755, Aortic Valve Repair Surgery | 112816004, Aortic Valve Replacement Surgery | 725351001, Aortic Valve Replacement - Transcatheter | 41873006, Coronary Artery Bypass Graft | 232717009, Implantable Cardioverter Defibrillator | 447365002, Mitral Valve Procedure | 112000001940, Mitral Valve Annuloplasty Ring Surgery | 323744004, Mitral Valve Repair Surgery | 384641003, Mitral Valve Replacement Surgery | 53059001, Mitral Valve Transcatheter Intervention | 112000001773, PCI | 415070008, Permanent Pacemaker | 449397007, Pulmonic Valve Procedure | 112000001769, Tricuspid Valve Procedure | 112000001741, Tricuspid Valve Repair Surgery | 384643000, Tricuspid Valve Replacement Surgery | 25236004, Tricuspid Valve Replacement - Transcatheter | 112000001977, Tricuspid Valve Transcatheter Intervention | 112000001779

Mitral Valve Replacement Type

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.734

Value Set Name: Mitral Valve Replacement Type

 Selections
 Selection Dependency

 Stented | 112000001758, Stentless | 112000001760
 TVT Pathway (13171) IN (TMVR)

Element: 14241Mitral Valve Replacement TypeValue Set Name: Mitral Valve Replacement TypeOID: 1.3.6.1.4.1.19376.1.4.1.6.5.734

Selections Selection Dependency

Machanical J 705004002 Stantad J 442000004759 Stantage J 442000004750 TVT Pathyray (42474) IN J 7AVP)

Mechanical | 705991002, Stented | 112000001758, Stentless | 112000001760 TVT Pathway (13171) IN (TAVR)

Element: 14273 Transcatheter Valve Therapy Procedure Type

Value Set Name: Transcatheter Valve Therapy Procedure OID: 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selections Selection Dependency

TAVR | 41873006 TVT Pathway (13171) IN (TAVR)

Element: 14273 Transcatheter Valve Therapy Procedure Type

Value Set Name: Transcatheter Valve Therapy ProcedureOID: 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selections Selection Dependency
TMVR | 112000001458 TVT Pathway (13171) IN (TMVR)

Element: 14273 Transcatheter Valve Therapy Procedure Type

Value Set Name: Transcatheter Valve Therapy Procedure OID: 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selections Selection Dependency
Tricuspid Valve Procedure | 112000001977 TVT Pathway (13171) IN (Tricuspid Valve Procedure)

Element: 14273 Transcatheter Valve Therapy Procedure Type

Value Set Name: Transcatheter Valve Therapy Procedure OID: 1.3.6.1.4.1.19376.1.4.1.6.5.695

Selections Selection Dependency
TMVr | 112000001801 TVT Pathway (13171) IN (TMVr)

Element: 13506 Reason for Aborting Procedure

Value Set Name: Transcatheter Valve Therapy Procedure Aborted Reasons OID: 1.3.6.1.4.1.19376.1.4.1.6.5.554

Selections Selection Dependency

Access Related | 112000001460, Navigation Issue After Successful Access | 112000001461, New Clinical Findings | 112000001462, Device or Delivery System

Transcatheter Valve Therapy Procedure Type (14273) IN (TMVr, TMVR)

Malfunction | 112000001463, Patient Clinical Status | 112000001464, Consent Issue | 112000001465, Transseptal Access Related | 112000001466, System Issue | 112000001467, Other | 100000351

Element: 13506

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Value Set Name: Transcatheter Valve Therapy Procedure Aborted Reasons

Reason for Aborting Procedure OID: 1 3 6 1 4 1 19376 1 4 1 6 5 554

Selection Dependency

Access Related | 112000001460, Navigation Issue After Successful Access | 112000001461, New Clinical Findings | 112000001462, Device or Delivery System Malfunction | 112000001463, Patient Clinical Status | 112000001464, Consent Issue | 112000001465, System Issue | 112000001467, Other | 100000351

Element: 13543 Reason for Conversion to Open Heart Surgery

Value Set Name: Reason for Conversion to Open Heart Surgery OID: 1.3.6.1.4.1.19376.1.4.1.6.5.513

Selection Dependency

Access Related | 112000001460, Cardiac Tamponade | 35304003, Inability to Position Device Transcatheter Valve Therapy Procedure Type (14273) IN (TMVr) | 112000001479, Device Embolization | 112000001324, Valve Injury | 762610001, Other |

100000351

Element: 14485

Reason for Conversion to Open Heart Surgery Element: 13543

Value Set Name: Reason for Conversion to Open Heart Surgery OID: 1.3.6.1.4.1.19376.1.4.1.6.5.513

Selection Dependency

Valve Dislodged to Aorta | 112000001328, Valve Dislodged to Left Ventricle | 112000001329, Transcatheter Valve Therapy Procedure Type (14273) IN (TAVR, TMVR, Tricuspid Valve Annulus Rupture | 112000001331, Ventricular Rupture | 112000001330, Aortic Dissection | 308546005, Coronary Occlusion | 63739005, Access Related | 112000001460, Cardiac Tamponade | 35304003, Inability to Position Device | 112000001479, Device Embolization |

Simulus FLX-O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic

112000001324, Valve Injury | 762610001, Other | 100000351

Transcatheter Valve Therapy Procedure Type (14273) IN (TAVR, Tricuspid Valve

Transcatheter Aortic Valve Replacement Device ID

Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

> Selections Selection Dependency

Mitral Valve | 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390, Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process | 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process | 4406, Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process | 4407, Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring | 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring |

4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex

Patient Enrolled in Research Study (3020) IN (No (or Not Answered)) AND Transcatheter Valve Therapy Procedure Type (14273) IN (TAVR)

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Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process | 4428, Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve | 4456, Biocor Stented Aortic Tissue Valve | 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System I 4483, Commander Delivery System I 4484, Edwards Expandable Introducer Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm | 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm x 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Sentinel Cerebral Protection System | 4522, EnVeo PRO Loading System | 4525, EnVeo PRO Loading System | 4526, EnVeo PRO Loading System | 4527, EnVeo PRO Delivery System with InLine Sheath | 4528, EnVeo PRO Delivery System with InLine Sheath | 4529, Lotus Edge | 4533, Evolut Pro Plus | 4534, Mosaic | 4535, Commander Delivery System | 4536, CG Future Band | 4587, INSPIRIS RESILIA Aortic Valve | 4592, PERCEVAL Aortic Valve | 4593, Evolut FX | 5156, Portico Transcatheter Heart Valve | 5162, Portico Transcatheter Heart Valve | 5163, Portico Transcatheter Heart Valve | 5164, Portico Transcatheter Heart Valve | 5165, SAPIEN 3 Ultra | 4341

Element: 14485
Value Set Name: TVT Procedure Devices

Transcatheter Aortic Valve Replacement Device ID

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Selections

Simulus FLX-O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic Mitral Valve | 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390,

Selection Dependency

Patient Enrolled in Research Study (3020) IN (Yes) AND Transcatheter Valve Therapy Procedure Type (14273) IN (TAVR)

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Data Dictionary v3.0

Value Set Member Constraints

Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process | 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process | 4406, Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process | 4407, Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring I 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring | 4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process | 4428, Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve | 4456, Biocor Stented Aortic Tissue Valve | 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System | 4483, Commander Delivery System | 4484, Edwards Expandable Introducer Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter | 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm \times 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Sentinel Cerebral Protection System | 4522, EnVeo PRO Loading System | 4525, EnVeo PRO Loading System | 4526, EnVeo PRO Loading System | 4527, EnVeo PRO Delivery System with InLine Sheath | 4528, EnVeo PRO Delivery System with InLine Sheath | 4529, Lotus Edge | 4533, Evolut Pro Plus | 4534, Mosaic | 4535, Commander Delivery System | 4536, SAPIEN 3 (research study device) | 4538, CoreValve

Evolut R (research study device) | 4539, CoreValve Evolut PRO (research study device) | 4540, CG Future Band | 4587, INSPIRIS RESILIA Aortic Valve | 4592, PERCEVAL Aortic Valve | 4593, Evolut FX | 5156, Portico Transcatheter Heart Valve | 5162, Portico Transcatheter Heart Valve | 5163. Portico Transcatheter Heart Valve | 5164. Portico Transcatheter Heart Valve | 5165, SAPIEN 3 Ultra | 4341

Element: 13795

Value Set Name: TVT Procedure Devices

Steerable Guide Cath Device ID OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Procedure Type (14273) IN (TMVr)

Selection Dependency

Valve Therapy Procedure Type (14273) IN (TMVr)

Valve Therapy Procedure Type (14273) IN (TMVr)

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Procedure Type (14273) IN (TMVr)

MitraClip NT Steerable Guide Catheter | 4342, MitraClipDeliverySystem | 4349, MitraClip NT Patient Enrolled in Research Study (3020) IN (Yes) AND Transcatheter Valve Therapy Clip Delivery System (research study device) | 4537, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, MitraClipClipDeliverySystem (research study device) | 4549, SteerableGuideCatheter (research study device) | 4550, MitraClip NT Steerable Guide Catheter (research study device) | 4551, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter | 4553, MitraClip G4 Clip Delivery System NT | 5151, MitraClip NT Clip Delivery System | 4343

Selections

Element: 13795 Steerable Guide Cath Device ID Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

> **Selection Dependency** Selections

MitraClip NT Steerable Guide Catheter | 4342, MitraClipClipDeliverySystem | 4349, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter | 4553, MitraClip G4 Clip Delivery

System NT | 5151, MitraClip NT Clip Delivery System | 4343

Mitral Repair Device ID Element: 13797 Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Selections **Selection Dependency**

MitraClip NT Steerable Guide Catheter | 4342, MitraClipDeliverySystem | 4349, MitraClip NT Patient Enrolled in Research Study (3020) IN (Yes) AND Transcatheter Valve Therapy Clip Delivery System (research study device) | 4537, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, MitraClipClipDeliverySystem (research study device) | 4549, SteerableGuideCatheter (research study device) | 4550, MitraClip NT

Steerable Guide Catheter (research study device) | 4551, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter | 4553, MitraClip G4 Clip Delivery System NT |

5151, MitraClip NT Clip Delivery System | 4343

Element: 13797 Mitral Repair Device ID

Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Selections **Selection Dependency** Patient Enrolled in Research Study (3020) IN (No (or Not Answered)) AND Transcatheter

MitraClip NT Steerable Guide Catheter | 4342, MitraClipClipDeliverySystem | 4349, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter | 4553, MitraClip G4 Clip Delivery

System NT | 5151, MitraClip NT Clip Delivery System | 4343

Value Set Name: TVT Procedure Devices

Transcatheter Mitral Valve Replacement Device ID Element: 14484

Selections **Selection Dependency**

Simulus FLX-O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic Mitral Valve | 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future

Patient Enrolled in Research Study (3020) IN (No (or Not Answered)) AND Transcatheter Valve Therapy Procedure Type (14273) IN (TMVR)

Patient Enrolled in Research Study (3020) IN (No (or Not Answered)) AND Transcatheter

Full Specifications

Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390, Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process | 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process | 4406, Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process I 4407. Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring | 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring | 4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process I 4428. Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve | 4456, Biocor Stented Aortic Tissue Valve | 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System | 4483, Commander Delivery System | 4484, Edwards Expandable Introducer Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter | 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm | 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm x 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Lotus Edge | 4533, Commander Delivery System | 4536, CG Future Band | 4587, SAPIEN 3 Ultra | 4341

Transcatheter Mitral Valve Replacement Device ID

Element: 14484

00

OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Selection Dependency

Value Set Name: TVT Procedure Devices Selections Simulus FLX-O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic Mitral Valve | 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390, Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process | 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process I 4406. Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process | 4407, Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring | 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring | 4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards

Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve | 4456, Biocor Stented Aortic Tissue Valve | 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor

Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System | 4483, Commander Delivery System | 4484, Edwards Expandable Introducer

Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards

Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process | 4428, Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary

Patient Enrolled in Research Study (3020) IN (Yes) AND Transcatheter Valve Therapy Procedure Type (14273) IN (TMVR)

Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter | 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm | 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm x 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Lotus Edge | 4533, Commander Delivery System | 4536, CG Future Band | 4587, SAPIEN 3 Ultra | 4341

Transcatheter Tricuspid Valve Device ID Element: 14483 Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

> Selections **Selection Dependency**

> > Valve Therapy Procedure Type (14273) IN (Tricuspid Valve Procedure)

MitraClip NT Steerable Guide Catheter | 4342, MitraClipDeliverySystem | 4349, Simulus FLX Patient Enrolled in Research Study (3020) IN (No (or Not Answered)) AND Transcatheter -O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic Mitral Valve | 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390, Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process I 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process | 4406, Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process | 4407, Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring | 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring | 4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process | 4428, Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic

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Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve | 4456, Biocor Stented Aortic Tissue Valve | 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System I 4483, Commander Delivery System I 4484, Edwards Expandable Introducer Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter | 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm | 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm x 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Sentinel Cerebral Protection System | 4522, EnVeo PRO Loading System | 4525, EnVeo PRO Loading System | 4526, EnVeo PRO Loading System | 4527, EnVeo PRO Delivery System with InLine Sheath | 4528, EnVeo PRO Delivery System with InLine Sheath | 4529, Lotus Edge | 4533, Evolut Pro Plus | 4534, Mosaic | 4535, Commander Delivery System | 4536, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, CG Future Band | 4587, INSPIRIS RESILIA Aortic Valve | 4592, PERCEVAL Aortic Valve | 4593, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter | 4553, MitraClip G4 Clip Delivery System NT | 5151, MitraClip NT Clip Delivery System | 4343, SAPIEN 3 Ultra | 4341

Element: 14483 Transcatheter Tricuspid Valve Device ID Value Set Name: TVT Procedure Devices OID: 1.3.6.1.4.1.19376.1.4.1.6.5.746

Selections Selection Dependency

MitraClip NT Steerable Guide Catheter | 4342, MitraClipDeliverySystem | 4349, Simulus FLX Patient Enrolled in Research Study (3020) IN (Yes) AND Transcatheter Valve Therapy -O Ring | 4339, Carpentier-Edwards Porcine Aortic Bioprosthesis | 4335, Epic Mitral Valve | Procedure Type (14273) IN (Tricuspid Valve Procedure) 4337, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4350, Sapien Valve Transfemoral, RetroFlex 3 Delivery | 4351, RetroFlex 3 | 4352, RetroFlex 3 | 4353, RetroFlex 3 | 4354, RetroFlex 3 | 4355, Sapien Valve Transapical | 4356, Ascendra Delivery System | 4357, Ascendra Delivery System | 4358, Ascendra Sheath Set | 4359, Ascendra Transapical | 4360, CoreValve System; DCS | 4361, CoreValve System; DCS | 4362, CoreValve System; CLS | 4363, Ascendra 3 Delivery System | 4364, Ascendra 3 Delivery System | 4365, Ascendra 3 Sheath Set | 4366, Ascendra 3 Transapical | 4367, Sapien XT Heart Valve | 4368, Simulus FLX-C Band | 4369, Simulus Semi-rigid Mitral Annuloplasty Ring | 4370, Simulus Adjustable Ring | 4371, Simulus Adjustable Band | 4372, Simulus Semi-rigid Ring | 4373, TriAd Tricuspid Annuloplasty Ring | 4374, 3f Aortic Bioprosthesis | 4375, 3f Enable Aortic Bioprosthesis | 4376, Hancock Apical Left Ventricle Connector | 4377, Contegra Unsupported Pulmonary Valve Conduit | 4378, Contegra Supported Pulmonary Valve Conduit | 4379, Mosaic Ultra Porcine Heart Valve | 4380, Mosaic Mitral Bioprosthesis | 4381, Duran Band | 4382, Duran Ring | 4383, Duran AnCore Band | 4384, Duran AnCore Band With Chordal Guide | 4385, Duran AnCore Ring | 4386, Duran Ancore Ring With Chordal Guide | 4387, CG Future Composite Ring | 4388, Profile 3D Ring | 4389, Contour 3D Annuloplasty Ring | 4390, Freestyle, Complete Subcoronary Aortic Bioprostheses | 4391, Freestyle, Modified Subcoronary Aortic Bioprostheses | 4392, Freestyle, Full Root - Aortic Bioprosthesis | 4393, Melody Transcatheter Pulmonary Valve | 4394, Hancock II Aortic Bioprosthesis | 4395, Hancock II Ultra Bioprosthesis Small Root System | 4396, Hancock II Mitral Bioprosthesis | 4397, Prima Aortic Stentless Bioprosthesis | 4398, Prima Plus Stentless Aortic Bioprosthesis | 4399, Carpentier-Edwards S.A.V. Aortic Porcine Bioprosthesis | 4400, Carpentier-Edwards Perimount Pericardial Aortic Bioprosthesis | 4401, Carpentier-Edwards Perimount Theon Pericardial Aortic Bioprosthesis with ThermaFix Process | 4402, Carpentier-Edwards Perimount RSR Pericardial Aortic Bioprosthesis | 4403, Carpentier-Edwards Perimount Theon

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Value Set Member Constraints

RSR Pericardial Aortic Bioprosthesis with ThermaFix Process | 4404, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis | 4405, Carpentier-Edwards Perimount Magna Pericardial Aortic Bioprosthesis with ThermaFix Process | 4406, Carpentier-Edwards Perimount Magna Ease Pericardial Aortic Bioprosthesis with ThermaFix Process | 4407, Carpentier-McCarthy-Adams IMR ETlogix Mitral Annuloplasty Ring | 4408, Edwards GeoForm Mitral Annuloplasty Ring | 4409, Carpentier-Edwards Bioprosthetic Valved Conduit | 4410, Carpentier-Edwards Classic Mitral Annuloplasty Ring | 4411, Carpentier-Edwards Classic Mitral Annuloplasty Ring with Duraflo Treatment | 4412, Carpentier-Edwards Physio Mitral Annuloplasty Ring | 4413, Carpentier-Edwards Physio Annuloplasty Ring with Duraflo Treatment | 4414, Carpentier-Edwards Classic Tricuspid Annuloplasty Ring | 4415, Carpentier -Edwards Classic Tricuspid Annuloplasty Ring with Duraflo Treatment | 4416, Cosgrove-Edwards Mitral/Tricuspid Annuloplasty Ring | 4417, Cosgrove-Edwards Annuloplasty System with Duraflo Treatment | 4418, MC3 Tricuspid Annuloplasty System | 4419, DETlogix Mitral Annuloplasty Ring | 4420, Myxomatous Annuloplasty Ring | 4421, Carpentier-Edwards Physio II Mitral Annuloplasty Ring | 4422, Carpentier-Edwards Physio Tricuspid Annuloplasty Ring | 4423, Carpentier-Edwards Porcine Mitral Bioprosthesis | 4424, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis with Extended Suture Ring | 4425, Carpentier-Edwards Duraflex Low Pressure Porcine Mitral Bioprosthesis | 4426, Carpentier-Edwards Perimount Plus Mitral Pericardial Bioprosthesis | 4427, Carpentier-Edwards Perimount Theon Mitral Pericardial Bioprosthesis with ThermaFix Process | 4428, Carpentier-Edwards Perimount Magna Mitral Pericardial Bioprosthesis | 4429, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis | 4430, Carpentier-Edwards Perimount Magna Mitral Ease Pericardial Bioprosthesis with ThermaFix Process | 4431, Cribier-Edwards Aortic Bioprosthesis | 4432, Cribier-Edwards Aortic Bioprosthesis | 4433, Intuity Transcatheter Heart Valve | 4434, CardioGraft Aortic Heart Valve - Large | 4435, CardioGraft Aortic Heart Valve - Medium | 4436, CardioGraft Aortic Heart Valve - Small | 4437, CardioGraft Pulmonary Heart Valve - Large | 4438, CardioGraft Pulmonary Heart Valve - Medium | 4439, CardioGraft Pulmonary Heart Valve - Small | 4440, Sovering Band (Mitral) | 4441, MEMO 3D Semi-rigid Annuloplasty Ring | 4442, Mitroflow Aortic Pericardial Heart Valve | 4443, Mitroflow Aortic Pericardial Heart Valve with PRT | 4444, Mitroflow Aortic Pericardial Heart Valve with PRT | 4445, Carbomedics AnnuloFlex Annuloplasty System | 4446, Carbomedics AnnuloFlo Annuloplasty System | 4447, Pericarbon Freedom Stentless | 4448, Soprano Armonia | 4449, Freedom Solo | 4450, Attune Flexible Adjustable Annuloplasty Ring | 4451, Biocor Aortic Valve | 4452, Biocor Aortic Valve | 4453, Biocor Mitral Valve | 4454, Biocor Mitral Valve | 4455, Trifecta Aortic Stented Tissue Valve I 4456, Biocor Stented Aortic Tissue Valve I 4457, Biocor Stented Mitral Tissue Valve | 4458, Biocor Porcine Stentless Bioprosthetic Heart Valve | 4459, Biocor Aortic Valve | 4460, Biocor Mitral Valve | 4461, Biocor Supra Aortic Stented Tissue Valve | 4462, Epic Aortic Stented Tissue Valve | 4463, Epic Mitral Stented Tissue Valve | 4464, Epic Aortic Valve | 4465, Epic Tissue Aortic Valve with Silzone Coating | 4466, Epic Tissue Mitral Valve with Silzone Coating | 4467, Epic Supra Aortic Stented Tissue Valve | 4468, Epic Stented Aortic Tissue Valve | 4469, Rigid Saddle Ring | 4470, Seguin Semi-Rigid Annuloplasty Ring | 4471, Seguin Annuloplasty Ring with Silzone Coating | 4472, Tailor Flexible Annuloplasty Band | 4473, Tailor Annuloplasty Ring with Silzone Coating | 4474, Tailor Flexible Annuloplasty Ring | 4475, Toronto SPV Valve | 4476, Toronto SPV II Bioprosthetic Heart Valve | 4477, CryoLife Aortic Valve and Conduit | 4478, CryoLife Aortic Valve without Conduit | 4479, Homograft valve (manufacturer not specified) | 4480, Commander Delivery System | 4481, Commander Delivery System | 4482, Commander Delivery System | 4483, Commander Delivery System | 4484, Edwards Expandable Introducer Sheath Set | 4485, Edwards Expandable Introducer Sheath Set | 4486, Edwards Transfemoral Balloon Catheter | 4487, Edwards Transfemoral Balloon Catheter | 4488, Edwards Transfemoral Balloon Catheter | 4489, Edwards Transfemoral Balloon Catheter | 4490, Crimper, Universal | 4491, Atrion QL2530 Inflation Device, 25 mL | 4492, Atrion QL38 Locking Syringe Device, 38 mL | 4493, Edwards Certitude Delivery System | 4494, Edwards Certitude Delivery System | 4495, Edwards Certitude Delivery System | 4496, Edwards Certitude Delivery System | 4497, Ascendra Balloon Aortic Valvuloplasty Catheter, 20mm | 4498, Edwards Certitude Introducer Sheath Set | 4499, Edwards Certitude Introducer Sheath Set | 4500, CoreValve System; TAV | 4501, CoreValve System; TAV | 4502, CoreValve Evolut R TAV | 4503, EnVeo TM R Delivery Catheter System | 4504, EnVeo TM R Loading System | 4505, EnVeo TM R Loading System | 4506, SAPIEN 3 | 4507, Commander Delivery System | 4508, Commander Delivery System | 4509, Commander Delivery System | 4510, Commander Delivery System | 4511, Edwards Expandable Introducer Sheath Set, 14F | 4512, Edwards Expandable Introducer Sheath Set, 16F | 4513, Edwards Balloon Catheter, 4 cm x 16 mm | 4514, Edwards Balloon Catheter, 4 cm x 20 mm | 4515, Edwards Balloon Catheter, 4 cm x 23 mm | 4516, Edwards Balloon Catheter, 4 cm x 25 mm | 4517, Crimper | 4518, INTUITY Elite Valve | 4520, Evolut PRO System | 4521, Sentinel Cerebral Protection System | 4522, EnVeo PRO Loading System | 4525, EnVeo PRO Loading System | 4526, EnVeo PRO Loading System | 4527, EnVeo PRO Delivery System with InLine Sheath | 4528, EnVeo PRO Delivery System with InLine Sheath | 4529, Lotus Edge | 4533, Evolut Pro Plus | 4534, Mosaic | 4535, Commander Delivery System | 4536, MitraClip NT Clip Delivery System (research study device) | 4537, SAPIEN 3 (research study device) | 4538, CoreValve Evolut R (research study device) | 4539, CoreValve Evolut PRO (research study device) | 4540, MitraClip NTR Clip Delivery System | 4541, MitraClip XTR Clip Delivery System | 4542, MitraClip G4 Clip Delivery System NT | 4543, MitraClip G4 Clip Delivery System NTW | 4544, MitraClip G4 Clip Delivery System XT | 4545, MitraClip G4 Delivery System XTW | 4546, MitraClipClipDeliverySystem (research study device) | 4549, CG Future Band | 4587, INSPIRIS RESILIA Aortic Valve | 4592, PERCEVAL Aortic Valve | 4593, SteerableGuideCatheter (research study device) | 4550, MitraClip NT Steerable Guide Catheter (research study device) | 4551, Steerable Guide Catheter | 4552, MitraClip G4 SGC Steerable Guide Catheter |

Element: 12153

Value Set Member Constraints

4553, MitraClip G4 Clip Delivery System NT | 5151, MitraClip NT Clip Delivery System | 4343, SAPIEN 3 Ultra | 4341

Value Set Name: Intra or Post Procedure Events

Intra or Post Procedure Events
OID: 1.3.6.1.4.1.19376.1.4.1.6.5.706

Selections

Annular Rupture | 11200001835, Atrial Fibrillation | 49436004, Bleeding - Access Site | 1000142440, Bleeding - Gastrointestinal | 74474003, Bleeding - Genitourinary | 417941003, Bleeding - Hematoma at Access Site | 385494008, Bleeding - Other | 1000142371, Bleeding - Retroperitoneal | 95549001, Cardiac Arrest | 410429000, Cardiac Perforation | 36191001:123005000=302509004, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, Coronary Artery Compression | 112000001837, COVID-19 Positive | 112000001982, Device Embolization | 112000001324, Device Migration | 370512004, Device Related Event - Other | 112000001828, Device Thrombosis | 112000001839, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Myocardial Infarction | 22298006, Pacemaker Lead Dislodgement or Dysfunction | 112000001884, Percutaneous Coronary Intervention | 415070008, Permanent Pacemaker | 449397007, Pulmonary Embolism | 59282003, Reintervention - Tricuspid Valve | 112000001820, Stroke - Hemorrhagic | 230706003, Stroke - Ischemic | 422504002, Stroke - Undetermined |

Selection Dependency
Transcatheter Valve Therapy Procedure Type (14273) IN (Tricuspid Valve Procedure) AND
TVT Pathway (13171) IN (Tricuspid Valve Procedure)

Element: 12153

Value Set Name: Intra or Post Procedure Events

Intervention - Unplanned | 112000000467

Intra or Post Procedure Events
OID: 1.3.6.1.4.1.19376.1.4.1.6.5.706

Selections

230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 11200000460, Vascular Complication - Minor | 112000001823, Vascular Surgery or

ASD Defect Closure due to Transseptal Catheterization | 112000001885, Atrial Fibrillation | 49436004, Bleeding - Access Site | 1000142440, Bleeding - Gastrointestinal | 74474003, Bleeding - Genitourinary | 417941003, Bleeding - Hematoma at Access Site | 385494008, Bleeding - Other | 1000142371, Bleeding - Retroperitoneal | 95549001, Cardiac Arrest | 410429000, Cardiac Perforation | 36191001:123005000=302509004, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, COVID-19 Positive | 112000001982, Device Embolization | 112000001324, Device Migration | 370512004, Device Related Event - Other | 112000001828, Device Thrombosis | 112000001839, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Left Ventricular Outflow Tract Obstruction | 253546004, Myocardial Infarction | 22298006, Permanent Pacemaker | 449397007, Reintervention - Mitral Valve | 112000001893, Stroke - Hemorrhagic | 230706003, Stroke - Ischemic | 422504002, Stroke - Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Transseptal Complication - Minor | 112000001823, Vascular Complication - Major | 11200000460, Vascular Complication - Minor | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Selection Dependency
Transcatheter Valve Therapy Procedure Type (14273) IN (TMVR) AND TVT Pathway
(13171) IN (TMVR)

Element: 12153

Value Set Name: Intra or Post Procedure Events

Intra or Post Procedure Events
OID: 1.3.6.1.4.1.19376.1.4.1.6.5.706

Selections

Annular Rupture | 112000001835, Aortic Dissection | 308546005, Atrial Fibrillation | 49436004, Bleeding - Access Site | 1000142440, Bleeding - Gastrointestinal | 74474003, Bleeding - Genitourinary | 417941003, Bleeding - Hematoma at Access Site | 385494008, Bleeding - Other | 1000142371, Bleeding - Retroperitoneal | 95549001, Cardiac Arrest | 410429000, Cardiac Perforation | 36191001:123005000=302509004, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, Coronary Artery Compression | 112000001837, COVID-19 Positive | 112000001982, Device Embolization | 112000001324, Device Migration | 370512004, Device Related Event - Other | 112000001828, Device Thrombosis | 112000001839, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Myocardial Infarction | 22298006, Percutaneous Coronary Intervention | 415070008, Permanent Pacemaker | 449397007, Reintervention - Aortic Valve | 112000001827, Stroke - Hemorrhagic | 230706003, Stroke - Ischemic | 422504002, Stroke - Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Selection Dependency
Transcatheter Valve Therapy Procedure Type (14273) IN (TAVR) AND TVT Pathway (13171) IN (TAVR)

Element: 12153

Value Set Name: Intra or Post Procedure Events

Intra or Post Procedure Events
OID: 1.3 6.1 4.1 19376.1 4.1 6.5 706

Selection

ASD Defect Closure due to Transseptal Catheterization | 112000001885, Atrial Fibrillation | 49436004, Bleeding - Access Site | 1000142440, Bleeding - Gastrointestinal | 74474003, Bleeding - Genitourinary | 417941003, Bleeding - Hematoma at Access Site | 385494008, Bleeding - Other | 1000142371, Bleeding - Retroperitoneal | 95549001, Cardiac Arrest | 410429000, Cardiac Perforation | 36191001:123005000=302509004, Cardiac Arrest | 112000001840, COVID-19 Positive | 112000001892, Complete Leaflet Clip Detachment | 112000001840, COVID-19 Positive | 112000001982, Delivery System Component Embolization | 112000001841, Device Embolization | 112000001324, Device Related Event - Other | 112000001828, Device Thrombosis | 112000001839, Dialysis (New Requirement) |

Selection Dependency
Transcatheter Valve Therapy Procedure Type (14273) IN (TMVr) AND TVT Pathway (13171)
IN (TMVr)

00

Other | 100000351

Selection Dependency

Selection Dependency

TVT Pathway (13171) IN (TMVr, TMVR, Tricuspid Valve Procedure)

Value Set Member Constraints

100014076, Endocarditis | 56819008, Mitral Leaflet or Subvalvular Injury | 112000001886, Myocardial Infarction | 22298006, Permanent Pacemaker | 449397007, Reintervention - Mitral Valve | 112000001893, Single Leaflet Device Attachment | 112000001538, Stroke -Hemorrhagic | 230706003, Stroke - Ischemic | 422504002, Stroke - Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Transseptal Complication | 112000001833, Vascular Complication - Major | 112000000460, Vascular Complication -Minor | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Discharge Location After Event Element: 14352 Value Set Name: Discharge Location OID: 1.3.6.1.4.1.19376.1.4.1.6.5.41

Selections Home | 01, Skilled Nursing Facility | 03, Extended Care/TCU/Rehab | 62, Other Discharge Status (14314) IN (Alive)

Location | 100001249

Element: 14361 Mitral Valve Reintervention Indication Value Set Name: Valve Reintervention Indication OID: 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selections **Selection Dependency** Regurgitation | 40445007, Stenosis | 44241007, Device Embolization | 112000001324, Transcatheter Valve Therapy Procedure Type (14273) IN (TMVr, TMVR) Endocarditis | 56819008, Device Thrombosis | 112000001839, Valve Injury | 762610001,

Element: 10200 Discharge Medication Code Value Set Name: Discharge Medication OID: 1.3.6.1.4.1.19376.1.4.1.6.5.165

Selection Dependency

Aldosterone Antagonist | 372603003, Angiotensin Converting Enzyme Inhibitor | 41549009, Angiotensin II Receptor Blocker | 372913009, Aspirin | 1191, Beta Blocker | 33252009, Direct Factor Xa Inhibitor | 112000000696, Direct thrombin inhibitor | 414010005, Diuretics Not Otherwise Specified | 112000001417, Loop Diuretics | 29051009, P2Y12 Antagonist |

112000001003, Thiazides | 372747003, Warfarin | 11289

Element: 10200 Discharge Medication Code Value Set Name: Discharge Medication OID: 1.3.6.1.4.1.19376.1.4.1.6.5.165

Selections Selection Dependency

Aspirin | 1191, Direct Factor Xa Inhibitor | 112000000696, Direct thrombin inhibitor | TVT Pathway (13171) IN (TAVR)

414010005, P2Y12 Antagonist | 112000001003, Warfarin | 11289

Element: 12933 Follow-up Event Name

Value Set Name: Follow Up Events OID: 1.3.6.1.4.1.19376.1.4.1.6.5.356

ASD Defect Closure due to Transseptal Catheterization | 112000001885, Atrial Fibrillation | Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TMVr) 49436004, Bleeding - Life Threatening | 112000000459, Bleeding - Major | 112000001889, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, COVID-19 Positive |

112000001982, Device Embolization | 112000001324, Device Thrombosis | 112000001839, Device Related Event - Other | 112000001828, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, Myocardial Infarction | 22298006, Permanent Pacemaker | 449397007, Readmission - Cardiac (Not Heart Failure) | 112000001897, Readmission - Heart Failure | 112000001896, Readmission - Non-Cardiac | 112000001898, Reintervention - Mitral Valve | 112000001893, Single Leaflet Device Attachment | 112000001538, Stroke - Ischemic | 422504002, Stroke - Hemorrhagic | 230706003, Stroke - Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 112000000460, Vascular Complication - Minor | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Selections

Element: 12933 Follow-up Event Name

Value Set Name: Follow Up Events OID: 1.3.6.1.4.1.19376.1.4.1.6.5.356

Selections Selection Dependency Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TAVR)

Atrial Fibrillation | 49436004, Bleeding - Life Threatening | 112000000459, Bleeding - Major | 112000001889, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, COVID-19 Positive | 112000001982, Device Embolization | 112000001324, Device Fracture |

112000001891, Device Thrombosis | 112000001839, Dialysis (New Requirement) |

100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Myocardial Infarction | 22298006, PCI | 415070008, Permanent Pacemaker | 449397007, Readmission - (Non-Valve Related) | 112000001895, Readmission (Valve Related) | 112000001894, Reintervention - Aortic Valve | 112000001827, Stroke - Ischemic | 422504002, Stroke - Hemorrhagic | 230706003, Stroke -

Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 112000000460, Vascular Complication - Minor | 112000001823,

Vascular Surgery or Intervention - Unplanned | 112000000467

Element: 12933 Follow-up Event Name

Value Set Name: Follow Up Events OID: 1.3.6.1.4.1.19376.1.4.1.6.5.356

> Selections **Selection Dependency**

Atrial Fibrillation | 49436004, Bleeding - Life Threatening | 112000000459, Bleeding - Major | 112000001889, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, COVID-19 Positive | 112000001982, Deep Vein Thrombosis | 128053003, Device Embolization | 112000001324, Device Fracture | 112000001891, Device Migration | 370512004, Device Thrombosis | 112000001839, Device Related Event - Other | 112000001828, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Myocardial Infarction | 22298006, PCI | 415070008, Permanent Pacemaker | 449397007, Pulmonary Embolism | 59282003, Readmission - (Non-Valve Related) | 112000001895, Readmission (Valve Related) | 112000001894, Reintervention - Tricuspid Valve | 112000001820, Stroke -Ischemic | 422504002, Stroke - Hemorrhagic | 230706003, Stroke - Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 112000000460, Vascular Complication - Minor | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Transcatheter Valve Therapy Reference Procedure Type (13705) IN (Tricuspid Valve Procedure)

Element: 12933 Follow-up Event Name

Value Set Name: Follow Up Events OID: 1.3.6.1.4.1.19376.1.4.1.6.5.356

> Selections Selection Dependency Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TMVR)

ASD Defect Closure due to Transseptal Catheterization | 112000001885, Atrial Fibrillation | 49436004, Bleeding - Life Threatening | 112000000459, Bleeding - Major | 112000001889, Cardiac Surgery or Intervention - Other Unplanned | 112000001892, COVID-19 Positive | 112000001982, Device Embolization | 112000001324, Device Fracture | 112000001891, Device Migration | 370512004, Device Thrombosis | 112000001839, Device Related Event -Other | 112000001828, Dialysis (New Requirement) | 100014076, Endocarditis | 56819008, ICD | ACC-NCDR-ICD, Myocardial Infarction | 22298006, Permanent Pacemaker | 449397007, Readmission - Cardiac (Not Heart Failure) | 112000001897, Readmission - Heart Failure | 112000001896, Readmission - Non-Cardiac | 112000001898, Reintervention - Mitral Valve | 112000001893, Stroke - Ischemic | 422504002, Stroke - Hemorrhagic | 230706003, Stroke -Undetermined | 230713003, Transient Ischemic Attack (TIA) | 266257000, Vascular Complication - Major | 112000000460, Vascular Complication - Minor | 112000001823, Vascular Surgery or Intervention - Unplanned | 112000000467

Discharge Location After Event Element: 14420 Value Set Name: Discharge Location OID: 1.3.6.1.4.1.19376.1.4.1.6.5.41

Selections **Selection Dependency**

Home | 01, Skilled Nursing Facility | 03, Extended Care/TCU/Rehab | 62, Other Discharge Status (14387) IN (Alive)

Location | 100001249

Element: 14406 Mitral Valve Reintervention Indication

Value Set Name: Valve Reintervention Indication OID: 1.3.6.1.4.1.19376.1.4.1.6.5.720

Selections **Selection Dependency** Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TMVr, TMVR)

Regurgitation | 40445007, Stenosis | 44241007, Device Embolization | 112000001324, Endocarditis | 56819008, Device Thrombosis | 112000001839, Valve Injury | 762610001, Other | 100000351

Follow-Up Medications Code Element: 11990

Value Set Name: Follow-up Medication OID: 2.16.840.1.113883.3.3478.6.5.203

Selection Dependency

Direct thrombin inhibitor | 414010005, Warfarin | 11289, P2Y12 Antagonist | 112000001003, Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TAVR) Aspirin | 1191, Direct Factor Xa Inhibitor | 112000000696

Element: 11990 Follow-Up Medications Code

Value Set Name: Follow-up Medication OID: 2.16.840.1.113883.3.3478.6.5.203

Selection Dependency Selections Transcatheter Valve Therapy Reference Procedure Type (13705) IN (TMVr, TMVR, Tricuspid

Valve Procedure)

Angiotensin Converting Enzyme Inhibitor | 41549009, Aldosterone Antagonist | 372603003, Direct thrombin inhibitor | 414010005, Warfarin | 11289, P2Y12 Antagonist | 112000001003, Aspirin | 1191, Angiotensin II Receptor Blocker | 372913009, Beta Blocker | 33252009, Thiazides | 372747003, Diuretics Not Otherwise Specified | 112000001417, Loop Diuretics |

29051009, Direct Factor Xa Inhibitor | 112000000696

Effective for Patient Discharged January 01, 2021

STS/ACC TVT Registry

Castion Cont	Section Containment Structure			cifications
Container Class	Section Structure	Section Code	Section Type	Cardinality
Container Class	Section	Jestion code	Section Type	Cardinality
patientContainer	Demographics	DEMOGRAPHICS	Section	1 1
episodeContainer	Episode of Care	EOC	Section	1 1
episodeContainer	Episode Information	EOCINFO	Section	1 1
episodeContainer	Admitting Providers	ADMTPROV	Section	01
episodeContainer	Attending Providers	ATTNPROV	Repeater Section	0 n
episodeContainer	Research Study	RSTUDY	Repeater Section	0 n
episodeContainer	History and Risk Factors	HISTORYANDRISK	Section	11
episodeContainer	Home Medications	HOMEMEDS	Repeater Section	0 n
episodeContainer	Condition History	CONDHIS	Repeater Section	1 n
episodeContainer	Condition History Details	CONDHISTDET	Section	01
episodeContainer	Atrial Fibrillation	AFib	Section	01
episodeContainer	Atrial Flutter	AFLUTTER	Section	01
episodeContainer	Carotid Artery Stenosis	CASTENOSIS	Section	01
episodeContainer	Cardiomyopathy	CARDIOM	Section	01
episodeContainer	Chronic Lung Disease	CLUNGD	Section	01
episodeContainer	Diabetes Therapy	DIABTHER	Section	0 1
episodeContainer	Endocarditis	ENDOCTIS	Section	0 1
episodeContainer	Myocardial Infarction	MITMEFME	Section	0 1
episodeContainer	Procedure History	PROCHIST	Repeater Section	1 n
episodeContainer	Procedure History Details	PROCHISTDET	Section	0 1
episodeContainer	Aortic Valve Replacement	AVREPL	Section	0 1
episodeContainer	Transcatheter AV Replacement	TRAVREPLIMP	Section	0 1
episodeContainer	ICD	ICD	Section	0 1
episodeContainer	Mitral Valve Annuloplasty	MVANUPLSTY	Section	0 1
episodeContainer	Mitral Valve Replacement	MVREPLC	Section	0 1
episodeContainer	Mitral Valve Transcatheter	MVTRANS	Section	0 1
episodeContainer	Permanent Pacemaker	PERMPACE	Section	0 1
episodeContainer	Tricuspid Valve Repair Surgery	TVREPAIR	Section	0 1
episodeContainer	Tricuspid Valve Intervention	TVINTVN	Section	0 1
episodeContainer	Tricuspid Valve Replacement Surgery	SURTVREPL	Section	0 1
episodeContainer	Transcatheter TV Replacement	TTVREPLC	Section	0 1
episodeContainer	Lab Visit	labvisit	Repeater Section	1 n
episodeContainer	Presentation and Evaluation	PREEVAL	Section	11
episodeContainer	STS Risk Score	STSRISK	Repeater Section	0 n
episodeContainer	Shared Decision Making	SDM	Section	01
episodeContainer	KCCQ12	BASEKCCQ	Section	01
episodeContainer	Five Meter Walk Test	FIVEMWT	Repeater Section	0 n
episodeContainer	Six Minute Walk Test	SIXMWT	Section	01
episodeContainer	Pre-Procedure Clinical Data	PREPROCLABS	Section	0 1
episodeContainer	Pre-Procedure ECG and Pulmonary Function	PREPROCPULMONARY	Section	0 1
episodeContainer	Pre-Procedure Medication(s)	PREPROCMED	Section	0 1
episodeContainer	Pre-Procedure Diagnostic Cath Findings	PREPROCDX	Section	0 1
episodeContainer	Pre-Procedure CTA Findings	PREPROCCTA	Section	0 1
episodeContainer	Pre-Procedure Echocardiogram Findings	PREPROCECHO	Section	0 1
episodeContainer	Left Ventricular Ejection	LVEF	Section	0 1
episodeContainer	Left Ventricular Dimension	LVEFDIM	Section	0 1
episodeContainer	Left Atrial Volume	LEFTATVOL	Section	0 1
episodeContainer	Aortic Valve Disease Etiology	ARVALETIOLOGY	Section	0 1
episodeContainer	Mitral Valve Disease	MVDisease	Section	0 1
episodeContainer	Mitral Valve Disease Etiology	MVEtiology	Section	0 1
episodeContainer	Tricuspid Valve Disease Etiology	TMVEtiology	Section	0 1
episodeContainer	Pre-Procedure Dobutamine Challenge	DOBUSTTST	Section	0 1
episodeContainer	Procedure Information	PROCINFO	Section	11
episodeContainer	Operator Information	OPRTRINFO	Repeater Section	0 n
episodeContainer	Radiation and Contrast	RADIATION	Section	0 1
episodeContainer	Post Implant Mitral Valve Data	POSTIMPMV	Section	01
episodeContainer	TAVR	TAVR	Section	0 1
episodeContainer	TAVR Devices	TAVRDEV	Repeater Section	0 n
episodeContainer	TMVr	TMVRpr	Section	01
episodeContainer	Mitral Leaflet Devices	MLEAFDEVICES	Repeater Section	0 n
episodeContainer	TMVR	TMVR	Section	01
episodeContainer	TMVR Devices	TMVRDEVICES	Repeater Section	0 n
episodeContainer	TTVP	TTVP	Section	01
episodeContainer	TTVP Pre-Implant	TTVPPREIMP	Section	01
episodeContainer	TTVP Post-Implant	TTVPPOSTIMP	Section	01
episodeContainer episodeContainer	TTVP Devices	TTVPPOSTIME	Repeater Section	0 n
episodeContainer	Post-Procedure - Intra or Post-Procedure Events	POPEVENTS	Repeater Section	1n
episodeContainer episodeContainer	In-Hospital Event Information	HOSPEVEADJ	Repeater Section	0 n
opioodeoonianiei	in Hoopital Event information	HOSI EVEADS	Nepeater Section	0 11

STS/ACC TVT Registry Data Dictionary v3.0 Full Specifications

Container Class	Section	Section Code	Section Type	Cardinality
episodeContainer	Stroke Or TIA	SRKRTIA	Section	0
episodeContainer	AV Re-Intervention	AVREINTVN	Section	0
episodeContainer	MV Re-Intervention	MVREINTVN	Section	0
episodeContainer	Tricuspid Valve Re-Intervention	TTVRREINTVN	Section	0
episodeContainer	Post-Procedure	POSTPROC	Section	0
episodeContainer	Post-Procedure Clinical Data	POPCLIDATA	Section	0
episodeContainer	Post-Procedure Hemoglobin	POSTPROCHEM	Section	0
episodeContainer	Post-Procedure 12 Lead	POSTPROC12L	Section	0
episodeContainer	Post-Procedure Creatinine	POSTPROCRT	Section	0
episodeContainer	Post-Procedure Highest Creatinine	POPROCHIGHCR	Section	0
episodeContainer	Post-Procedure Echocardiogram Findings	POSTPROCECHO	Section	0
episodeContainer	Post-Procedure AV Regurgitation	POPAVREG	Section	0
episodeContainer	Post-Procedure MV Regurgitation	POPMVREG	Section	0
episodeContainer	Post-Procedure TV Regurgitation	POPTVREG	Section	0
episodeContainer	Discharge	DISCHARGE	Section	1
episodeContainer	Discharge Medications	DISCMED	Repeater Section	0
followupContainer	Follow Up	FOLLOWUP	Section	1
followupContainer	Follow-Up Clinical Assessment	FUPCLINASMT	Section	0
followupContainer	Follow-Up Echocardiogram	FUPECHO	Section	0
episodeContainer	Follow-Up Imaging	IMGPERF	Section	0
episodeContainer	Follow-Up Aortic Valve	AVVALVE	Section	0
followupContainer	Follow-Up AV Regurgitation	FPOPAVREG	Section	0
episodeContainer	Follow-Up MV Imaging	MVIMG	Section	0
followupContainer	Follow-Up MV Regurgitation	FPOPMVREG	Section	0
episodeContainer	Follow-Up TV Imaging	TVREG	Section	0
episodeContainer	Follow-Up TV Regurgitation	FPOPTVREG	Section	0
followupContainer	Follow-Up 4DCTA	FCTAFindings	Section	0
followupContainer	Follow-Up Six Minute Walk Test	FSIXMIN	Section	0
followupContainer	Follow-Up KCCQ	FKCCQ	Section	0
followupContainer	Follow-Up Events	FUPEVENTS	Repeater Section	0
followupContainer	Follow-Up Event Information	FADJ	Repeater Section	0
followupContainer	Follow-Up Stroke or TIA	FSTRKTIA	Section	0
followupContainer	Follow-Up AV Re-Intervention	FAVREINTVN	Section	0
followupContainer	Follow-Up MV Re-Intervention	FMVREINTVN	Section	0
followupContainer	Follow-up Readmission	FREADMISSION	Section	0
followupContainer	Follow-Up Tricuspid Valve Re-Intervention	FTTVRREINTVN	Section	0
followupContainer	Follow-Up Medications	FUPMEDS	Repeater Section	0
submissionInfoContair	•	ADMIN	Section	1



STS/ACC TVT Registry

Reference Code System Listing

Code System	Oid
ACC NCDR	2.16.840.1.113883.3.3478.6.1
United States Social Security Number (SSN)	2.16.840.1.113883.4.1
HL7 Race	2.16.840.1.113883.5.104
HL7 Ethnicity	2.16.840.1.113883.5.50
SNOMED CT	2.16.840.1.113883.6.96
LOINC	2.16.840.1.113883.6.1
ACC NCDR EP Devices	2.16.840.1.113883.3.3478.6.1.21
ACC NCDR Lead Devices	2.16.840.1.113883.3.3478.6.1.20
ACC NCDR Catheter Ablation Devices	2.16.840.1.113883.3.3478.6.1.22
PHDSC	2.16.840.1.113883.3.221.5
HL7 Administrative Gender	2.16.840.1.113883.5.1
HL7NullFlavor	2.16.840.1.113883.5.1008
HL7 Discharge disposition	2.16.840.1.113883.12.112
RxNorm	2.16.840.1.113883.6.88
USPostalCodes	2.16.840.1.113883.6.231
ACC NCDR Intracoronary Devices	2.16.840.1.113883.3.3478.6.1.101
Center for medicare and medicaid services, MBI	2.16.840.1.113883.4.927
clinicaltrials.gov	2.16.840.1.113883.3.1077