

Integrating the Healthcare Enterprise



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IHE Cardiology Technical Framework Supplement

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Cath Report Content (CRC)

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Trial Implementation

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Foreword

This is a supplement to the IHE Cardiology Technical Framework V4.0. Each supplement undergoes a process of public comment and trial implementation before being incorporated into the volumes of the Technical Frameworks.

30 This supplement is published for Trial Implementation on July 25, 2012 and may be available for testing at subsequent IHE Connectathons. The supplement may be amended based on the results of testing. Following successful testing it will be incorporated into the Cardiology Technical Framework. Comments are invited and may be submitted at <http://www.ihe.net/cardiology/cardiologycomments.cfm>.

35 This supplement describes changes to the existing technical framework documents and where indicated amends text by addition (**bold underline**) or removal (~~**bold strikethrough**~~), as well as addition of new sections introduced by editor’s instructions to “add new text” or similar, which for readability are not bolded or underlined.

40 “Boxed” instructions like the sample below indicate to the Volume Editor how to integrate the relevant section(s) into the relevant Technical Framework volume:

<i>Replace Section X.X by the following:</i>
--

General information about IHE can be found at: www.ihe.net

45 Information about the IHE Cardiology domain can be found at: <http://www.ihe.net/Domains/index.cfm>

Information about the structure of IHE Technical Frameworks and Supplements can be found at: <http://www.ihe.net/About/process.cfm> and <http://www.ihe.net/profiles/index.cfm>

The current version of the IHE Technical Framework can be found at: http://www.ihe.net/Technical_Framework/index.cfm

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Introduction to this Supplement

180 This Supplement introduces a new Profile to the IHE Cardiology Technical Framework, with the overall design in Volume 1 and specific content in Volume 3. This Profile relies heavily on Content Profile concepts specified in the IHE Patient Care Coordination Technical Framework and CDA templates specified in the HL7 Implementation Guide for CDA Release 2: IHE Health Story Consolidation, Release 1 DSTU (December 2011) (C-CDA).

185 This content profile is motivated by cardiologists, who face an increasing demand from patient-care, data-quality and legislative perspectives to increase the usefulness and actionability of (discrete) clinical data across the various care-settings and stakeholders.

A solution for such interoperability is, however, not a simple undertaking. Unstructured textual data forms remains the predominate mechanism for information exchange among health care
190 providers, and a good majority of data needed by physicians and other health care providers to make good clinical decisions is embedded in this free text. Efficient and effective interoperability therefore begins by identifying the most relevant clinical data.

Clinically-relevant Cardiac Cath lab data is the key value proposition of this profile. The approach is to:

- 195 1. reuse the distribution and structuring work from the XDS (ITI domain), Medical summaries (PCC domain), and the HL7 Implementation Guide for CDA Release 2: IHE Health Story Consolidation, Release 1 DSTU (C-CDA) for exchangeable procedure notes (HL7/IHE)
- 200 2. extend it through adding and codifying the ACC-NCDR Cath/PCI dataset and the ACC NCDR Cath/PCI version 4.4 to clinical data standards like ICD9/10, Snomed and LOINC
3. evaluate it as it applies to the work in progress by the ACC and AHA to create an “ACCF-AHA Cardiac Cath Reporting – Report template”

The aim is to enable collection and distribution of the most clinically-relevant discrete data on
205 the cardiac catheterization procedures common in cardiology. The usage of the discrete data is three-fold:

1. To enable individual Cath/PCI procedures to be more easily shared and used between care givers and systems
- 210 2. To enable population-based outcomes-based research on procedure effectiveness
3. To provide the ability to interact with data registries for data exchange.

These Cath/PCI procedures are used as key constituents of the patient’s treatment during cardiac encounters and disease management. Allowing a means to extract and exchange key cardiac measures across providers and their systems will be a huge advantage to providing a complete, accessible and actionable cardiac data set in front of cardiologists.

- 215 There are very successful quality-improvement programs in place by the professional bodies such as the ACC, AHA, and state registries concentrating on the most invasive, and expensive cardiac procedures. However, the effort in translating and extracting the discrete data required by these registries still involves significant manual work and inefficiencies due to the absence of a standardized structuring of information at the point of clinical reporting.
- 220 This supplement provides a framework to make progress in these areas. This profile codifies representative areas of procedure indications, procedures, medications, observations, complications and findings for invasive cardiac procedures and specifies how this discrete data can be organized to be used by both care-providers and automated data processing systems.

225 **Relationship to Workflow Profiles**

Cath Report Content (CRC) is a *content* profile – it is agnostic with respect to the workflow or data exchange mechanism in which the data is produced and handled.

- 230 Content Profiles define how the content used in a transaction is structured. The binding of the Content to an IHE transaction that is part of an IHE Workflow Profile specifies how this payload may influence the metadata or the behavior of the transaction. Content modules within the Content Profile then define the payloads. Content modules are transaction neutral, in that what they describe is independent of the transaction in which they are used, whereas content bindings explain how the payload influences the transaction metadata and/ or behavior.

- 235 The CRC content is intended to be deployed, for example in the Displayable Reports (DRPT) workflow profile for in-patient environments, or the Cross-Enterprise Document Sharing (XDS) profile to propagate the content across organizational boundaries.

It is important to note that that key report-generation/distribution workflow aspects such as physician identification, insurance preauthorization, report routing and acknowledgement, and patient consent, are **out of scope** for this Content Profile.

240

Cardiac Cath Report Content (CRC) Profile

The Cardiac Cath Report Content (CRC) Profile specifies the content structure for a clinical report of a Cardiac Cath imaging exam, recorded in a DICOM Study. Such exams include:

- Diagnostic Cath
- 245 • Angiography
- PCI

The CRC Profile specifies the use of an HL7 Clinical Document Architecture (CDA) format for the report. This format supports both the human readable narrative historically used for clinical reports, as well as a substantial set of discrete data elements that may be used for longitudinal or population analysis or other computer processing.

Note: It is expected that future evolution of cardiology reporting will incorporate more robust and extensive sets of data elements. This Profile is a first step beyond simple narrative to interoperable discrete data elements.

Not included in the scope of this profile are electrophysiology procedures, and non-cardiology procedures (e.g., peripheral angiography). Pediatric Cath cases are also excluded. Such use cases may be supported by other similar content profiles.

It is assumed that there is a DICOM Study associated with the exam. If there is not a DICOM Study, this report content may not be appropriate. In addition to reference images, the DICOM Study data may include discrete data elements encoded in DICOM Structured Report information objects that may be transcoded into the discrete data elements specified in this Profile. (See the Evidence Documents Profile and its cardiology options in Section 7.)

The CRC Profile does not presume to describe the complete content of a cardiac catheterization laboratory report. It does provide the framework of high level section titles and a set of discrete data elements. Within that framework reports can be created with the clinical content desired by their authors, including additional discrete data elements. In general, there are no constraints on the narrative text and figures that the cardiologist could include in the report document, although there are requirements on minimum data elements reflecting expert consensus (ACC-NCDR Cath/PCI, ACCF-AHA Cardiac Cath Reporting workgroup).

This profile also does not provide all of the details necessary to construct a CDA compliant document. Please refer to the HL7 CDA Release 2 Standard.

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Open Issues and Questions

#	Open Issue Description
2	Generally, certain concepts represented by SNOMED compositions have been submitted for pre-coordinated codes. In CIRC, compositional coding was used to represent some concepts and there are still some in this profile. Requests were submitted to SNOMED for pre-coordinated codes that represent these compositions with a single code and we're

#	Open Issue Description
	waiting for these single codes to be provided by SNOMED. A: There is a need for one code (mid right coronary artery)– and it should be available in July 2012. This issue will remain open until the code is available.
38	Depth vs. breadth presentation of sections and entries – the sections and entries are presented in Volume 3 per the order they are included in a clinical CRC report (as an instance of a CRC CDA document) (depth view) rather than organizing the sections by listing them ordered alphabetically and having the complete set of entries used within this document organized in a separate list ordered alphabetically (breadth view). Is this approach useful or preferred vs. the breadth approach?

Closed Issues

#	Closed Issue Description/ Resolution
1	<i>Which document do we use on which to base this profile, HL7 Consolidated CDA DSTU or the IHE_CARD_Suppl_CIRC_Rev1.1_TI 2011-06-24 content profile.</i> A: The committee agreed to use the HL7 Implementation Guide for CDA Release 2: IHE Health Story Consolidation, Release 1 DSTU (Dec 2011 edition).
2a	Need to handle multiple performers for header, including authors and legal authenticators. A: Header either has allowances for multiple performers and has been expanded to include multiple authors and authenticators as need but is limited to a single legal authenticator as constrained by CDA.
3	How do we associate medical equipment with Procedures. A: Medical devices are described in the Procedures Description Cardiac Section.
4	What to do with US realm race code(7263), ethnicity(5323), do we really need race or ethnicity? A: All US realm specifics should be reflected in section (or Volume?) 4 of this content profile which will be done at a later date.
5	How do we associate procedures and findings, particularly coronary segments? A: We are using a Lesion ID (as an instance of a result observation “id”) to link findings to procedures. The Lesion ID is associated with a coronary segment.
6	In medical history, do we need past and present illness? A: NO. The ACCF-AHA Cardiac Cath Reporting - ReportTemplate does not separate illness by past and present, and committee experience shows no real reason to separate these out, especially since there is an allowance to record if a problem is active in the Medical History Section.

#	Closed Issue Description/ Resolution
7	<p>How do we record more than one piece of information for a data concept, like a table format versus straight text in the rendered report?</p> <p>A: We can have structured elements in the coded data and if these elements appear in the narrative, they must be accurately rendered. In the narrative text, any valid HTML markup can be used, including markup used to render a table in HTML.</p>
8	<p>Q - Does the text narrative for every CDA construct need to completely contain the full coded content.</p> <p>A: If the ACT relationship is DRIV, then the narrative is based solely on the coded content. But does this need to include all the coded content?</p> <p>Technically it is not required, but all coded content SHOULD be included in the narrative. There SHALL be no conflicts between the narrative and the coded content. The coded content may not be an equivalent of the narrative.</p> <p>This has been added as a note in section 6.3.1.4 (Conventions).</p>
8a	<p>What do we do with data fields that have coded values potentially cover multiple Snomed codes, for example New York Class and Angina Class.</p> <p>A: We can use multiple SNOMED codes to describe the value needed based on CCDA.</p>
9	<p>CIRC has an Indications and Planned Procedure section and CCDA separates these into separate sections.</p> <p>A: We have adopted the C-CDA approach and will have a separate Procedure Indications Section and a Planned Procedure Section.</p>
10	<p>Some of these concepts identified in the NCDR CathPCI Registry v4.4 Coder’s Data Dictionary do not seem to be “Yes/No” valued. Do they belong in the Complications section?</p> <p>A: No, only concepts that are SNOMED CT “findings” should be included in this section. This also includes disorders (which are also findings).</p> <ol style="list-style-type: none"> a. Concepts that are procedures should be included in other sections b. Items that are measurements should be associated with the appropriate procedures c. If there are appropriate findings concepts for these procedures, these will be included in the complications list <ol style="list-style-type: none"> i. Renal failure ii. Anemia due to blood loss
11	<p>Is there a need to create a specialization of the Problem Observation entry to reflect that there will not be either a Health Status entry or an Age Observation entry?</p> <p>A: We created a specialization of the problem observation entry to also include severity so we can set the cardinality of these other entries as needed for this specific use.</p>

#	Closed Issue Description/ Resolution
12	<p>Is this content profile for US realm only or for Universal realm?</p> <p>A: This CRC profile is realm agnostic and could be further extended for US-realm, Universal realm or any national extension.</p>
13	<p>How do we link Procedure Findings.Result Observation and Procedures.Procedure Activity Procedure entries?</p> <p>A: There is a new lesion ID which is intended to be used to link a Procedure Findings.Result Observation and Procedures.Procedure Activity Procedure entries which are for the same lesion.</p>
14	<p>Q - Can we use the segmental wall analysis, from IHE_CARD_Suppl_CIRC_Rev1.1_TI 2011-06-24 in a revised format to fit drawings from ACCF-AHA Cardiac Cath Reporting - ReportTemplate and other drawings performed by vendors?</p> <p>A: Yes, it is possible to include graphical representations (e.g., drawings) of coronary anatomy and segmental wall analysis to be embedded in the CDA document to be either embedded in-line or referenced via a URL. These are allowed at the document summary level.</p>
15	<p>Q - Which document content section should be used to record bruits(femoral,carotid)?</p> <p>A: These are recorded in the Vital Signs Section, see Value Set 1.3.6.1.4.1.19376.1.4.1.5.36</p>
16	<p>Q - Do we need an anesthesia section (e.g., for aortic valve replacement done in cath lab under general anesthesia) ?</p> <p>A: We have the ability to record Local anesthetics and sedation administration in the Medications Administered Section if needed and have included an Anesthesia section to handle all other types of anesthesia</p>
17	<p>Q - The ACCF-AHA Cardiac Cath Reporting - ReportTemplate shows ICD9 coding sections for both pre and post diagnosis. How are postprocedure diagnoses determined and are the ICD-9 codes actually available at the time of producing this procedure note?</p> <p>A: After discussion, it was determined that some systems will have this information available at the time of the Procedure Note, so we have included language in preprocedure and postprocedure diagnosis to allow the inclusion of ICD9 coding.</p>

#	Closed Issue Description/ Resolution
18	<p>If grafts were performed, the ACC coding only requires stenosis to be recorded for each of these systems. In the real world, a methodology should be used that uniquely identifies a graft and its related stenosis which can then be used for both Cath Lab and OR cardiac procedures.</p> <p>Q – do you agree?, and is this addressed in the ACCF-AHA Cardiac Cath Reporting – ReportTemplate?</p> <p>A: The ACCF-AHA Cardiac Cath Reporting – ReportTemplate does not reflect grafts in the samples but discussion centered on including graft descriptions that include an origin, type of conduit, and insert site. This has been allowed for by including these descriptions in text format by the Content Creator as part of the narrative for the Procedure Description Section (or any other section).</p>
19	<p>Q - In the Medical History section, is “health status observation” for the patient required or used? Or is “problem status” sufficient for each problem observation?</p> <p>A: C-CDA 1.1 provides a new value set for health status observation that is meaningful. For this initial CRC version, this will be available for use, if needed/desired.</p>
19a	<p>Q – can we have a shared single code for identifying cath and PCI document types?</p> <p>A: We have adopted Cath, PCI, and Cath/PCI document types for this content profile.</p>
20	<p>In the Medical History Section, should prior procedures be moved to the history section of Procedures. Logically they fit in this section but CCDA has a section for history in the procedures area.</p> <p>Q – does it make most clinical sense to put prior procedures in the medical history section of the report?</p> <p>A: Prior procedures have been included in the Medical History section.</p>
21	<p>Q – is there a good source of Cath Procedure Findings you can recommend?</p> <p>A: We have created a list of Procedure Findings that is extensible, which means it can be expanded as needed.</p>
22	<p>Q - Is this the complete set of complications that should be included in this profile? Are there other specific complications that should be added?(Complication Section)?</p> <p>A: We have included ACC-NCDR Cath/PCI complication values as a starting point in this extensible table. Expansion to other complications is at the Content Creator's discretion.</p>
23	<p>Q -Is There a need for the Procedures Specimens Taken section?</p> <p>A: Yes, the Section has been added and is based on the C-CDA definition. It can be used to handle biopsy and other specimens.</p>

#	Closed Issue Description/ Resolution
24	<p>Q - Is there a need for the Procedure Implants section?</p> <p>A: No. This is required for EP and is out of scope for this CRC profile.</p>
25	<p>Q - For the Medical History section, is there a need to identify the “severity” of the problem?</p> <p>A: Yes, possibly so we have added a “Severity Observation” entry to the Problem Observation entry in this section.</p>
26	<p>We are working with draft documents from the ACCF-AHA Cardiac Cath Reporting - ReportTemplate. We will need to revise our profile when this document is final.</p> <p>Q - When is expected completion?</p> <p>A: Per discussions at the PC F2F @ACC it will be available in 6 months from 3/25/2012.</p>
27	<p>Q - How is Hematoma size best represented clinically ? This is not a complication but could be related to a complication.(Complication Section)</p> <p>A: This should be treated a result observation related to the particular Problem Observation in the Complication Section. This is out of scope for this profile version.</p>
28	<p>Vol 1 - Section 12.3.1 – Should there be a binding to the IEO profile? Technically there could be a binding to IEO, but practically it is questionable. IEO is targeted for cardiology practice offices.</p> <p>A:This profile is not targeted for the ambulatory setting so there should not be a binding to the IEO profile. Text referencing IEO has been removed.</p>
29	<p>Need to remove C-CDA sections/entries that are unchanged. Also need to harmonize definitions of entries across sections within this profile.</p> <p>A: Done</p>
30	<p>Need to assign IHE Card specific template IDs for new/changed entries and vocabulary constraints. Entries highlighted in table 6.3.3-1 need IHE Card specific template IDs.</p> <p>A: New template IDs were created for the CRC specific sections and entries.</p>
31	<p>Need better xml samples for all sections/entries. It would be useful to create XML of a complete sample report.</p> <p>A: Done</p>
32	<p>IVUS/IVOCT procedures are mention in the intro section as being in scope. No specific measurements are described elsewhere in the content specification. Should there be additional specific measurements for IVUS/IVOCT or should the intro section be modified to remove this from the scope for this profile?</p> <p>A:Removed references to IVUS/IVOCT from this specification. It is believed that</p>

#	Closed Issue Description/ Resolution
	support for IVUS/IVOCT requires additional vocabulary which could be added (via a CP) after complete analysis is done.
33	Need to assign unique constraint IDs to CRC specific constraints. A: This will be done by tooling from MDHT (when available) as part of the CDA template development process.
34	The intention is to have the CRC specific vocabulary sets be extensible and also not be based on a specific version of the vocabulary standard (e.g., SNOMED). This profile specifies all the CRC specific vocabulary sets as STATIC. The value sets can be extended where they are designated for use for a specific element specified as a CWE data type.
35	There are problems with C-CDA specification that should be submitted to HL7 and addressed by HL7. Need to compile list of these problems and IHE Card will submit them. A: A list of C-CDA errata was developed and submitted to HL7.
36	Q: Should the cardinality for Legal Authenticator be [0..1] or [1..1]. A:[1..1] because this profile does not support the exchange of preliminary unapproved reports (non – legally authenticated).
37	There is no code in Table 6.3.6.11 selected for Antiarrhythmics: Azimilide A: this is a general class 3 Antiarrhythmics (potassium channel antagonist), but no specific code was found. This entry was removed from the table.

Volume 1 – Profiles

12 Cath Report Content Profile (CRC)

280 The Cath Report Content (CRC) Profile specifies the content structure for a clinical report of a cardiology procedure recorded in a Cardiac Cath Lab. Such procedures include:

- Diagnostic Cath
- Angiography
- PCI

285 The CRC Profile specifies the use of an HL7 Clinical Document Architecture (CDA) format for the report.

Not included in the scope of this profile are imaging studies (e.g., echocardiography), electrophysiology procedures, and non-cardiology procedures (e.g., peripheral angiography). Such use cases may be supported by other similar content profiles.

290 This profile also does not provide all of the details necessary to construct a CDA compliant document. Please refer to the HL7 CDA Release 2 Standard.

12.1 CRC Actors, Transactions, and Content Modules

295 Figure 12.1-1 shows the actors directly involved in the CRC Profile and the relevant transactions between them. There are two actors in this profile, the Content Creator and the Content Consumer. Content is created by a Content Creator and is to be consumed by a Content Consumer. The sharing or transmission of content from one actor to the other is addressed by the appropriate use of other IHE profiles, and is out of scope of this profile; hence there is no transaction per se defined for this profile.



Figure 12.1-1: Cath Report Template Actor Diagram

300 Note: The primary intended transmission mechanism in the intra-institutional context is the IHE Displayable Reports Profile (DRPT), and in the inter-institutional context the IHE Portable Data for Imaging (PDI) or IHE Cross Enterprise Document Sharing Profiles (XDS, XDM, and XDR). A Report Creator, Document Source or a Portable Media Creator of those profiles may embody the Content Creator Actor. A Document Consumer, a Document Recipient or a Portable Media Importer may embody the Content Consumer Actor.

305 **12.1.1 Actor Descriptions and Actor Profile Requirements**

12.1.1.1 Content Creator

1. A Content Creator shall be able to create a Cardiac Cath Lab Report Document according to the specifications for that content profile found in CARD TF-3.
- 310 2. A Content Creator shall be grouped with the Time Client Actor, and shall synchronize its clock with a Time Server.

12.1.1.2 Content Consumer

1. A Content Consumer shall be able to consume (receive and process) a Cardiac Cath Lab Report document.
- 315 2. A Content Consumer shall implement the View Option or Discrete Data Import option, or both.
3. A Content Consumer that implements the Document Import or Section Import Option shall implement the View Option as well.
4. A Content Consumer that implements the View option shall be able to:
- 320 a. Demonstrate rendering of the document for display.
- b. Print the document.
- c. Display the document with its original style sheet.
- d. Support traversal of any links contained within the document.
5. A Content Consumer that implements the Document Import Option shall:
- 325 a. Store the document.
- b. Demonstrate the ability to access the document again from that storage.
6. A Content Consumer that implements the Section Import Option shall offer a means to import one or more document sections into the patient record as free text.
7. A Content Consumer that implements the Discrete Data Import Option shall offer a means to import structured data from one or more sections of the document.

330 **12.2 CRC Actor Options**

Options that may be selected for this Content Profile are listed in Table 12.2-1 along with the Actors to which they apply. Dependencies between options when applicable are specified in notes.

Table X.2-1: CRC Profile Options

Actor	Option Name	Optionality	Section
Content Consumer	View Option	O (see 12.2.1)	PCC TF-2 :3.1.1
	Document Import Option	O (see 12.2.1)	PCC TF-2 :3.1.2
	Section Import Option	O (see 12.2.1)	PCC TF-2 :3.1.3

Actor	Option Name	Optionality	Section
	Discrete Data Import Option	O (see 12.2.1)	PCC TF-2 :3.1.4
Content Creator	<i>No options defined</i>		

335

12.2.1 Content Consumer Options

The Content Consumer actor is required to support at least one of the View or Discrete Data Import options. The Document Import and Section Import options, if implemented, also require the View option. These options as specified in the PCC Technical Framework assume use of XDS or related profiles for transport; this Profile specifies bindings to other workflow profiles (see Section 12.5), and these options should be interpreted as applicable with any binding. See also Section 12.6.2.

340

12.3 CRC Actor Required Groupings

The Content Creator shall be grouped with Time Client actor of the IHE IT Infrastructure Consistent Time Profile, as specified in ITI TF-1:7. This allows the Legal Authentication timestamp to be accurate.

345

Content modules describe the content of a payload found in an IHE transaction. Content profiles are transaction neutral. They do not have dependencies upon the transaction that they appear in.

12.4 CRC Document Content Module

There is often a very clear distinction between the transactions in a messaging framework used to package and transmit information, and the information content actually transmitted in those messages. This is especially true when the messaging framework begins to move towards mainstream computing infrastructures being adopted by the healthcare industry.

350

In these cases, the same transactions may be used to support a wide variety of use cases in healthcare, and so more and more the content and use of the message also needs to be profiled, sometimes separately from the transaction itself. Towards this end IHE has developed the concept of a Content Profile.

355

Content Profiles specify how the payload of a transaction fits into a specific use of that transaction. A content profile has three main parts. The first part describes the use case (this is found in Volume 1 in the definition of each Profile). The second part is a Content Module (found in this Volume 3), which describes the payload of the transaction; a content module is specified so as to be independent of the transaction in which it appears. The third part is binding to a specific IHE transaction, which describes how the content affects the transaction. The binding of CDA-based medical documents to workflow transactions is described in the Profile definition in Volume 1 (e.g., see IHE CARD TF-1:12.7).

360

365

12.5 CRC Overview

The Cardiac Cath Lab Report Content (CRC) Profile specifies the content structure for a clinical report of a cardiology Cath Lab Visit. Such procedures include:

- Diagnostic Cath
- 370 • Angiography
- PCI

The CRC Profile specifies the use of an HL7 Clinical Document Architecture (CDA) format for the cath physician clinical report. This format supports both the human readable narrative historically used for clinical reports, as well as a substantial set of discrete data elements that 375 may be used for longitudinal or population analysis or other computer processing.

Note: It is expected that future evolution of cardiology reporting will incorporate more robust and extensive sets of data elements. This Profile is a first step beyond simple narrative to interoperable discrete data elements.

Not included in the scope of this profile are imaging studies, ECG, electrophysiology procedures, and non-cardiology procedures (e.g., peripheral angiography). Such use cases may be supported 380 by other similar content profiles. This profile does not address how discrete data is collected and transmitted for registry data collection and other secondary usages.

It is assumed that there is a DICOM Study associated with the exam. If there is not a DICOM Study, this report content may not be appropriate. In addition to reference images, the DICOM Study data may include discrete data elements encoded in DICOM Structured Report 385 information objects that may be transcoded into the discrete data elements specified in this Profile. (See the Evidence Documents Profile and its cardiology options in Section 7.)

The CRC Profile does not presume to describe the complete content of an imaging study report. It does provide the framework of high level section titles and a set of discrete data elements. Within that framework reports can be created with the clinical content desired by their authors, 390 including additional discrete data elements. In general, there are no constraints on the narrative text and figures that the cardiologist could include in the report document, although there are requirements on minimum data elements reflecting expert consensus (ACC-NCDR Cath PCI data elements)

This profile also does not provide all of the details necessary to construct a CDA compliant 395 document. Please refer to the HL7 CDA Release 2 Standard.

12.5.1 Concepts

Not Applicable

12.5.2 Use Case #1: Compile and Transfer Report of Cardiac Cath Lab Procedure with Use of ACC-NCDR Cath/PCI Data Elements

400 **12.5.2.1 Compile and Transfer Report of Cardiac Cath Lab Procedure with Use of ACC-NCDR Cath/PCI Data Elements Use Case Description**

This use case addresses the generation and transfer of a cardiac cath lab report based on the NCDR CathPCI Registry v4.4 Coder's Data Dictionary data elements (see http://www.ncdr.com/WebNCDR/NCDRDocuments/CathPCI_v4_CodersDictionary_4.4.pdf).

405 The initial content, structure and coding of the report to support this use case are detailed as part of this profile (see IHE CARD TF-3: 6 Content Modules). However various reporting system implementations, institute reporting guidelines and individual Reporting Physician usage may result in some variability in the specific report content provided.

410 **12.5.2.2 Compile and Transfer Report of Cardiac Cath Lab Procedure with Use of ACC-NCDR Cath/PCI Data Elements Process Flow**

Pre conditions

415 The systems underlying the data collection and management for the various elements of the procedure report have all the *mandatory* data elements identified using codes, and are expected to be the source for the information used in creating the *majority* of the structured cath lab report document.

Main Flow

- Cardiologist reviews and/or records the codified
 - procedures and protocols used in the procedure
 - Data generated from the various modalities and monitoring equipment used during the procedure so that the key physiological measures, acquired and derived (pre, during and post intervention) are present in line with the ACC-NCDR Cath/PCI guidelines.
 - Other relevant patient characteristics
 - Medications documented for the patient both pre and during procedure.
 - Equipment used and implanted in the patient
 - Indications and observations/complications noticed during the procedure.
 - Findings, assessment and plan
- Cardiologist approves the procedure report and this marks it ready for distribution
- The content creator system will format the report appropriately (this profile) and send it via one of the IHE mechanisms to a content consumer system (an appropriate workflow profile).

430 **Post conditions**

The subsequent clinical stakeholder (system) receives the Document for import, processing and optionally viewing of the data.

12.5.3 Use Case #2: Perform Discrete Data-analysis on Procedure Report Content

435 **12.5.3.1 Perform Discrete Data-analysis on Procedure Report Content Use Case Description**

The goal of this use case is to assist data collection for comparative and research purposes. Based on a report generated in the previous use cases an advanced medical data analysis system collects discrete data from multiple patients and their procedure, e.g., for cardiac Clinical Decision Support or for advanced lifetime patient records.

440 **12.5.3.2 Perform Discrete Data-analysis on Procedure Report Content Process Flow**

Pre conditions

445 The content consumer (e.g., an advanced medical data analysis system) received a Cardiac Cath Lab Report with coded/structured content as defined in IHE CARD TF-3: 6 CDA Release 2 Content Modules.

Main Flow

The consuming system collects and processes the data from the various reports it receives and extracts those relevant data for either:

- 450 • A specific clinical concern for a population e.g., pre-populating a procedure-specific registry; extracting a data subset for a specific research question.
- A more comprehensive longitudinal patient record (e.g., an EMR) which can provide trending over time on an individual patient’s key cardiac measures.

Post conditions

455 The content consumer generated new (derived) data for use by others. The type of data generated is out of the scope of this *profile*.

12.5.4 Use Case #3: Review Procedure Report

12.5.4.1 Review Procedure Report Use Case Description

A secondary use-case addressed by this profile involves the direct human use of the procedure report. In most practical cases this will be:

- 460 • The referring physician who instigated/ordered the procedure, and other healthcare providers who manage subsequent patient care activities
- Another person involved in downstream clinical or administrative data processing e.g., someone validating/source-checking for QA the original report as part of JCAHO audits, or pre-submission checking on the original reporting data against the case-data imported in the
- 465 ACC-NCDR Cath/PCI registry-submission application

12.5.4.2 Review Procedure Report Process Flow

Pre conditions

- The Reviewing Physician consumer has a system (EMR or other) capable of importing and displaying the received report in a clinically useful format

- 470
- The Cath Lab Report has been received at this system

Note: This profile does not assume any explicitly specified relationship between the creator and consumer.

Main Flow

- The reviewing physician selects the report of his patient and opens it for review
- The system displays the human readable content for the reviewing physician to review

475

Post conditions

The Reviewer has extracted (visually) the necessary information from the report.

12.6 CRC Security Considerations

Security considerations are dealt with by the transport mechanism (e.g., XDS, DRPT) and are outside the scope of this content profile. See PCC TF-1: 3.8

480

12.7 CRC Cross Profile Considerations

A Content Creator or Content Consumer should be grouped with appropriate actors from workflow profiles that manage interchange of clinical data. Such groupings are described in this section.

- 485
- Content profiles may impose additional requirements on the transactions used when grouped with actors from other IHE Profiles. The metadata sent in the document sharing or interchange messages has specific relationships to the content of the clinical document described in the content profile. These mappings between the workflow metadata and the content attributes are described in IHE PCC TF-2:4.

12.7.1 Content Bindings for Displayable Reports (DRPT) Profiles

- 490
- CDA documents using the CRC content may be exchanged between a Report Creator and a Report Manager, as defined in the Displayable Reports (DRPT) Profile using the Encapsulated Report Submission [CARD-7] transaction. In this case, the CRC Content Creator actor is grouped with the DRPT Report Creator actor, and the CRC Content Consumer actor is grouped with the DRPT Report Manager actor.

495

12.7.2 Content Bindings for XDS, XDM, XDR, XDS-I, and XDR-I

It is expected that the transfers of care will occur in an environment where the physician offices and hospitals have a coordinated infrastructure that serves the information sharing needs of this community of care. Several mechanisms are supported by IHE profiles:

- 500
- A registry/repository-based infrastructure is defined by the IHE Cross Enterprise Document Sharing (XDS) and other IHE Integration Profiles such as patient identification (PIX & PDQ)

and notification of availability of documents (NAV). An extension for imaging study exchange is Cross Enterprise Document Sharing for Imaging (XDS-I).

- A media-based infrastructure is defined by the IHE Cross Enterprise Document Media Interchange (XDM) profile.
 - 505 • A reliable messaging-based infrastructure is defined by the IHE Cross Enterprise Document Reliable Interchange (XDR) profile. An extension for imaging study exchange is Cross Enterprise Document Reliable Interchange for Imaging (XDR-I).
 - All of these infrastructures support Security and privacy through the use of the Consistent Time (CT) and Audit Trail and Node Authentication (ATNA) profiles.
- 510 For more details on these profiles, see the IHE IT Infrastructure Technical Framework, and the IHE Radiology Technical Framework for XDS-I and XDR-I.

Document Source and Document Consumer Actors from the ITI XDS, XDM and XDR profiles are logically grouped with the CRC Content Creator and Content Consumer actors, respectively.

12.7.3 Binding for Portable Data for Imaging (PDI)

- 515 CDA documents using the CRC content may be exchanged on interchange media in accordance with the Portable Data for Imaging (PDI) profile. Such documents may be encapsulated within DICOM SOP Instances, or may be native CDA documents, as described in the IHE Radiology Technical Framework. In this case, the CRC Content Creator actor is grouped with the PDI Portable Media Creator actor, and the CRC Content Consumer actor is grouped with the PDI
- 520 Display or Portable Media Importer actors.

12.7.4 Content Binding for Retrieve Form for Data Capture (RFD)

- 525 A CDA document may be used for pre-population of a data entry form managed by actors of the Retrieve Form for Data Capture (RFD) Profile. In particular, the CRC content, as a carrier of discrete encoded data, may be used to pre-populate data entry forms for cardiovascular data registries. The CRC profile has been developed with key data elements that support common research related data fields. This profile, however, does not provide mapping between CRC field content and any specific registry field content. In this case, the CRC Content Consumer actor is grouped with the RFD Form Manager actor for the purpose of extracting discrete data from the report to pre-populate the data capture form.

530 12.7.5 Relationship to Document Digital Signature (DSG)

When a Content Creator Actor needs to digitally sign a document in a submission set, it may support the Digital Signature (DSG) Content Profile as a Document Source. When a Content Consumer Actor needs to verify a Digital Signature, it may retrieve the digital signature document and may perform the verification against the signed document content.

535

Appendices

Actor Summary Definitions

540

<i>Add the following terms to the IHE TF General Introduction Namespace list of Actors:</i>

None

Transaction Summary Definitions

<i>Add the following terms to the IHE TF General Introduction Namespace list of Transactions:</i>

None

545

Glossary

Add the following terms to the IHE Technical Frameworks General Introduction Glossary:

None

550

Volume 3 – Content Modules

Add section 6.3

6 Content Modules

6.3 Cath Report Content Modules

6.3.1 Cath Report Content Specification 1.3.6.1.4.1.19376.1.4.1.1.2

555 This is the template for Cardiac Cath/PCI Reports with discrete data elements as described in the NCDR CathPCI Registry version 4.4 Coder’s Data Dictionary. The Template ID for conformance to this template is OID = 1.3.6.1.4.1.19376.1.4.1.1.2.

This CDA document is not a direct specialization of any existing CDA document template ID. However, some parts were based on the IHE Card CIRC document specification and the HL7
560 Implementation Guide for CDA Release 2: IHE Health Story Consolidation, Release 1 DSTU - December 2011 (C-CDA) Procedure Note document specification.

6.3.1.1 Format Code

The XDSDocumentEntry format code for this content is **urn:ihe:card:CRC:2012**

565 The mapping of CDA header attributes to XDS metadata shall be identical to the XDS-MS mapping specified in PCC TF-2: 4.1.1.

6.3.1.2 Relationship to the IHE Cardiology CIRC Profile

This CRC document is inconsistent with the existing Cardiac Imaging Report Content (CIRC) content profile that was published for Trial Implementation in 2011.

These inconsistencies include:

- 570
- Overall document structure
 - This is not based on the IHE PCC section and entry templates but is based on the C-CDA section and entry templates.

6.3.1.3 Relationship to C-CDA

575 Some CDA section and entries used within this CRC document were based on the HL7 Implementation Guide for CDA Release 2: IHE Health Story Consolidation, Release 1 DSTU (C-CDA) section and entry definitions.

- a. Where constraints defined in C-CDA were not modified, the constraint remains as the C-CDA constraint identifier (e.g., CONF:5361). If only the value set was modified, then the constraint is considered unchanged.
- 580 b. Where constraints defined in C-CDA were modified, the original constraint ID is also modified by appending “-CRC” (e.g., CONF:5253-CRC). Modifications could include changes in the cardinality.

- c. Where new constraints were introduced, a new constraint identifier was defined (e.g., CONF:CRC-xxx)

585 If there are no new or modified constraints for a section or entry or if only the value sets are constrained, then the definition of the section or entry is considered unchanged from the C-CDA definition and the C-CDA template Id will be used. These unchanged sections/entries are referenced directly from the C-CDA specification and are not included in this specification.

590 If there are new or modified constraints for a section or entry, then that section or entry is assigned a new IHE Card specific template Id.

The description of the type of modification to affected section or entry content modules are outlined with boxes.

6.3.1.4 Conventions

6.3.1.4.1 Conformance Terms

595 The definitions of the conformance verbs, the terms *optional* and *required* and the cardinality indicator are as defined in C-CDA Section 1.8 – Conformance Conventions.

6.3.1.4.2 Narrative requirements

600 There is no general requirement for the section text narrative to completely contain the full coded content of all the elements of the section and its contained entries. However for this profile, it is recommended that all coded content in the section and its contained entries SHOULD be included in the narrative for each section. In any case, there SHALL be no conflicts between the narrative and the coded content.

605 In the case where the section ACT relationship is specified to be “DRIV” (derived), then the section narrative SHALL be based solely on the coded content. This narrative content SHOULD include as much of the coded content as possible.

The coded content may not be an equivalent of the narrative.

6.3.1.5 Standards

The following table identifies the standards upon which this specification is based.

Table 6.3.1.4-1: Reference Standards

Standard Name (short)	Standard Name (full)	Reference to Published Standard
CathPCI Registry	NCDR CathPCI Registry v4.4 Coder’s Data Dictionary	http://www.ncdr.com/WebNCDR/NCDRDocuments/CathPCI_v4_CodersDictionary_4.4.pdf
CDAR2	HL7 CDA Release 2.0	http://www.hl7.org/documentcenter/private/standards/cda/r2/cda_r2_normativewebedition2010.zip

Standard Name (short)	Standard Name (full)	Reference to Published Standard
C-CDA	HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation, Release 1 - US Realm	http://www.hl7.org/documentcenter/public/standards/dstu/CDAR2_IG_IHE_CONSOL_R1_DSTU_2011DEC.zip
DICOM	NEMA PS3.16 – DICOM Part 16: Content Mapping Resource	ftp://medical.nema.org/medical/dicom/2009/09_16pu.pdf

610

6.3.2 Cath Report Content Header Element Constraints

The header for the Cardiac Report Content document shall support the following header constraints as noted in this section. Note that this content profile is realm agnostic. These header constraints are based on the C-CDA header constraints but all references to US Realm specific types have been removed.

615

1. SHALL contain exactly one [1..1] **typeId** (CONF:5361).
 - a. This typeId SHALL contain exactly one [1..1] **@root**="2.16.840.1.113883.1.3" (CONF:5250).
 - b. This typeId SHALL contain exactly one [1..1] **@extension**="POCD_HD000040" (CONF:5251).
2. SHALL contain exactly one [1..1] **templateId** (CONF:5252) such that it
 - a. SHALL contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.1.2" for the Cath Report Content document template (CONF:CRC-xxx).
3. SHALL contain exactly one [1..1] **id** (CONF:5363).
 - a. This id SHALL be a globally unique identifier for the document (CONF:9991).
4. SHALL contain exactly one [1..1] **code** (CONF:5253).
 - a. SHALL be selected from ValueSet `ProcedureNoteDocumentTypeCodes 2.16.840.1.113883.11.20.6.1 DYNAMIC` (CONF:8497). Either of the following codes should be included:

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Value Set: ProcedureNoteDocumentTypeCodes 2.16.840.1.113883.11.20.6.1 DYNAMIC			
Code System: LOINC 2.16.840.1.113883.6.1			
LOINC Code	Type of Service 'Component'	Setting 'System'	Specialty/Training/Professional Level 'Method_Type'
18745-0	Study report	Heart	Cardiac catheterization
34896-1	Interventional procedure note	{Setting}	Cardiology

- 635 5. SHALL contain exactly one [1..1] **title** (CONF:5254).
- a. Can either be a locally defined name or the display name corresponding to clinicalDocument/code (CONF:5255).
- 640 6. SHALL contain exactly one [1..1] **effectiveTime** (CONF:5256).
- a. Signifies the document creation time, when the document first came into being. Where the CDA document is a transform from an original document in some other format, the ClinicalDocument.effectiveTime is the time the original document was created. The time when the transform occurred is not currently represented in CDA (CONF:9995).
- 645 7. SHALL contain exactly one [1..1] **confidentialityCode**, which SHOULD be selected from ValueSet HL7 BasicConfidentialityKind 2.16.840.1.113883.1.11.16926 STATIC 2010-04-21 (CONF:5259).

```

<typeId root="2.16.840.1.113883.1.3" extension="POCD_HD000040"/>
<!--CRC Template -->
<templateId root="1.3.6.1.4.1.19376.1.4.1.1.2"/>
650 <id extension="999021" root="2.16.840.1.113883.19"/>
<code codeSystem="2.16.840.1.113883.6.1"
      codeSystemName="LOINC" code="18745-0"
      displayName="Cardiac catheterization study report"/>
655 <title>Cardiac catheterization study report</title>
<effectiveTime value="20050329171504+0500"/>
<confidentialityCode code="N" codeSystem="2.16.840.1.113883.5.25"/>

```

Figure 6.3.2-1: header example

- 660 8. SHALL contain exactly one [1..1] **recordTarget** (CONF:5266-CRC). The **recordTarget** records the patient whose health information is described by the clinical document; the clinical document must contain exactly one **patientRole** element.
- a. Such recordTargets SHALL contain exactly one [1..1] **patientRole** (CONF:5267).
- 665 i. This patientRole SHALL contain at least one [1..*] **id** (CONF:5268)
- ii. This patientRole SHALL contain at least one [1..*] **addr** (CONF:5271).
1. This addr SHALL contain at least one [1..*] **postalCode** (CONF:CRC-xxx).
- 670 iii. This patientRole SHALL contain at least one [1..*] **telecom** (CONF:5280).
- iv. This patientRole SHALL contain exactly one [1..1] **patient** (CONF:5283).
1. This patient SHALL contain exactly one [1..1] **name** (CONF:5284).
- 675 a. This name SHALL contain exactly one [1..1] **family** (CONF:7159).

- b. This name **SHALL** contain at least one [1..*] **given** (CONF:7157).
 - i. The second occurrence of given (given[2]) if provided, **SHALL** include middle name or middle initial (CONF:7163).
- 2. This patient **SHALL** contain exactly one [1..1] **administrativeGenderCode**, which **SHALL** be selected from ValueSet Administrative Gender (HL7 V3) 2.16.840.1.113883.1.11.1 **DYNAMIC** (CONF:6394).
- 3. This patient **SHALL** contain exactly one [1..1] **birthTime** (CONF:5298).
 - a. **SHALL** be precise to year (CONF:5299).
 - b. **SHOULD** be precise to day (CONF:5300).

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```

<recordTarget>
  <patientRole>
    <id extension="12345" root="2.16.840.1.113883.19"/>
    <addr use="HP">
      <streetAddressLine>17 Daws Rd.</streetAddressLine>
      <city>Blue Bell</city>
      <state>MA</state>
      <postalCode>02368</postalCode>
      <country>US</country>
    </addr>
    <telecom value="tel:(781)555-1212" use="HP"/>
    <patient>
      <name>
        <prefix>Mr.</prefix>
        <given>Adam</given>
        <given>Frankie</given>
        <family>Everyman</family>
      </name>
      <administrativeGenderCode code="M"
        codeSystem="2.16.840.1.113883.5.1" displayName="Male"/>
      <birthTime value="19541125"/>
    </patient>
  </patientRole>
</recordTarget>
    
```

715

Figure 6.3.2-2: recordTarget example

720

- 9. **SHALL** contain at least one [1..*] **author** (CONF:5444-CRC). The **author** element represents the person who created the clinical document. If there are multiple procedures performed, there may be multiple authors for the content of this document.
 - a. Such authors **SHALL** contain exactly one [1..1] **time** (CONF:5445). This is the time the author started to contribute to this document.

b. Such authors SHALL contain exactly one [1..1] **assignedAuthor** (CONF:5448).

- 725 i. This assignedAuthor SHALL contain at least one [1..*] **id** (CONF:5449).
 i. This assignedAuthor SHALL contain at least one [1..*] **addr** (CONF:5452).
 ii. This assignedAuthor SHALL contain at least one [1..*] **telecom** (CONF:5428).
 730 iii. This assignedAuthor SHALL contain exactly one [1..1] **assignedPerson** (CONF:5430-CRC).
 1. This assignedPerson SHALL contain at least one [1..*] **name** (CONF:5431-CRC).

735

```

<author>
  <time value="20120329224411+0500"/>
  <assignedAuthor>
    <id extension="KP00017" root="2.16.840.1.113883.19.5"/>
    <addr>
      <streetAddressLine>21 North Ave.</streetAddressLine>
      <city>Burlington</city>
      <state>MA</state>
      <postalCode>02368</postalCode>
      <country>US</country>
    </addr>
    <telecom use="WP" value="tel:(555)555-1003"/>
    <assignedPerson>
      <name>
        <given>Henry</given>
        <family>Seven</family>
      </name>
    </assignedPerson>
  </assignedAuthor>
</author>
    
```

740

745

750

Figure 6.3.2-3: Person author example

755

10. SHALL contain exactly one [1..1] **custodian** (CONF:5519).

a. This custodian SHALL contain exactly one [1..1] **assignedCustodian** (CONF:5520).

- 760 i. This assignedCustodian SHALL contain exactly one [1..1] **representedCustodianOrganization** (CONF:5521).
 1. This representedCustodianOrganization SHALL contain at least one [1..*] **id** (CONF:5522).
 2. This representedCustodianOrganization SHALL contain exactly one [1..1] **name** (CONF:5524).
 765 3. This representedCustodianOrganization SHALL contain exactly one [1..1] **telecom** (CONF:5525).

4. This `representedCustodianOrganization` SHALL contain at least one `[1..*]` `addr` (CONF:5559).

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```

<custodian>
  <assignedCustodian>
    <representedCustodianOrganization>
      <id root="2.16.840.1.113883.19.5"/>
      <name>Good Health Clinic</name>
      <telecom value="tel:(555)555-1212" use="WP"/>
      <addr use="WP">
        <streetAddressLine>17 Daws Rd.</streetAddressLine>
        <city>Blue Bell</city>
        <state>MA</state>
        <postalCode>02368</postalCode>
        <country>US</country>
      </addr>
    </representedCustodianOrganization>
  </assignedCustodian>
</custodian>

```

Figure 6.3.2-4: custodian example

790

11. SHALL contain exactly one `[1..1]` `legalAuthenticator` (CONF:5579-CRC).

- a. The `legalAuthenticator` SHALL contain exactly one `[1..1]` `time` (CONF:5580).
- b. This `legalAuthenticator`, if present, SHALL contain exactly one `[1..1]` `signatureCode` (CONF:5583).

795

- i. This `signatureCode` SHALL contain exactly one `[1..1]` `@code="S"` (CodeSystem: Participationsignature 2.16.840.1.113883.5.89) (CONF:5584).

- c. This `legalAuthenticator`, if present, SHALL contain exactly one `[1..1]` `assignedEntity` (CONF:5585).

800

- i. This `assignedEntity` SHALL contain at least one `[1..*]` `id` (CONF:5586).
- ii. This `assignedEntity` MAY contain zero or one `[0..1]` `code` (CONF:9949-CRC).
- ii. This `assignedEntity` SHALL contain at least one `[1..*]` `addr` (CONF:5589).

805

- iii. This `assignedEntity` SHALL contain at least one `[1..*]` `telecom` (CONF:5595).

1. Such telecoms SHOULD contain `@use` (CONF:7999-CRC).

- iv. This `assignedEntity` SHALL contain exactly one `[1..1]` `assignedPerson` (CONF:5597).

810

1. This `assignedPerson` SHALL contain at least one `[1..*]` `name` (CONF:5598).

The **legalAuthenticator** identifies the single person legally responsible for the document and must be present if the document has been legally authenticated.

```

815 <legalAuthenticator>
      <time value="20050329224411+0500"/>
      <signatureCode code="S"/>
      <assignedEntity>
820         <id extension="KP00017" root="2.16.840.1.113883.19"/>
         <addr>
           <streetAddressLine>21 North Ave.</streetAddressLine>
           <city>Burlington</city>
           <state>MA</state>
           <postalCode>02368</postalCode>
825           <country>US</country>
         </addr>
         <telecom use="WP" value="tel:(555)555-1003"/>
         <assignedPerson>
           <name>
830             <given>Henry</given>
             <family>Seven</family>
           </name>
         </assignedPerson>
       </assignedEntity>
     </legalAuthenticator>

```

835 **Figure 6.3.2-5: legalAuthenticator example**

12. MAY contain zero or more [0..*] **authenticator** (CONF:5607).
- a. Such authenticators, if present, SHALL contain exactly one [1..1] **time** (CONF:5608).
 - 840 b. Such authenticators, if present, SHALL contain exactly one [1..1] **signatureCode** (CONF:5610).
 - i. This signatureCode SHALL contain exactly one [1..1] @code="S" (CodeSystem: Participationsignature 2.16.840.1.113883.5.89) (CONF:5611).
 - 845 c. Such authenticators, if present, SHALL contain exactly one [1..1] **assignedEntity** (CONF:5612).
 - i. This assignedEntity SHALL contain at least one [1..*] **id** (CONF:5613).
 - iii. This assignedEntity SHALL contain at least one [1..*] **addr** (CONF:5616).
 - 850 ii. This assignedEntity SHALL contain at least one [1..*] **telecom** (CONF:5622).
 - iii. This assignedEntity SHALL contain exactly one [1..1] **assignedPerson** (CONF:5624).
 - 855 1. This assignedPerson SHALL contain at least one [1..*] **name** (CONF:5625).

The authenticator identifies a participant or participants who attested to the accuracy of the information in the document. There may be one authenticator for the content for each of the Cath procedures – e.g., diagnostic cath and PCI.

860

865

870

875

880

```

<authenticator>
  <time value="20050329224411+0500"/>
  <signatureCode code="S"/>
  <assignedEntity>
    <id extension="KP00017" root="2.16.840.1.113883.19"/>
    <addr>
      <streetAddressLine>21 North Ave.</streetAddressLine>
      <city>Burlington</city>
      <state>MA</state>
      <postalCode>02368</postalCode>
      <country>US</country>
    </addr>
    <telecom use="WP" value="tel:(555)555-1003"/>
    <assignedPerson>
      <name>
        <given>Henry</given>
        <family>Seven</family>
      </name>
    </assignedPerson>
  </assignedEntity>
</authenticator>

```

Figure 6.3.2-6: Authenticator example

13. MAY contain zero or one [0..1] **inFulfillmentOf** (CONF:9952-CRC).

885

a. Such **inFulfillmentOf** elements, if present, SHALL contain exactly one [1..1] **order** (CONF:9953-CRC).

890

- i. This order SHALL contain at least one [1..*] **id** and one value SHALL be the Accession Number used in the DICOM imaging data, with the root representing the Assigning Authority (Issuer of Accession Number) (CONF:9954-CRC).
- ii. This order SHALL contain at least one [1..*] **priorityCode** with values from coding system HL7 ActPriority (OID = 2.16.840.1.113883.5.7) (CONF:CRC-xxx).

```

895 <inFulfillmentOf>
      <order>
        <id root="1.2.3.4.5.6" extension="acc#1" />
        <priorityCode code="R" codeSystem=" 2.16.840.1.113883.5.7"
900       codeSystemName="ActPriority" displayName="Routine">
      </order>
    </inFulfillmentOf>

```

Figure 6.3.2-7: inFulfillmentOf example

14. **MAY** contain zero or one [0..1] **authorization** (CONF:CRC-xxx).
- 905 a. Such authorization elements, if present, **MAY** contain zero or more [0..*] **consent**. (CONF:CRC-xxx).
- b. A consent **MAY** be provided for the procedure and a consent **MAY** be provided for the anesthesia. (CONF:CRC-xxx)
- 910 i. The following LOINC codes **SHOULD** be used
1. 64293-4 – procedure consent
 2. 61359-6 – anesthesia consent

```

915 <authorization typeCode="AUTH">
      <consent classCode="CONS" moodCode="EVN">
        <id root="629deb70-5306-11df-9879-0800200c9a66" />
        <code codeSystem=" 2.16.840.1.113883.6.1" codeSystemName="LOINC"
920       code="64293-4" displayName="Procedure Consent"/>
        <statusCode code="completed"/>
      </consent>
      <consent classCode="CONS" moodCode="EVN">
        <id root="629deb70-5306-11df-9879-0800200c9a66" />
        <code codeSystem=" 2.16.840.1.113883.6.1" codeSystemName="LOINC"
925       code="61359-6" displayName="Anesthesia Consent"/>
        <statusCode code="completed"/>
      </consent>
    </authorization>

```

Figure 6.3.2-8: consent example

15. **SHALL** contain exactly one [1..1] **componentOf** (CONF:9955-CRC)
- 930 a. This componentOf element **SHALL** contain exactly one [1..1] **encompassingEncounter** (CONF:9956)
- i. **SHALL** contain at least one [1..*] **id** (CONF:9959).
 - ii. **SHALL** contain exactly one [1..1] **effectiveTime** (CONF:9958).
 1. **SHALL** be accurate to the day and **MAY** be accurate to the
 935 second (CONF:CRC-xxx).
 - iii. This componentOf/encompassingEncounter **SHALL** contain exactly one [1..1] **code** (CONF:8501).

- 940
- iv. This componentOf/encompassingEncounter SHALL contain at least one [1..*] **location/healthCareFacility** (CONF:8500-CRC).
1. This **SHALL** contain at least one [1..*] **code** representing the type of location (CONF:8500-CRC).
 2. This **SHALL** contain at least one [1..*] **id** (CONF:8500-CRC).
 3. This **SHOULD** contain at least one [1..*] **serviceProviderOrganization** (CONF:8500-CRC-xxx).
 - 945 a. This **SHALL** contain at least one [1..*]**name** (CONF:8500-CRC-xxx).
 - b. This **SHALL** contain at least one [1..*]**addr** (CONF:8500-CRC-xxx).
 - 950 c. This assignedEntity **SHALL** contain at least one [1..*] **telecom** (CONF:8500-CRC-xxx).
 4. This **MAY** contain zero or more [0..*] **location** (CONF:8500-CRC-xxx).
 - 955 a. If present, this **SHALL** contain at least one [1..*]**name** and/or **addr** to identify the place of the encounter (CONF:8500-CRC-xxx).
- v. This componentOf/encompassingEncounter MAY contain zero to four [0..4] **encounterParticipant** (CONF:8502-CRC) such that it
- 960 1. **MAY** contain at most two [0..2] **@typeCode="REF"** Referrer (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90) for the referring cardiologist and referring physician (CONF:8503-CRC).
 2. **MAY** contain zero or one [0..1] **@typeCode="ATND"** Physician of Record (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90) (CONF:8503-CRC).
 - 965 3. **MAY** contain zero or one [0..1] **@typeCode="RESP"** Responsible Party (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90) (CONF:8503-CRC).
- 970

```

975 <componentOf>
    <encompassingEncounter>
      <id extension="KP00017" root="2.16.840.1.113883.19"/>
      <effectiveTime value="20110407"/>
980      <code code="1234097013" codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT"
        displayName="Diagnostic Coronary Angiography ">
      <location>
        <healthCareFacility>
985          <id root="1.2.3.4.5.6.7" extension="facility ID"/>
          <code code="CARD" codeSystem="2.16.840.1.113883.5.111"
            codeSystemName="roleCode" displayName="Cardiology Clinic">
          <serviceProviderOrganization>
            <name>My Favorite Cardiac Care Organization</name>
990            <addr>
              <streetAddressLine>Healthcare Lane</streetAddressLine>
              <city>East Town</city>
              <state>OH</state>
              <country>US</country>
995            </addr>
            <telecom value="1-800-555-1212" use="WP"/>
          </serviceProviderOrganization>
          <location>
            <name>My Cardiac Hospital</name>
1000            <addr>
              <streetAddressLine>Healthcare Lane</streetAddressLine>
              <city>East Town</city>
              <state>OH</state>
              <country>US</country>
1005            </addr>
          </location>
        </healthCareFacility>
      </location>
      <encounterParticipant typeCode="REF">
1010        <assignedEntity>
          <id root="2.16.840.1.113883.4.6" extension="12345"/>
          <code code="xyz" codeSystem="2.16.840.1.113883.6.101"
            codeSystemName="nuccProviderCodes"
            displayName="Referring cardiologist"/>
          <addr>Referring Physician Lane, USA</addr>
1015          <telecom value="1-800-555-1212" use="WP"/>
          <assignedPerson>
            <name>Dr. Referring Physician</name>
          </assignedPerson>
        </assignedEntity>
1020      </encounterParticipant>
    </encompassingEncounter >
  </componentOf>

```

Figure 6.3.2-9: componentOf/encompassingEncounter example

1025 16. SHALL contain exactly one [1..1] **documentationOf** (CONF:8510-CRC).

- 1030 a. Such documentationOf SHALL contain exactly one [1..1] **serviceEvent** (CONF:10061).
- 1035 i. The value of Clinical Document /documentationOf/serviceEvent/code **SHOULD** be selected from code system 2.16.840.1.113883.6.96 SNOMED CT and **MAY** be selected from a localized procedure coding system for a given country such as 2.16.840.1.113883.6.104 ICD9 CM Procedures or 2.16.840.1.113883.6.12 CPT-4 in the U.S (CONF:CRC-xxx).
- 1040 ii. This serviceEvent SHALL contain at least one [1..*] **id** values including the Study Instance UID used in the DICOM imaging data, with the UID value in the root attribute (CONF:CRC-xxx).
- 1045 iii. This serviceEvent SHALL contain exactly one [1..1] **effectiveTime** (CONF:10062).
- 1050 1. The serviceEvent/effectiveTime SHALL be present with effectiveTime/low (CONF:8513).
- 1055 2. If a width is not present, the serviceEvent/effectiveTime SHALL include effectiveTime/high. (CONF:8514)
- 1060 3. When only the date and the length of the procedure are known a width element SHALL be present and the serviceEvent/effectiveTime/high SHALL not be present. (CONF:8515).
- 1065 4. The serviceEvent/effectiveTime SHALL be accurate to the day and **MAY** be accurate to the second (CONF:CRC-xxx).
- 1070 iv. This serviceEvent SHALL contain at least one [1..*] **performer** (CONF:8520-CRC) such that it
- 1075 1. SHALL contain one or two [1..2] **@typeCode="PPRF"** Primary Performer (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90). This is for the case where both a cath and PCI are performed in the same procedure. (CONF:8521-CRC).
- 1080 2. **SHALL** contain exactly one [1..1] **assignedEntity** (CONF:14911).
- 1085 a. This assignedEntity **SHOULD** contain zero or one [0..1] **code** (CONF:14912).
- 1090 i. The code, if present, **SHOULD** contain zero or one [0..1] **@code**. (CONF:14913-CRC).
- 1095 3. Any assistants **SHALL** be identified and **SHALL** be identified as secondary performers (SPRF). (CONF:8524).

1065

```

1070 <documentationOf>
      <serviceEvent classCode="PROC">
        <code code="70051019" codeSystem="2.16.840.1.113883.6.96"
          codeSystemName="SNOMED CT"
          displayName="Diagnostic catheterization"/>
        <id root="DICOM study instance UID" extension="acc1"/>
1075 <effectiveTime>
        <low value="201003292240" />
        <width value="15" unit="m"/>
      </effectiveTime>
      <performer typeCode="PPRF">
1080 <assignedEntity>
        <id extension="IO00017" root="2.16.840.1.113883.19.5" />
        <code code="17561000"
          codeSystem="2.16.840.1.113883.6.96"
          codeSystemName="SNOMED CT"
          displayName="Cardiologist" />
1085 <addr>
        <streetAddressLine>1001 Hospital Lane</streetAddressLine>
        <city>Ann Arbor</city>
        <state>MI</state>
        <postalCode>99999</postalCode>
1090 <country>US</country>
      </addr>
      <telecom value="tel:(999)555-1212" />
      <assignedPerson>
        <name>
1095 <prefix>Dr.</prefix>
        <given>Tony</given>
        <family>Tum</family>
      </name>
      </assignedPerson>
1100 </assignedEntity>
    </performer>
  </serviceEvent>
</documentationOf>

```

Figure 6.3.2-10: documentationOf/serviceEvent example

1105 6.3.3 Cath Report Content Body Containment

The body for the Cardiac Report Content document shall include section content modules. The section content modules will be specified by a set of constraints.

1. SHALL contain exactly one [1..1] **component** (CONF:9588).
 - 1110 a. A Cath Report Content SHALL have a structuredBody (CONF:9589-CRC).
 - 1115 i. A Cath Report Content SHALL conform to CDA Level 3 (structuredBody containing sections that contain a narrative block and coded entries). In this template (templateId 1.3.6.1.4.1.19376.1.4.1.1.2), some coded entries are used. (CONF:9590-CRC).

- b. The component/structuredBody **SHALL** conform to the section constraints below (CONF:9595-CRC).
 - i. **EACH section SHALL HAVE A title AND THE title SHALL NOT BE EMPTY** (CONF:9937).

1120

Table 6.3.3-1 identifies the set of specific *section content modules* that may be required, recommended, or allowed to be included in the CRC document. This table also identifies the most important *entry content modules* contained within those section content modules. The containment relationship among the section and entry content modules in the body of a Cath Report Content document is represented in the “Template Title” column as noted by the indentation relative to the other content modules.

1125

1. Section content modules

- a. Any section content module that is used exactly as specified in C-CDA shall not have the C-CDA constraints replicated in this specification.
- b. If the section content module is used in this profile but with different vocabulary constraints, then the vocabulary constraints shall be listed in the “Constraints” columns of the table and shall be included in this specification.
- c. Sample XML shall be included for each section content module and should include XML for each entry contained within the section.

1130

1135

2. Entry content modules

- a. Any entry content module that is used exactly as specified in C-CDA shall not have the specific constraints replicated in this specification.
- b. If the entry content module is relevant to this CRC profile, it shall be included in Table 6.3.3-1 following the section content module it is contained within.
- c. If the entry content module has CRC specific vocabulary constraints, the constraints shall be identified in the “Constraints” columns of the table and shall be documented in this specification.
- d. Sample XML should be included for the entries within the section content module where it is used.

1140

1145

Table 6.3.3-1: Template Containment for a Cath Report Content document

Template Title	Cardinality	Template Type	templateId	Specification Document	Constraints
Cath Report Content	R[1..1]	document	1.3.6.1.4.1.19376.1.4.1.1.2	CARD TF-3 6.3	
Document Summary Section	O[0..1]	section	1.3.6.1.4.1.19376.1.4.1.2.16	CARD TF-3 6.3.4.1	CARD TF-3 6.3.4.1

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Template Title	Cardinality	Template Type	templateId	Specification Document	Constraints
Medical History - Cardiac Section	R[1..1]	section	1.3.6.1.4.1.19376.1.4.1.2.17	CARD TF-3 6.3.4.2 (C-CDA 4.31 - parent)	CARD TF-3 6.3.4.2
Procedure Activity Observation	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.13	C-CDA 5.48	CARD TF-3 6.3.4.2.2
Procedure Activity Procedure	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.14	C-CDA 5.49	CARD TF-3 6.3.4.2.3
Problem Observation - Cardiac	O[0..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.9	CARD TF-3 6.3.5.1 (C-CDA 5.45 – parent)	CARD TF-3 6.3.4.2.1
Age Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.31	C-CDA 5.3	
Health Status Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.5	C-CDA 5.19	
Problem Status	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.6	C-CDA 5.46	
Severity Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.8	C-CDA 5.60	
Allergies Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.6	C-CDA 4.2	CARD TF-3 6.3.4.3
Allergy Problem Act	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.30	C-CDA 5.5	
Allergy Observation	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.7	C-CDA 5.4	CARD TF-3 6.3.4.3.1
Allergy Status Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.28	C-CDA 5.6	
Reaction Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.9	C-CDA 5.54	
Severity Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.8	C-CDA 5.60	
Family History – Cardiac Section	O[0..1]	section	1.3.6.1.4.1.19376.1.4.1.2.18	CARD TF-3 6.3.4.4	CARD TF-3 6.3.4.4
Problem Observation	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.4	C-CDA 5.45	CARD TF-3 6.3.4.4.1
Social History Section	O[0..1]	section	2.16.840.1.113883.10.20.22.2.17	C-CDA 4.57	CARD TF-3 6.3.4.5
Social History Observation	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.38	C-CDA 5.61	CARD TF-3 6.3.4.5.1
Physical Exam Section	R[1..1]	section	2.16.840.1.113883.10.20.2.10	C-CDA 4.38	CARD TF-3 6.3.4.6
Vital Signs Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.4	C-CDA 4.60	CARD TF-3 6.3.4.7
Vital Signs Organizer	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.26	C-CDA 5.66	

IHE Cardiology Technical Framework Supplement – Cath Report Content (CRC)

Template Title	Cardinality	Template Type	templateId	Specification Document	Constraints
Vital Sign Observation	R[2..*]	entry	2.16.840.1.113883.10.20.22.4.27	C-CDA 5.65	CARD TF-3 6.3.4.7.1
Pre-Procedure Results – Cardiac Section	R[1..1]	section	1.3.6.1.4.1.19376.1.4.1.2.23	CARD TF-3 6.3.4.8 (C-CDA 4.55 – parent)	CARD TF-3 6.3.4.8
Result Organizer - Cardiac	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.11	CARD TF-3 6.3.4.8.1 (C-CDA 5.57 – parent)	CARD TF-3 6.3.4.8.1
Result Observation	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.2	C-CDA 5.56	CARD TF-3 6.3.4.8.2
Planned Procedure Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.30	C-CDA 4.40	CARD TF-3 6.3.4.9
Plan of Care Activity Procedure	R[1..2]	entry	2.16.840.1.113883.10.20.22.4.41	C-CDA 5.36	
Procedure Indications Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.29	C-CDA 4.50	CARD TF-3 6.3.4.10
Indication	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.19	C-CDA 5.25	CARD TF-3 6.3.4.10.1
Anesthesia Section	O[0..1]	section	2.16.840.1.113883.10.20.22.2.25	C-CDA 4.3	CARD TF-3 6.3.4.11
Procedure Activity Procedure	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.14	C-CDA 5.49	CARD TF-3 6.3.4.11.1
Medication Activity	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.16	C-CDA 5.27	

IHE Cardiology Technical Framework Supplement – Cath Report Content (CRC)

Template Title	Cardinality	Template Type	templateId	Specification Document	Constraints
Medications Administered Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.38	C-CDA 4.32	CARD TF-3 6.3.4.12
Medication Activity	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.16	C-CDA 5.27	
Medication Information	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.23	C-CDA 5.29	CARD TF-3 6.3.4.12.1
Procedure Description - Cardiac Section	R[1..1]	section	1.3.6.1.4.1.19376.1.4.1.2.19	CARD TF-3 6.3.4.13 (C-CDA 4.45 – parent)	CARD TF-3 6.3.4.13
Lesion Observation	O[0..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.10	CARD TF-3 6.3.5.2	
Procedure Device Organizer - Cardiac	O[0..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.12	CARD TF-3 6.3.4.13.2	CARD TF-3 6.3.4.13.2
Device Observation	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.13	CARD TF-3 6.3.4.13.3	CARD TF-3 6.3.4.13.3
Procedure Activity Procedure - Cardiac	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.14	CARD TF-3 6.3.4.13.1 (C-CDA 5.49 – parent)	CARD TF-3 6.3.4.13.1
Lesion Observation	O[0..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.10	CARD TF-3 6.3.5.2	
Product Instance	O[1..*]	entry	2.16.840.1.113883.10.20.22.4.37	C-CDA 5.51	
Procedure Device Organizer - Cardiac	O[0..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.12	CARD TF-3 6.3.4.13.2	CARD TF-3 6.3.4.13.2
Device Observation	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.13	CARD TF-3 6.3.4.13.3	CARD TF-3 6.3.4.13.3
Procedure Specimens Taken Section	O[0..1]	section	2.16.840.1.113883.10.20.22.2.31	C-CDA 4.51	CARD TF-3 6.3.4.14
Procedure Disposition Section	R[1..1]	section	2.16.840.1.113883.10.20.18.2.12	C-CDA 4.46	CARD TF-3 6.3.4.15
Procedure Results - Cardiac Section	R[1..1]	section	1.3.6.1.4.1.19376.1.4.1.2.20	CARD TF-3 6.3.4.16 (C-CDA 4.48 – parent)	CARD TF-3 6.3.4.16
Procedure Results Organizer - Cardiac	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.15	CARD TF-3 6.3.4.16.1 (C-CDA 5.57 – parent)	
Result Observation - Cardiac	R[1..*]	entry	1.3.6.1.4.1.19376.1.4.1.4.16	CARD TF-3 6.3.4.16.2 (C-CDA 5.56 – parent)	

Template Title	Cardinality	Template Type	templateId	Specification Document	Constraints
Severity Observation	O[0..1]	entry	2.16.840.1.113883.10.20.22.4.8	C-CDA 5.60	
Complications Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.37	C-CDA 4.8	CARD TF-3 6.3.4.17
Problem Observation	O[0..*]	entry	2.16.840.1.113883.10.20.22.4.4	C-CDA 5.45	CARD TF-3 6.3.4.17.1
Postprocedure Diagnosis Section	R[1..1]	section	2.16.840.1.113883.10.20.22.2.36	C-CDA 4.42	CARD TF-3 6.3.4.18
Postprocedure Diagnosis	R[1..1]	entry	2.16.840.1.113883.10.20.22.4.51	C-CDA 5.40	
Problem Observation	R[1..*]	entry	2.16.840.1.113883.10.20.22.4.4	C-CDA 5.45	CARD TF-3 6.3.4.18.1
Plan of Care – Cardiac Section	O[0..1]	section	1.3.6.1.4.1.19376.1.4.1.2.22	CARD TF-3 6.3.4.19 (C-CDA 4.39 - parent)	CARD TF-3 6.3.4.19
Plan of Care Activity Act - Cardiac	R[1..1]	entry	1.3.6.1.4.1.19376.1.4.1.4.17	CARD TF-3 6.3.4.19.1 (C-CDA 5.33 – parent)	CARD TF-3 6.3.4.19.1
Key Images – Cardiac Section	O[0..1]	section	1.3.6.1.4.1.19376.1.4.1.2.21	CARD TF-3 6.3.4.20	CARD TF-3 6.3.4.20
Sop Instance Observation	R[1..*]	entry	2.16.840.1.113883.10.20.6.2.8	C-CDA 5.62	
DICOM Object Catalog Section	O[0..1]	section	2.16.840.1.113883.10.20.6.1.1	C-CDA 4.9	
Study Act	R[1..*]	entry	2.16.840.1.113883.10.20.6.2.6	C-CDA 5.63	
Series Act	R[1..*]	entry	2.16.840.1.113883.10.20.6.4.63	C-CDA 5.58	
Sop Instance Observation	R[1..*]	entry	2.16.840.1.113883.10.20.6.2.8	C-CDA 5.62	

1150 6.3.4 Cath Report Content Document Section/Entry Constraints

6.3.4.1 Document Summary Section 55112-7

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.16 (open)]

The Document Summary section content module includes a summary of most significant aspects of the procedures in a narrative form. It is a condensed form of the full narrative report whose structure has no constraint.

1155

This Document Summary section content module is a new content module that has no equivalent in C-CDA. The complete set of constraints for the Document Summary section content module are listed below.

- 1160 1. **SHALL** contain exactly one [1..1] **templateId** (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] **@root="1.3.6.1.4.1.19376.1.4.1.2.16"** (CONF:CRC-xxx).
2. **SHALL** contain exactly one [1..1] **code/@code="55112-7"** Document Summary (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:CRC-xxx).
- 1165 3. **SHALL** contain exactly one [1..1] **title** (CONF:CRC-xxx).
4. **SHALL** contain exactly one [1..1] **text** (CONF:CRC-xxx).
- a. The text element **MAY** contain one or more **renderMultiMedia** element representing an in-line graphic. The related observationMedia entry may be within the summary section structured entries or may be referenced from another section.
- 1170 5. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] **ObservationMedia** element (CONF:CRC-xxx) such that it
- i. **SHALL** contain exactly one [1..1] **@classCode="OBS"** (CONF:CRC-xxx).
- ii. **SHALL** contain exactly one [1..1] **@moodCode="EVN"** Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:CRC-xxx).
- iii. **SHALL** contain at least one [1..*] **id** (CONF:CRC-xxx).
- iv. **SHALL** contain at least one [1..*] **value** with **@xsi:type="ED"** (CONF:CRC-xxx)
1. **SHALL** contain exactly one [1..1] **@mediaType** drawn from the value set 1.3.6.1.4.1.19376.1.4.1.5.45 SupportedFileFormats **STATIC** 20100512 (CONF:CRC-xxx).
2. **MAY** contain zero or one [0..1] **reference** (CONF:CRC-xxx).
- a. The URL of a referenced graphic element **MAY** be present (CONF:CRC-xxx).
3. An encapsulated data value may have both inline data and a reference. The reference must point to the same data as provided inline as per HL7 v3 Data Types – Abstract Specification, Release 1, Section 2.4.5 (CONF:CRC-xxx).
- 1185
- 1190
- 1195

```

1200 <!-- example with external content referenced by file name -->
1205 <section>
1210   <templateId root="1.3.6.1.4.1.19376.1.4.1.2.16"/>
1215   <code codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"
1220     code="55112-7" displayName="DOCUMENT SUMMARY"/>
1225   <title>CATH PROCEDURE SUMMARY</title>
1230   <text>
1235     <paragraph>A cath procedure was performed. The following image shows the
1240     region of interest.</paragraph>
1245     <renderMultiMedia referencedObject="CRC-image1"/>
1250     <paragraph>The patient needed no further interventions.</paragraph>
1255   </text>
1260   <entry>
1265     <observationMedia classCode="OBS" moodCode="EVN" ID="CRC-image1">
1270       <id root="2.16.840.1.113883.19.2.1"/>
1275       <value xsi:type="ED" mediaType="image/jpeg">
1280         <reference value="sample cath image.jpeg"/>
1285       </value>
1290     </observationMedia>
1295   </entry>
1300 </section>

1305 <!-- alternative example - embed the content within the reference element value
1310 attribute -->
1315 <section>
1320   ...
1325   <text mediaType="image/jpeg" representation="B64">elxydGY...</text>
1330   <entry>
1335     <observationMedia classCode="OBS" moodCode="EVN" ID="CRC-embedded">
1340       <id root="2.16.840.1.113883.19.2.1"/>
1345       <value xsi:type="ED" mediaType="image/jpeg" reference="B64">
1350         <reference value="elxydGY...">
1355       </value>
1360     </observationMedia>
1365   </entry>
1370 </section>

```

Figure 6.3.4.1-1: Document Summary – Cardiac section example

1235 6.3.4.2 Medical History - Cardiac Section 11329-0

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.17 (open)]

([section: templateId 2.16.840.1.113883.10.20.22.2.39 (open)] - parent)

1240 The Medical History – Cardiac section content module describes all aspects of the medical history of the patient even if not pertinent to the current procedure, and may include chief complaint, past medical history, social history, family history, surgical or procedure history, medication history, and other history information. The history may be limited to information pertinent to the current procedure or may be more comprehensive. The history may be reported as a collection of random clinical statements or it may be reported categorically. Entries for

1245 History of Past Illness and History of Present Illness have been consolidated into this section. Social and Family History are discussed in their own sections. For this Cath Report Content profile, this Medical History – Cardiac section content module may also contain history about specific relevant problems as problem observations.

1250 In the event that the patient was transferred from another facility where there was a problem indication that the patient was determined to need a cath procedure, this will be noted as a problem observation in this medical history section as text in the narrative for now until there is a code representing this.

This Medical History – Cardiac section content module extends the Medical (General History Section (C-CDA 4.31). The additional constraints are listed below.

- 1255 1. **SHALL** contain exactly one [1..1] **templateId** (CONF:8160) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.17" (CONF:10403-CRC).
2. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] [Problem Observation - Cardiac](#) (1.3.6.1.4.1.19376.1.4.1.4.9) (CONF:CRC-xxx).
- 1260 3. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] [Procedure Activity Observation](#) (2.16.840.1.113883.10.20.22.4.13) (CONF:CRC-xxx).
- 1265 4. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] [Procedure Activity Procedure](#) (2.16.840.1.113883.10.20.22.4.14) (CONF:CRC-xxx).

```

1270 <section>
      <templateId root="1.3.6.1.4.1.19376.1.4.1.2.17"/>
      <templateId root="2.16.840.1.113883.10.20.22.2.39"/>
      <code code="11329-0" codeSystem="2.16.840.1.113883.6.1"
1275         codeSystemName="LOINC"
            displayName="MEDICAL (GENERAL) HISTORY"/>
      <title>MEDICAL (GENERAL) HISTORY</title>
      <text>
        <list listType="ordered">
1280           <item>Patient has had a recent issue with chest pain that does
              not seem to be related to any particular cause.</item>
           <item>Previous concerns of heart disease were actually
related to other causes.</item>
           <item>Patient had recent weight gain due to sedentary lifestyle and
1285             new job.</item>
        </list>
      </text>
      <entry>
        <observation classCode="OBS" moodCode="EVN">
1290           <templateId root="1.3.6.1.4.1.19376.1.4.1.4.9"/>
           <!-- Problem Observation - Cardiac template -->
           <id root="d11275e7-67ae-11db-bd13-0800200c9a66"/>
           <code code="55607006" codeSystem="2.16.840.1.113883.6.96"
              codeSystemName="SNOMED CT"
1295             displayName="Problem"/>
           <text>There was history of hypertension.</text>
           <statusCode code="completed"/>
           <effectiveTime>
</effectiveTime>
           <value xsi:type="CD" code="38341003"
1300             codeSystem="1.2.840.10008.6.1.253" codeSystemName="SNOMED CT"
              displayName="Hypertension"/>
           <entryRelationship typeCode="SUBJ" inversionInd="true">
             <observation classCode="OBS" moodCode="EVN">
1305               <templateId root="2.16.840.1.113883.10.20.22.4.31"/>
               <!-- Age observation template -->
               <code code="445518008" codeSystem="2.16.840.1.113883.6.96"
                  displayName="Age At Onset"/>
               <statusCode code="completed"/>
               <value xsi:type="PQ" value="57" unit="a"/>
1310             </observation>
           </entryRelationship>
           <entryRelationship typeCode="SUBJ" inversionInd="true">
             <observation classCode="OBS" moodCode="EVN">
1315               <templateId root="2.16.840.1.113883.10.20.22.4.6"/>
               <!--Problem status template -->
               <code code="33999-4" codeSystem="2.16.840.1.113883.6.1"
                  displayName="Status"/>
               <statusCode code="completed"/>
               <value xsi:type="CD" code="55561003"
1320             codeSystem="2.16.840.1.113883.6.96"
                  codeSystemName="SNOMED CT" displayName="Active"/>
             </observation>
           </entryRelationship>

```


1325

```

<entryRelationship typeCode="REFR" inversionInd="true">
  <observation classCode="OBS" moodCode="EVN">
    <templateId root="2.16.840.1.113883.10.20.22.4.5"/>
    <!-- Health status observation template -->
    <code code="11323-3"
      codeSystem="2.16.840.1.113883.6.1"
      codeSystemName="LOINC"
      displayName="Health status"/>
    <statusCode code="completed"/>
    <value xsi:type="CE" code="413322009"
      codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT"
      displayName="Resolved"/>
  </observation>
</entryRelationship>

```

1330

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```

<entryRelationship typeCode="REFR" inversionInd="true">
  <observation classCode="OBS" moodCode="EVN">
    <templateId root="2.16.840.1.113883.10.20.22.4.8"/>
    <!-- Severity observation template -->
    <code code="SEV" displayName="Severity Observation"
      codeSystem="2.16.840.1.113883.5.4"
      codeSystemName="ActCode"/>
    <text>
      <reference value="#severity"/>
    </text>
    <statusCode code="completed"/>
    <value xsi:type="CD" code="371924009" displayName="Moderate to severe"
      codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT"/>
  </observation>
</entryRelationship>
</observation>
</entry>
</entry>

```

1360

1365

1370

```

<procedure classCode="PROC" moodCode="EVN">
  <templateId root="2.16.840.1.113883.10.20.22.4.14"/>
  <!-- Procedure Activity Procedure template -->
  <id root="e401f340-7be2-11db-9fe1-0800200c9a66"/>
  <code code="500786010" codeSystem="1.3.6.1.4.1.19376.1.4.1.5.40"
    displayName="Left Heart Cath Procedure">
    <originalText>Left Heart Cath Procedure
      <reference value="procedure1"/></originalText>
    </code>
    <text>
      <reference value="procedure1"/>
    </text>
    <statusCode code="completed"/>
    <effectiveTime value="1998"/>
    <targetSiteCode code="41879009" codeSystem="2.16.840.1.113883.6.96"
      displayName="Distal RCA"/>
  </procedure>

```

1375

`</procedure>``</entry>``</entry>`

1380

`<observation classCode="OBS" moodCode="EVN">``<templateId root="2.16.840.1.113883.10.20.22.4.13"/>``<!-- Procedure Activity Observation template -->``<id extension="proc1" root="2.16.840.1.113883.19"/>``<code code="500786010" codeSystem="2.16.840.1.113883.6.96"``displayName="Left Heart Cath Procedure" codeSystemName="SNOMED CT">``<originalText>``<reference value="#procedure1"/>`

1385

`</originalText>``</code>``<statusCode code="aborted"``codeSystem="2.16.840.1.113883.5.14"``codeSystemName="ActStatus"/>`

1390

`<effectiveTime value="20110203"/>``<priorityCode code="CR" codeSystem="2.16.840.1.113883.5.7"``codeSystemName="ActPriority"``displayName="Callback results"/>`

1395

`<value xsi:type="CD" code="" codeSystem="2.16.840.1.113883.6.96"/>``<methodCode nullFlavor="UNK"/>``<targetSiteCode code="91083009" codeSystem="2.16.840.1.113883.6.96"``codeSystemName="SNOMED CT"``displayName=" Proximal Right Coronary Artery" />`

1400

`<performer>``<assignedEntity>``<id root="1.2.3.4" extension="1234"/>``<addr>``<streetAddressLine>17 Daws Rd.</streetAddressLine>``<city>Blue Bell</city>``<state>MA</state>``<postalCode>02368</postalCode>``<country>US</country>``</addr>``<telecom use="WP" value="1(555)555-1234"/>`

1410

`<representedOrganization>``<id root="2.16.840.1.113883.19.5"/>``<name>Good Health Clinic</name>``<telecom nullFlavor="UNK"/>``<addr nullFlavor="UNK"/>`

1415

`</representedOrganization>``</assignedEntity>``</performer>``</observation>`

1420

`</entry>``</section>`**Figure 6.3.4.2-1: Medical History – Cardiac section example**

6.3.4.2.1 Problem Observation – Cardiac Constraints

1425 [observation: templateId 1.3.6.1.4.1.19376.1.4.1.9 (open)]

This Problem Observation – Cardiac entry is used exactly as specified in the CRC Common Entry Content Modules - section 6.3.5.1, except for vocabulary constraints.

1430 A Content Creator SHALL be able to include a Problem Observation – Cardiac Entry for each of the conditions identified in Value Set 1.3.6.1.4.1.19376.1.4.1.5.31 Cardiac Problems/Concerns. The value set for CONF:9058 (**value**) SHOULD be selected from ValueSet Cardiac Problems/Concerns (1.3.6.1.4.1.19376.1.4.1.5.31) **STATIC**.

A Content Creator SHALL be able to indicate the absence of the condition for the patient using the negation indicator.

1435 A Content Creator SHALL be able to include a Problem Observation – Cardiac Entry with code {160245001, SNOMED CT, “No current problems or disability”}.

6.3.4.2.2 Procedure Activity Observation - Constraints

[observation: templateId 2.16.840.1.113883.10.20.22.4.13 (open)]

This entry is used exactly as specified in C-CDA - section 5.48, except for vocabulary constraints for targetSiteCode.

1440 This entry is used to document the prior procedures for this patient that may be relevant to this cath procedure.

The value set for CONF:10121 (**targetSiteCode**) SHOULD be selected from ValueSet Body Site Value Set 1.3.6.1.4.1.19376.1.4.1.5.32 **STATIC**.

6.3.4.2.3 Procedure Activity Procedure - Constraints

1445 [procedure: templateId 2.16.840.1.113883.10.20.22.4.14 (open)]

This entry is used exactly as specified in C-CDA - section 5.49, except for vocabulary constraints for code and targetSiteCode.

This entry is used to document the prior procedures for this patient that may be relevant to this cath procedure. Prior procedures can include but are not limited to Cath, PCI and CABG.

1450 The value set for CONF:7657 (**code**) SHOULD be selected from ValueSet Cardiac Activity Procedures (1.3.6.1.4.1.19376.1.4.1.5.40) **STATIC**.

The value set for CONF:7683 (**targetSiteCode**) SHOULD be selected from ValueSet Body Site (1.3.6.1.4.1.19376.1.4.1.5.32) **STATIC**.

1455 **6.3.4.3 Allergies Section 48765-2**

[section: templateId 2.16.840.1.113883.10.20.22.2.6 (open)]

1460 This Allergies section content module lists and describes any medication allergies, adverse reactions, idiosyncratic reactions, anaphylaxis/anaphylactoid reactions to food items, and metabolic variations or adverse reactions/allergies to other substances (such as latex, iodine, tape adhesives) used to assure the safety of health care delivery. At a minimum, it should list currently active and any relevant historical allergies and adverse reactions.

This Allergies section content module is used exactly as specified in C-CDA - section 4.2.

1465 Within this Allergies section content module the Content Creator SHALL be able to create an Allergy Observation entry for each of the cardiac imaging agent classes identified in Value Set Contrast Agents Classes for Adverse Reactions (1.3.6.1.4.1.19376.1.4.1.5.34).

```

1470 <section>
    <templateId root="2.16.840.1.113883.10.20.22.2.6"/>
    <code code="48765-2"
        displayName="Allergies, adverse reactions, alerts"
        codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
    <title>Allergies</title>
    <text>
1475     The patient has allergies to penicillin based products
    </text>
    <entry typeCode="DRIV">
        <act classCode="ACT" moodCode="EVN">
1480         <templateId root="2.16.840.1.113883.10.20.22.4.30"/>
         <!-- Allergy Problem Act template -->
         ...
        </act>
        </entry>
    </section>

```

Figure 6.3.4.3-1: Allergies section example

1485 **6.3.4.3.1 Allergy Observation - Constraints**

[observation: templateId 2.16.840.1.113883.10.20.22.4.7 (open)]

This Allergy Observation entry content module is used exactly as specified in C-CDA - section 5.4, except for vocabulary constraints on CONF:10083.

1490 If the allergy is to Contrast Agents, the value set for CONF:10083 (**code**) SHALL be selected from ValueSet Contrast Agents Classes for Adverse Reactions (1.3.6.1.4.1.19376.1.4.1.5.34) **STATIC**.

6.3.4.4 Family History – Cardiac Section 10157-6

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.18 (open)]

1495 The Family History - Cardiac section content module contains data defining the patient’s genetic relatives in terms of possible or relevant health risk factors that have a potential impact on the patient’s healthcare risk profile.

This Family History – Cardiac section content module modifies the constraints defined for the C-CDA Family History Section (C-CDA 4.55). The complete set of constraints for this Family History – Cardiac section content module are listed below.

- 1500 1. **SHALL** contain exactly one [1..1] **templateId** (CONF:7932) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.18" (CONF:10388-CRC).
2. **SHALL** contain exactly one [1..1] **code/@code**="10157-6" Family History (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:7933).
3. **SHALL** contain exactly one [1..1] **title** (CONF:7934).
- 1505 4. **SHALL** contain exactly one [1..1] **text** (CONF:7935).
5. **SHALL** contain at least one [1..*] **Problem Observation**. (CONF:CRC-XXX)

```

1510 <section>
    <templateId root="1.3.6.1.4.1.19376.1.4.1.2.18"/>
    <!-- Family history section template -->
    <code code="10157-6" codeSystem="2.16.840.1.113883.6.1"/>
    <title>Family history</title>
    <text>No Family History of Cardiovascular Disease</text>
    <entry>
1515     <observation classCode="OBS" moodCode="EVN">
        <templateId root="2.16.840.1.113883.10.20.22.4.4"/>
        <id root="d11275e7-67ae-11db-bd13-0800200c9a66"/>
        <code code="404684003" codeSystem="2.16.840.1.113883.6.96"
1520             codeSystemName="SNOMED CT" displayName="Finding"/>
        <text>There was no family history of cardiovascular disease.</text>
        <statusCode code="completed"/>
        <effectiveTime>
        </effectiveTime>
        <value xsi:type="CD" code="160270001"
1525             codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                displayName=" No family history of cardiovascular disease"/>
        </observation>
    </entry>
</section>

```

1530 **Figure 6.3.4.4-1: Family History – Cardiac section example**

6.3.4.4.1 Problem Observation - Constraints

[Observation: templateId 2.16.840.1.113883.10.20.22.4.4 (open)]

1535 The Problem Observation entry content module is used exactly as specified in C-CDA - section 5.45, except for vocabulary constraints on CONF:9058.

The value set for CONF:9058 (**code**) **SHOULD** be selected from ValueSet Cardiovascular Family History (1.3.6.1.4.1.19376.1.4.1.5.33) **STATIC**.

6.3.4.5 Social History Section 29762-2

1540 [section: templateId 2.16.840.1.113883.10.20.22.2.17 (open)]

The Social History section content module is used exactly as specified in C-CDA - section 4.57.

1545 This Social History section content module contains data defining the patient's occupational, personal (e.g., lifestyle), social, and environmental history and health risk factors, as well as administrative data such as marital status, race, ethnicity and religious affiliation. Social history can have significant influence on a patient's physical, psychological and emotional health and wellbeing so should be considered in the development of a complete record.

```

1550 <section>
    <templateId root="2.16.840.1.113883.10.20.22.2.17"/>
    <!-- ** Social history section template ** -->
    <code code="29762-2" codeSystem="2.16.840.1.113883.6.1"
        displayName="Social History"/>
    <title>Social History</title>
    <text>
1555     The patient was a former smoker.
    </text>
    <entry typeCode="DRIV">
        <observation classCode="OBS" moodCode="EVN">
1560         <templateId root="2.16.840.1.113883.10.20.22.4.38"/>
        <!-- ** Social history observation template ** -->
        <id root="45efb604-7049-4a2e-ad33-d38556c9636c"/>
        <code code="229819007" codeSystem="2.16.840.1.113883.6.96"
            displayName="Tobacco use and exposure"/>
1565         <statusCode code="completed"/>
        <effectiveTime>
            <low value="1973"/>
        </effectiveTime>
        <value xsi:type="CD" code="8517006"
            codeSystem="2.16.840.1.113883.6.96"
            displayName="Former Smoker"/>
1570     </observation>
    </entry>
</section>

```

Figure 6.3.4.5-1: Social History section example

1575 **6.3.4.5.1 Social History Observation - Constraints**

[observation: templateId 2.16.840.1.113883.10.20.22.4.38 (open)]

The Social History Observation entry content module is used exactly as specified in C-CDA - section 5.61.

1580 To include smoking status observations, the values allowed for CONF:8559 (**value**) SHOULD be selected from ValueSet Smoking History 1.2.840.10008.6.1.225 STATIC.

To include cocaine misuse behavior, the value allowed for CONF:8559 (**value**) SHOULD be “78267003” from SNOMED CT “Cocaine abuse”.

6.3.4.6 Physical Exam Section 29545-1

[section: templateId 2.16.840.1.113883.10.20.2.10 (open)]

1585 The Physical Exam section content module is used exactly as specified in C-CDA - section 5.5.

1590 The Physical Exam section content module includes direct observations made by the clinician. The examination may include the use of simple instruments and may also describe simple maneuvers performed directly on the patient’s body. This Physical Exam section includes only observations made by the examining clinician using inspection, palpation, auscultation, and percussion; it does not include laboratory or imaging findings. The exam may be limited to pertinent body systems based on the patient’s chief complaint or it may include a comprehensive examination. The examination may be reported as a collection of random clinical statements or it may be reported categorically.

1595

```
<section>
  <templateId root="2.16.840.1.113883.10.20.2.10"/>
  <code codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"
    code="29545-1" displayName="PHYSICAL FINDINGS"/>
  <title>PHYSICAL EXAMINATION</title>
  <text>
    <paragraph>All normal to examination.</paragraph>
  </text>
</section>
```

1600

Figure 6.3.4.6-1: Physical Exam section example

1605 **6.3.4.7 Vital Signs Section 8716-3**

[section: templateId 2.16.840.1.113883.10.20.22.2.4 (open)]

The Vital Signs Section content module is used exactly as specified in C-CDA - section 4.60.

The Vital Signs section content module is intended to include vital sign measurements taken at admission and at the time of procedure, if feasible.

1610

```

1615 <section>
      <templateId root="2.16.840.1.113883.10.20.22.2.4"/>
      <code code="8716-3"
1620         codeSystem="2.16.840.1.113883.6.1"
            codeSystemName="LOINC"
            displayName="VITAL SIGNS" />
      <title>Vital Signs</title>
      <text>These are the vital signs related to the procedure </text>
      <entry typeCode="DRIV">
1625         <organizer classCode="CLUSTER" moodCode="EVN">
              <templateId root="2.16.840.1.113883.10.20.22.4.26"/>
              <!-- Vital signs organizer template -->
              <id root="c6f88320-67ad-11db-bd13-0800200c9a66"/>
              <code code="46680005" codeSystem="2.16.840.1.113883.6.96"
1630                 codeSystemName="SNOMED CT" displayName="Vital signs"/>
              <statusCode code="completed"/>
              <effectiveTime value="19991114"/>
              <component>
1635                 <observation classCode="OBS" moodCode="EVN">
                      <templateId root="2.16.840.1.113883.10.20.22.4.27"/>
                      <!-- Vital sign observation for height -->
                      <id root="c6f88321-67ad-11db-bd13-0800200c9a66"/>
                      <code code="8302-2"
1640                          codeSystem="2.16.840.1.113883.6.1"
                              codeSystemName="LOINC"
                              displayName="Height"/>
                      <text><reference value="#height1"/></text>
                      <statusCode code="completed"/>
                      <effectiveTime value="19991114"/>
                      <value xsi:type="PQ" value="177" unit="cm"/>
                      <interpretationCode code="N" codeSystem="2.16.840.1.113883.5.83"/>
1645                  </observation>
                </component>
              </organizer>
            </entry>
1650 </section>

```

Figure 6.3.4.7-1: Vital Signs section example

6.3.4.7.1 Vital Sign Observation - Constraints

1650 [observation: templateId 2.16.840.1.113883.10.20.22.4.27 (open)]

The Vital Sign Observation entry content module is used exactly as specified in C-CDA - section 5.65, except for vocabulary constraints.

The value set for CONF:7301 (**code**) SHOULD be selected from ValueSet Vital Sign Result Type 1.3.6.1.4.1.19376.1.4.1.5.36 STATIC.

1655 **6.3.4.8 Pre-Procedure Results – Cardiac Section 30954-2**

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.23 (open)]

(([section: templateId 2.16.840.1.113883.10.20.22.2.3 (open)] – parent)

This Pre-Procedure Results – Cardiac Section content module is based on the C-CDA Results Section content module as specified in C-CDA - section 4.55.

1660 The Pre-Procedure Results – Cardiac section content module contains the results of pre-procedure tests that are required to prepare for the cath procedure. Results from prior diagnostic cath procedures are included here if they provide indications for the current interventional procedure. There may also be a reference to an optional external document in the result organizer.

1665 This Pre-Procedure Results – Cardiac section content module contains the results of observations generated by laboratories, imaging procedures, and other procedures. The scope includes observations such as hematology, chemistry, serology, virology, toxicology, microbiology, plain x-ray, ultrasound, CT, MRI, angiography, echocardiography, nuclear medicine, pathology, and procedure observations. The section often includes notable results such as abnormal values or relevant trends, and could contain all results for the period of time being documented.

1670 Imaging results are typically generated by a clinician reviewing the output of an imaging procedure, such as where a cardiologist reports the left ventricular ejection fraction based on the review of a cardiac echocardiogram.

1675 This Pre-Procedure Results – Cardiac section content module modifies the Results Section (C-CDA 4.55). The complete set of constraints for the Pre-Procedure Results – Cardiac section content module are defined below. **The substitutions are highlighted in yellow.** This Pre-Procedure Results – Cardiac section content module is also conformant to the C-CDA Results Section content module.

- 1680 1. **SHALL** contain two or more [2..*] **templateId** (CONF:7108-CRC) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="2.16.840.1.113883.10.20.22.2.3" (CONF:9136).
 - b. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.23" (CONF:CRC-xxx).
- 1685 2. **SHALL** contain exactly one [1..1] **code/@code**="30954-2" Relevant diagnostic tests and/or laboratory data (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:7110).
3. **SHALL** contain exactly one [1..1] **title** (CONF:8892).
4. **SHALL** contain exactly one [1..1] **text** (CONF:7111).
- 1690 5. **SHALL** contain at least one [1..*] **entry** (CONF:7112) such that it
 - a. **SHALL** contain exactly one [1..1] **Result Organizer - Cardiac** (1.3.6.1.4.1.19376.1.4.1.4.11) (CONF:7113-CRC).

1695	<pre> <section> <templateId root="1.3.6.1.4.1.19376.1.4.1.2.23"/> <templateId root="2.16.840.1.113883.10.20.22.2.3"/> <code code="30954-2" </pre>
1700	<pre> codeSystem="2.16.840.1.113883.6.1"/> codeSystemName="LOINC" displayName="RESULTS" /> <title>Results</title> <text> </pre>
1705	<pre> ... </text> <entry typeCode="DRIV"> <organizer classCode="CLUSTER" moodCode="EVN"> </pre>
1710	<pre> <templateId root="1.3.6.1.4.1.19376.1.4.1.4.11"/> <id root="7d5a02b0-67a4-11db-bd13-0800200c9a66"/> <code code="57021-8" displayName="CBC W Auto Differential panel" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> <statusCode code="completed"/> <component> </pre>
1715	<pre> <observation classCode="OBS" moodCode="EVN"> <!-- Result observation template --> <templateId root="2.16.840.1.113883.10.20.22.4.2"/> <id root="107c2dc0-67a5-11db-bd13-0800200c9a66"/> <code code="30313-1" displayName="HGB" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> <statusCode code="completed"/> <effectiveTime value="200003231430"/> <value xsi:type="PQ" value="13.2" unit="g/dl"/> <interpretationCode code="N" codeSystem="2.16.840.1.113883.5.83"/> <methodCode/> <targetSiteCode/> <referenceRange> </pre>
1720	<pre> <observationRange> <text>M 13-18 g/dl; F 12-16 g/dl</text> </observationRange> </referenceRange> </observation> </component> <reference typeCode="REFR"> </pre>
1735	<pre> <externalDocument> <id root="b50b7910-7ffb-4f4c-bbe4-177ed68cbbf3"/> <text mediaType="application/pdf"> <reference value="PreProcedureResults.pdf"/> </text> </externalDocument> </reference> </organizer> </entry> </section> </pre>
1740	
1745	

Figure 6.3.4.8-1: Pre-Procedure Results section example

6.3.4.8.1 Result Organizer - Cardiac

[organizer: templateId 1.3.6.1.4.1.19376.1.4.1.4.11 (open)]

([organizer: templateId 2.16.840.1.113883.10.20.22.4.1 (open)] – parent)

1750

This clinical statement identifies a set of result observations. It contains information applicable to all of the contained result observations. Result type codes categorize a result into one of several commonly accepted values (e.g., "Diagnostic Cath", "PCI", "Diagnostic Cath and PCI"). These values are often implicit in the Result Organizer code (e.g., an Organizer/code of "complete blood count" implies a Result Observation code of "Hematology").

1755

An appropriate nullFlavor can be used when a single result observation is contained in the organizer, and organizer/code or organizer/id is unknown.

There may also be a reference to an optional external document in the result organizer.

1760

This Result Organizer – Cardiac entry content module extends the C-CDA Result Organizer entry definition (C-CDA 5.57) by adding the constraints listed below.

1. **SHALL** contain exactly one [1..1] **templateId** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@root="1.3.6.1.4.1.19376.1.4.1.4.11"** (CONF:CRC-xxx).
2. **MAY** contain zero or more [0..*] **reference** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@typeCode="REFR"** Refers to (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:CRC-xxx).
 - b. **SHALL** contain exactly one [1..1] **externalDocument** (CONF:CRC-xxx).
 - i. This externalDocument **SHALL** contain at least one [1..*] **id** (CONF:CRC-xxx).
 - ii. This externalDocument **MAY** contain zero or one [0..1] **text** (CONF:CRC-xxx).
 1. The text, if present, **MAY** contain zero or one [0..1] **@mediaType** (CONF:CRC-xxx).
 2. The text, if present, **MAY** contain zero or one [0..1] **reference** (CONF:CRC-xxx).
 - a. The URL of a referenced pre-procedure results document **MAY** be present, and **SHALL** be represented in organizer/reference/ExternalDocument/text/reference (CONF:CRC-xxx).
 - b. If a URL is referenced, then it **SHOULD** have a corresponding linkHTML element in narrative block (CONF:CRC-xxx).

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6.3.4.8.2 Result Observation - Constraints

[observation: templateId 2.16.840.1.113883.10.20.22.4.2 (open)]

1790 This clinical statement represents details of a lab, radiology, or other study performed on a patient.

This Result Observation entry is used exactly as specified in C-CDA - section 5.56 except for vocabulary constraints for the code and value elements.

1795 The value set for CONF:7166 (**code**) **SHOULD** be selected from LOINC (CodeSystem: 2.16.840.1.113883.6.1) or SNOMED CT using the Value Set Result Observation (1.3.6.1.4.1.19376.1.4.1.5.38) **STATIC**.

The value set for CONF:9109 (**code**) **SHOULD** be selected from ValueSet Cardiac Lab Results (1.3.6.1.4.1.19376.1.4.1.5.35) **STATIC**.

The value set for CONF:7153 (**value**) **SHOULD** be selected from ValueSet Body Site (1.3.6.1.4.1.19376.1.4.1.5.32) **STATIC**.

1800 6.3.4.9 Planned Procedure Section 59772-4

[section: templateId 2.16.840.1.113883.10.20.22.2.30 (open)]

The Planned Procedure section is used exactly as specified in C-CDA - section 4.40.

1805 The Planned Procedure section records the procedure(s) that a physician or clinician thought would need to be done based on the preoperative assessment. Procedures include but are not limited to Diagnostic Cath, Angiography, and PCI. It may be important to record the procedure(s) that were originally planned for, consented to, and perhaps pre-approved by the payor, particularly if different from the actual procedure(s) and procedure details, to provide evidence to various stakeholders that the providers are aware of the discrepancy and the justification can be found in the procedure details.

1810

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```

<section>
  <templateId root="2.16.840.1.113883.10.20.22.2.30"/>
  <!-- ***** Planned Procedure Section template ***** -->
  <code code="59772-4" codeSystem="2.16.840.1.113883.6.1"
    codeSystemName="LOINC" displayName="Planned Procedure"/>
  <title>Planned Procedure</title>
  <text>
    A diagnostic catheterization is planned.
  </text>
  <entry>
    <procedure moodCode="RQO" classCode="PROC">
      <templateId root="2.16.840.1.113883.10.20.22.4.41"/>
      <!-- ** Plan of Care Activity Procedure Template ** -->
      <id root="9a6d1bac-17d3-4195-89c4-1121bc809b5a"/>
      <code code="41976001" codeSystem="2.16.840.1.113883.6.96"
        displayName="Diagnostic Catheterization"/>
      <statusCode code="new"/>
      <effectiveTime>
        <center value="20000421"/>
      </effectiveTime>
    </procedure>
  </entry>
</section>

```

Figure 6.3.4.9-1: Planned Procedure section example

1835 6.3.4.10 Procedure Indications Section 59768-2

[section: templateId 2.16.840.1.113883.10.20.22.2.29 (open)]

The Procedure Indications section content module is used exactly as specified in C-CDA - section 4.50.

1840

The Procedure Indications section content module records details about the reason for this Cath/PCI procedure. This Procedure Indications section content module may include the pre-procedure diagnosis or diagnoses as well as one or more symptoms that contribute to the reason the procedure is being performed.

1845

```

<section>
  <templateId root="2.16.840.1.113883.10.20.22.2.29"/>
  <code code="59768-2" codeSystem="2.16.840.1.113883.6.1"
    codeSystemName="LOINC" displayName="PROCEDURE INDICATIONS"/>
  <title>Procedure Indications</title>
  <text>The procedure is performed for screening in a low risk individual.
</text>
</entry>
  <observation classCode="OBS" moodCode="EVN">
    <!-- Indication Entry -->
    <templateId root="2.16.840.1.113883.10.20.22.4.19"/>
    <code code="409586006"
      codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT" displayName="Complaint"/>
    <statusCode code="completed"/>
    <value xsi:type="CD"
      code="29857009" codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT" displayName="Chest pain"/>
  </observation>
</entry>
</section>

```

1850

1855

1860

1865

Figure 6.3.4.10-1: Procedure Indications section example

6.3.4.10.1 Indication - Constraints

[observation: templateId 2.16.840.1.113883.10.20.22.4.19 (open)]

This Indication entry content module is used exactly as specified in C-CDA - section 5.25 except for vocabulary constraints.

1870

The Indication entry content module documents the rationale for an activity. It can do this with the id element to reference a problem recorded elsewhere in the document or with a code and value to record the problem type and problem within the Indication. For example, the indication for Diagnostic Catheterization might be chest pain.

1875

The value set for CONF:7657 (**value**) **SHOULD** be selected from ValueSet Procedure Indications (1.3.6.1.4.1.19376.1.4.1.5.37) **STATIC**.

6.3.4.11 Anesthesia Section 59774-0

[section: templateId 2.16.840.1.113883.10.20.22.2.25 (open)]

The Anesthesia section content module is used exactly as specified in C-CDA - section 4.3.

1880

The Anesthesia section content module briefly describes the general anesthesia used and may state the actual agent used. The Procedure Activity Procedure entry content module describes the anesthesia procedure. The Medication Activity entry content module may describe the general anesthesia medication used during this cath procedure.

```

1885 <section>
      <templateId root="2.16.840.1.113883.10.20.22.2.25"/>
      <code code="59774-0"
1890         codeSystem="2.16.840.1.113883.6.1"
            codeSystemName="LOINC"
            displayName="PROCEDURE ANESTHESIA"/>
      <title>Procedure Anesthesia</title>
      <text> Conscious sedation with propofol 200 mg IV </text>
      <entry>
1895         <procedure classCode="PROC" moodCode="EVN">
            <!-- Procedure activity procedure template -->
            <templateId root="2.16.840.1.113883.10.20.22.4.14"/>
            <id root="e401f340-7be2-11db-9fe1-0800200c9a66"/>
            <code code="415070008" codeSystem="2.16.840.1.113883.6.96"
1900                 displayName="PCI">
                <originalText> PCI <reference value="procedure1"/></originalText>
            </code>
            <text>
1905                 <reference value="procedure1"/>
            </text>
            <statusCode code="completed"/>
            <effectiveTime value="201109261015"/>
            <targetSiteCode code="41879009"
1910                 codeSystem="2.16.840.1.113883.6.96"
                    displayName="Distal RCA"/>
            <participant typeCode="DEV">
                <participantRole classCode="MANU">
1915                 <!-- Product instance template -->
                    <templateId root="2.16.840.1.113883.10.20.22.4.37"/>
                    ...
                </participantRole>
            </participant>
            <entryRelationship typeCode="COMP" inversionInd="true">
                <substanceAdministration classCode="SBADM" moodCode="INT">
1920                 <!-- Medication activity template -->
                    <templateId root=" 2.16.840.1.113883.10.20.22.4.16"/>
                    ...
                </substanceAdministration>
            </entryRelationship>
            </procedure>
1925 </entry>
        <entry>
            <substanceAdministration classCode="SBADM" moodCode="EVN">
                <!-- Medication activity template -->
1930                 <templateId root="2.16.840.1.113883.10.20.22.4.16"/>
                    ...
            </substanceAdministration>
        </entry>
    </section>

```

Figure 6.3.4.11-1: Anesthesia section example

1935

6.3.4.11.1 Procedure Activity Procedure - Constraints

[procedure: templateId 2.16.840.1.113883.10.20.22.4.14 (open)]

The Procedure Activity Procedure entry content module is used exactly as specified in C-CDA - section 5.49, except for vocabulary constraints.

1940 The Procedure Activity Procedure entry content module describes the anesthesia procedure.

The value set for CONF:7657 (**code**) SHOULD be selected from ValueSet Cardiac Activity Procedures (1.3.6.1.4.1.19376.1.4.1.5.40) **STATIC**.

The value set for CONF:7683 (**targetSiteCode**) SHOULD be selected from ValueSet Body Site Value Set 1.3.6.1.4.1.19376.1.4.1.5.32 **STATIC**.

1945 6.3.4.12 Medications Administered Section 29549-3

[section: templateId 2.16.840.1.113883.10.20.22.2.38 (open)]

This Medications Administered section content module is used exactly as specified in C-CDA - section 4.32 except for vocabulary constraints.

1950 The Medications Administered section content module defines medications and fluids administered during the procedure, encounter, or other activity excluding general anesthetic medications.

1955 A Content Creator SHALL be able to create a Medications Activity entry with a Medication Information Entry for each of the cardiac medication classes identified in Value Set 1.3.6.1.4.1.19376.1.4.1.5.41 Drug Classes and Specific Cardiac Drugs Used in Cardiac Procedures.

A Content Creator SHALL be able to create a Medications Activity entry with a Medication Information entry for the relevant cardiac contrast agents identified in Value Set 1.3.6.1.4.1.19376.1.4.1.5.39 Contrast Agents.

1960 The set of contrast agents implemented may be limited to a subset of the Value Set, based on the types of procedures for which the Content Creator creates reports, hence the term “*relevant cardiac contrast agents*”.

1965	<section>
	<templateId root="2.16.840.1.113883.10.20.22.2.38" />
	<code code="29549-3"
	codeSystem="2.16.840.1.113883.6.1"
	codeSystemName="LOINC"
	displayName="MEDICATIONS ADMINISTERED" />
1970	<title>Medications Administered</title>
	<text>Aspirin, other antiplatelet agents</text>
	<entry>
	<substanceAdministration classCode="SBADM" moodCode="EVN">
	<templateId root="2.16.840.1.113883.10.20.22.4.16"/>
1975	<!-- Medication Activity template -->
	<id root="cdbd33f0-6cde-11db-9fe1-0800200c9a66"/>
	<text>
	<reference value="#med1"/>
	Aspirin, other antiplatelet agents
1980	</text>
	<statusCode code="completed"/>
	<effectiveTime xsi:type="IVL_TS">
	<low value="20110926"/>
	<high value="20111014"/>
	</effectiveTime>
1985	<effectiveTime xsi:type="PIVL_TS" institutionSpecified="true"
	operator="A">
	<period value="6" unit="h"/>
	</effectiveTime>
	<doseQuantity value="1"/>
1990	<consumable>
	<manufacturedProduct classCode="MANU">>
	<templateId root="2.16.840.1.113883.10.20.22.4.23"/>
	<!-- Medication Information template -->
1995	<id/>
	<manufacturedMaterial>
	<code code="7947003"
	codeSystem="2.16.840.1.113883.6.96"
	displayName="Aspirin"/>
	</manufacturedMaterial>
2000	<manufacturerOrganization>...</manufacturerOrganization>
	</manufacturedProduct>
	</consumable>
	<performer>
	</substanceAdministration>
2005	</entry>
	</section>

Figure 6.3.4.12-1: Medications administered section example

6.3.4.12.1 Medication Information - Constraints

[manufacturedProduct: templateId 2.16.840.1.113883.10.20.22.4.23 (open)]

2010 This Medication Information entry is used exactly as specified in C-CDA - section 5.29 except for vocabulary constraints.

The value set for CONF:7412 ([manufacturedMaterial/code@code](#)) **SHOULD** be selected from ValueSet Medication Clinical Drug (1.3.6.1.4.1.19376.1.4.1.5.41) **STATIC** or **SHOULD** be selected from ValueSet Contrast Agents (1.3.6.1.4.1.19376.1.4.1.5.39) **STATIC**.

2015 **6.3.4.13 Procedure Description – Cardiac Section 29554-3**

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.19 (open)]

(([section: templateId 2.16.840.1.113883.10.20.22.2.27 (open)] – parent)

2020 The Procedure Description – Cardiac section content module records the details of the cardiac procedures and may include procedure site preparation, surgical site preparation, pertinent details related to sedation/anesthesia, pertinent details related to measurements and markings, procedure times, medications administered, estimated blood loss, specimens removed, instrumentation, sponge counts, tissue manipulation, wound closure, sutures used, vital signs and other monitoring data. Local practice often identifies the level and type of detail required based on the procedure or specialty.

2025 This Procedure Description – Cardiac section content module may include a device organizer to record information about each device used during the procedures. All devices should be defined at this section level within a Procedure Device Organizer – Cardiac entry. Additional characteristics inherent to these devices, like length and diameter, should be defined using an additional Procedure Device Organizer – Cardiac entry within this section. In addition, dynamic attributes of these devices, like balloon inflation atmospheres, should be recorded in the
 2030 Procedure Activity Procedure – Cardiac entry within this section content module.

2035 For PCI procedures, individual lesions will be defined in this section as separate lesion observations identified by a unique “lesion ID”. Only the location of the lesion will be identified here. Procedures, procedure findings, and results can then reference to the lesion to which it is related by creating an entryRelationship of type=”REFR” to the lesion observation based on the “lesion ID” within the Procedure Activity Procedure – Cardiac entry.

This Procedure Description – Cardiac section content module extends the C-CDA Procedure Description section (C-CDA 4.45) by adding the constraints listed below.

- 2040
1. **SHALL** contain exactly one [1..1] **templateId** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.19" (CONF:CRC-xxx).
 2. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] [Procedure Device Organizer - Cardiac](#) (1.3.6.1.4.1.19376.1.4.1.4.12) (CONF:CRC-xxx).
 - 2045 3. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] [Lesion Observation](#) (1.3.6.1.4.1.19376.1.4.1.4.10) (CONF:CRC-xxx). These identify the lesions including where they are located.
 4. **SHALL** contain at least one [1..*] **entry** (CONF:CRC-xxx) such that it

2050

- a. **SHALL** contain exactly one [1..1] [Procedure Activity Procedure - Cardiac](#) (1.3.6.1.4.1.19376.1.4.1.4.14) (CONF:CRC-xxx).

```

2055 <section>
      <templateId root="1.3.6.1.4.1.19376.1.4.1.2.19"/>
      <templateId root="2.16.840.1.113883.10.20.22.2.27"/>
      <!-- Procedure Description - Cardiac section template -->
      <code code="29554-3"
2060         codeSystem="2.16.840.1.113883.6.1"
         codeSystemName="LOINC"
         displayName="PROCEDURE DESCRIPTION" />
      <title>Procedures</title>
      <text>
        This is the narrative for this section...
      </text>
2065 <entry>
      <!-- Procedure Device Organizer for device inventory - this could include
the guide wire, the balloon and the stent... -->
      <organizer classCode="CLUSTER" moodCode="EVN">
2070         <templateId root="1.3.6.1.4.1.19376.1.4.1.4.12"/>
         <participant typecode="SUBJ">
           <participantRole classCode="MANU">
             <id root=" eb936010-7b17-11db-9fe1-0800200c9b66">
2075               <playingDevice> <!-- guidewire -->
                 <code code="272224001" codeSystem="2.16.840.1.113883.6.96"
                   displayName="guide wire"
                   <id root=" eb936010-7b17-11db-9fe1-0800200c9b67">
                 </playingDevice>
                 <scopingEntity>
2080                   <id root="eb936010-7b17-11db-9fe1-0800200c9b65"/>
                 </scopingEntity>
               </participantRole>
             </participant>
           </organizer>
         </entry>
2085 <entry>
      <!-- Organizer for specific device with observations (e.g. size/dimensions)
-->
      <organizer classCode="CLUSTER" moodCode="EVN">
2090       <!-- Procedure Device Organizer template -->
       <templateId root="1.3.6.1.4.1.19376.1.4.1.4.12"/>
       <participant typecode="SUBJ">
         <participantRole classCode="MANU">
           <id root=" eb936010-7b17-11db-9fe1-0800200c9b68">
2095             <playingDevice> <!-- stent -->
               <code code="3831886012"
                 codeSystem="2.16.840.1.113883.6.96"
                 displayName="JJ-stent"
                 <id root="eb936010-7b17-11db-9fe1-0800200c9b69">
             </playingDevice>
             <scopingEntity>
2100               <id root="eb936010-7b17-11db-9fe1-0800200c9b65"/>
             </scopingEntity>
           </participantRole>
         </participant>
2105 <component>
       <observation classCode="SUBJ" moodCode="EVN">

```

2110

```

    <!-- Device Observation template -->
    <templateId root="1.3.6.1.4.1.19376.1.4.1.4.13"/>
    <id root=" eb936010-7b17-11db-9fe1-0800200c9b6a">
    <code code="408706001" displayName="vascular stent diameter"
          codeSystem="2.16.840.1.113883.6.96"
          codeSystemName="SNOMED CT"/>

```

2115

```

    <statusCode code="completed"/>
    <effectiveTime value="201109261015"/>
    <value xsi:type="PQ" value="13.2" unit="mm"/>
  </observation>
</component>
<component>

```

2120

```

  <observation classCode="SUBJ" moodCode="EVN">>
  <!-- Device Observation template -->
  <templateId root="1.3.6.1.4.1.19376.1.4.1.4.13"/>
  <id root=" eb936010-7b17-11db-9fe1-0800200c9b6a">
  <code code="408703009" displayName="vascular stent length"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT"/>

```

2125

```

  <statusCode code="completed"/>
  <effectiveTime value="201109261015"/>
  <value xsi:type="PQ" value="11.8" unit="mm"/>
  </observation>
</component>
</organizer>
</entry>

```

2130

```

<!-- define lesions by indicating the targetSiteCodes where located -->
<entry>
  <observation classCode="OBS" moodCode="EVN">
  <!-- Lesion Observation template -->
  <templateID root="1.3.6.1.4.1.19376.1.4.1.4.10" />
  <id root="2.840.110893.98120.74.8" ext="lesion #1" />
  <code code="404684003"
        codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
        displayName="Finding" />
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="40415009"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2135

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2140

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2145

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2150

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2155

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

2160

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

```

  <value code="40415009"
        codeSystem="2.16.840.1.113883.6.96"
        codeSystemName="SNOMED CT" displayName="proximal" />
  </qualifier>
</targetSiteCode>
  <targetSiteCode code="56322004"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Left PDA">
  <qualifier>
    <value code="255562008"
           codeSystem="2.16.840.1.113883.6.96"
           codeSystemName="SNOMED CT" displayName="mid" />
  </qualifier>
</targetSiteCode>

```

```

2165     </observation>
2170   </entry>
2175   <entry typeCode="DRIV">
2180     <procedure classCode="PROC" moodCode="EVN">
2185       <!-- Procedure Activity Procedure - Cardiac template -->
2190       <templateId root="1.3.6.1.4.1.19376.1.4.1.4.14"/>
2195       <templateId root="2.16.840.1.113883.10.20.22.4.14"/>
2200       <id root="CPAC1"/>
2205       <code code="415070008" codeSystem="2.16.840.1.113883.6.96"
2210         displayName="PCI">
2215         <originalText>PCI<reference value="procedure1"/></originalText>
2220       </code>
2225       <text>
2230         <reference value="procedure1"/>
2235       </text>
2240       <statusCode code="completed"/>
2245       <effectiveTime value="201109261015"/>
2250       <targetSiteCode code="41879009" codeSystem="2.16.840.1.113883.6.96"
2255         displayName="Left PDA"/>
2260     <participant typeCode="PRD">
2265       <participantRole classCode="MANU">
2270         <!-- Product instance template -->
2275         <templateId root="2.16.840.1.113883.10.20.22.4.37"/>
2280         <playingDevice>
2285           <id root="P123">
2290             <code code="102319006" codeSystem="2.16.840.1.113883.6.96"
2295               displayName="Percutaneous transluminal angioplasty
2300                 balloon, device (physical object)"/>
2305           </code>
2310         </playingDevice>
2315         <scopingEntity>
2320           <id root="eb936010-7b17-11db-9fe1-0800200c9b65"/>
2325         </scopingEntity>
2330       </participantRole>
2335     </participant>
2340   <participant typeCode="PRD">
2345     <participantRole classCode="MANU">
2350       <!-- Product instance template -->
2355       <templateId root="2.16.840.1.113883.10.20.22.4.37"/>
2360       <playingDevice>
2365         <id root="G456">
2370           <code code="272224001" codeSystem="2.16.840.1.113883.6.96"
2375             displayName="guide wire"/>
2380         </code>
2385       </playingDevice>
2390       <scopingEntity>
2395         <id root="eb936010-7b17-11db-9fe1-0800200c9b65"/>
2400       </scopingEntity>
2405     </participantRole>
2410   </participant>
2415 </entry>
2420   <organizer classCode="CLUSTER" moodCode="EVN">
2425     <participant typeCode="PRD">
2430       <participantRole classCode="MANU">
2435         <playingDevice>
2440           <id root="p123">

```

```

2215         <code code="102319006"
                codeSystem="2.16.840.1.113883.6.96"
                displayName=" Percutaneous transluminal angioplasty
                    balloon, device (physical object)"/>
2220     </playingDevice>
     <scopingEntity>
2225         <id root="eb936010-7b17-11db-9fe1-0800200c9b65"/>
     </scopingEntity>
     </participantRole>
</participant>
2225 <component>
     <observation classCode="OBS" moodCode="EVN">
2230         <!-- Device observation template -->
         <templateId root="1.3.6.1.4.1.19376.1.4.1.4.13"/>
         <id root=" eb936010-7b17-11db-9fe1-0800200c9b6b"/>
         <code code="371851006" displayName="angioplasty inflation
pressure"
                codeSystem="2.16.840.1.113883.6.96"
                codeSystemName="SNOMED CT"/>
2235         <statusCode code="completed"/>
         <effectiveTime value="201109261015"/>
         <value xsi:type="PQ" value="13.2" unit="[ATM]"/>
     </observation>
     </component>
     <component>
2240         <observation classCode="OBS" moodCode="EVN">
         <!-- Device observation template -->
         <templateId root="1.3.6.1.4.1.19376.1.4.1.4.13"/>
         <id root=" eb936010-7b17-11db-9fe1-0800200c9b6c"/>
         <code code="371852004"
2245             displayName="angioplasty inflation duration"
             codeSystem="2.16.840.1.113883.6.96"
             codeSystemName="SNOMED CT"/>
         <statusCode code="completed"/>
         <effectiveTime value="201109261015"/>
2250         <value xsi:type="PQ" value="11.6" unit="s"/>
     </observation>
     </component>
     </organizer>
2255 </entry>

<!-- link to the lesion for this procedure which was defined previously in this
section -->
     <entryRelationship typeCode="REFR">
2260         <observation classCode="OBS" moodCode="EVN">
         <templateID root="1.3.6.1.4.1.19376.1.4.1.10" />
         <id root="2.840.110893.98120.74.8" ext="lesion #1" />
         <code code="404684003"
2265             codeSystem="2.16.840.1.113883.6.96"
             codeSystemName="SNOMED CT" displayName="Finding" />
     </observation>
     </entryRelationship>
</procedure>
</entry>

```

</section>

2270

Figure 6.3.4.13-1: Procedure Description - Cardiac section example

6.3.4.13.1 Procedure Activity Procedure - Cardiac

[procedure: templateId 1.3.6.1.4.1.19376.1.4.1.4.14 (open)]

([procedure: templateId 2.16.840.1.113883.10.20.22.4.14 (open)] – parent)

2275

This clinical statement represents procedures whose immediate and primary outcome (post-condition) is the alteration of the physical condition of the patient. Examples of these procedures are a diagnostic cardiac catheterization and PCI.

2280

This Procedure Activity Procedure – Cardiac entry content module may also include a device organizer to record specific properties of the devices as observed during the procedure. Dynamic attributes of these devices, like balloon inflation atmospheres, should be recorded in this Procedure Activity Procedure – Cardiac entry.

Within this Procedure Activity Procedure – Cardiac entry content module, Product Instances are used to document the devices used. Record as many devices as needed unless the cath lab procedure is aborted. In this case, there may be no devices used.

2285

Developers using this CRC content profile will map specific equipment using appropriate inventory numbering and product descriptions provided by the hemodynamic monitoring system or equivalents. If this CRC content profile is to be consumed and used in a CVIS, it is up to the developer to map the actual codes to the appropriate ACC NCDR-Cath/PCI codes.

2290

This Procedure Activity Procedure – Cardiac entry content module is used exactly as specified in C-CDA - section 5.49 except for the modifications to the constraints highlighted in yellow below. This Procedure Activity Procedure – Cardiac entry content module is also conformant to the C-CDA Procedure Activity Procedure entry content module.

2295

1. **SHALL** contain exactly one [1..1] @classCode="PROC" Procedure (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:7652).
2. **SHALL** contain exactly one [1..1] @moodCode, which **SHALL** be selected from ValueSet MoodCodeEvnInt 2.16.840.1.113883.11.20.9.18 **STATIC** 2011-04-03 (CONF:7653).

2300

3. **SHALL** contain two or more [2..*] templateId (CONF:7654-CRC) such that
 - a. **SHALL** contain exactly one [1..1] @root="2.16.840.1.113883.10.20.22.4.14" (CONF:10521).
 - b. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.4.1.4.14" (CONF:7654-CRC-xxx).

2305

4. **SHALL** contain at least one [1..*] id (CONF:7655).
5. **SHALL** contain exactly one [1..1] code (CONF:7656).
 - a. This code in a procedure activity **SHOULD** be selected from LOINC (codeSystem 2.16.840.1.113883.6.1) or SNOMED CT (CodeSystem:

- 2310 2.16.840.1.113883.6.96), and **MAY** be selected from CPT-4 (CodeSystem: 2.16.840.1.113883.6.12), ICD9 Procedures (CodeSystem: 2.16.840.1.113883.6.104), ICD10 Procedure Coding System (CodeSystem: 2.16.840.1.113883.6.4). The recommended Value Set can be found in Cardiac Activity Procedures (1.3.6.1.4.1.19376.1.4.1.5.40). (CONF:7657-CRC)
- 2315 b. This code **SHOULD** contain zero or one [0..1] **originalText** (CONF:7658).
 i. The originalText, if present, **SHOULD** contain zero or one [0..1] **reference/@value** (CONF:7659).
 1. This reference/@value **SHALL** begin with a '#' and **SHALL** point to its corresponding narrative (using the approach defined in CDA Release 2, section 4.3.5.1) (CONF:7660).
- 2320 6. **SHALL** contain exactly one [1..1] **statusCode**, where the @code **SHALL** be selected from ValueSet ProcedureAct statusCode 2.16.840.1.113883.11.20.9.22 **DYNAMIC** (CONF:7661).
 7. **SHOULD** contain zero or one [0..1] **effectiveTime** (CONF:7662).
 8. **MAY** contain zero or one [0..1] **priorityCode**, where the @code **SHALL** be selected from ValueSet ActPriority 2.16.840.1.113883.1.11.16866 **DYNAMIC** (CONF:7668).
- 2325 9. **MAY** contain zero or one [0..1] **methodCode** (CONF:7670).
 a. methodCode **SHALL NOT** conflict with the method inherent in Procedure / code (CONF:7890).
- 2330 10. **SHALL** contain at least one [1..*] **targetSiteCode** (CONF:7683-CRC).
 a. The targetSiteCode, if present, **SHALL** contain exactly one [1..1] **code**, where the @code **SHALL** be selected from ValueSet Body Site 1.3.6.1.4.1.19376.1.4.1.5.32 **STATIC** (CONF:10122-CRC).
 b. This code **SHOULD** contain zero or one [0..1] **originalText** (CONF:CRC-xxx).
 i. The originalText, if present, **SHOULD** contain zero or one [0..1] **reference/@value** (CONF:CRC-xxx).
 1. This reference/@value **SHALL** begin with a '#' and **SHALL** point to its corresponding narrative (using the approach defined in CDA Release 2, section 4.3.5.1) (CONF:CRC-xxx).
 2. This text is used to describe the native or graft structures in the patient's coronary anatomy.
- 2340 11. **MAY** contain zero or more [0..*] **specimen** (CONF:7697).
 a. This specimen is for representing specimens obtained from a procedure (CONF:8008).
 b. The specimen, if present, **SHALL** contain exactly one [1..1] **specimenRole** (CONF:7704).
 i. This specimenRole **SHOULD** contain zero or more [0..*] **id** (CONF:7716).
- 2345

- 2350 1. If you want to indicate that the Procedure and the Results are referring to the same specimen, the Procedure/specimen/specimenRole/id **SHOULD** be set to equal an Organizer/specimen/ specimenRole/id (CONF:7717).
- 2355 12. **SHOULD** contain zero or more [0..*] **performer** (CONF:7718) such that it
- a. **SHALL** contain exactly one [1..1] **assignedEntity** (CONF:7720).
 - i. This assignedEntity **SHALL** contain at least one [1..*] **id** (CONF:7722).
 - ii. This assignedEntity **SHALL** contain exactly one [1..1] **addr** (CONF:7731).
 - iii. This assignedEntity **SHALL** contain exactly one [1..1] **telecom** (CONF:7732).
 - iv. This assignedEntity **SHOULD** contain zero or one [0..1] **representedOrganization** (CONF:7733).
 - 1. The representedOrganization, if present, **SHOULD** contain zero or more [0..*] **id** (CONF:7734).
 - 2. The representedOrganization, if present, **MAY** contain zero or more [0..*] **name** (CONF:7735).
 - 3. The representedOrganization, if present, **SHALL** contain exactly one [1..1] **addr** (CONF:7736).
 - 4. The representedOrganization, if present, **SHALL** contain exactly one [1..1] **telecom** (CONF:7737).
- 2360
- 2365 13. **MAY** contain zero or more [0..*] **participant** (CONF:7751) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="DEV"** Device (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:7752).
 - b. **SHALL** contain exactly one [1..1] **Product Instance** (2.16.840.1.113883.10.20.22.4.37) (CONF:7754).
- 2370
- 2375 14. **MAY** contain zero or more [0..*] **participant** (CONF:7765) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="LOC"** Location (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90) (CONF:7766).
 - b. **SHALL** contain exactly one [1..1] **Service Delivery Location** (2.16.840.1.113883.10.20.22.4.32) (CONF:7767).
- 2380
- 2385 15. **MAY** contain zero or more [0..*] **entryRelationship** (CONF:7768) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="COMP"** Has Component (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:7769).
 - b. **SHALL** contain exactly one [1..1] **@inversionInd="true"** true (CONF:8009).
 - c. **SHALL** contain exactly one [1..1] **encounter** (CONF:7770).
 - i. This encounter **SHALL** contain exactly one [1..1] **@classCode="ENC"** Encounter (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:7771).
 - ii. This encounter **SHALL** contain exactly one [1..1] **@moodCode="EVN"** Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:7772).
- 2390

- iii. This encounter **SHALL** contain exactly one [1..1] **id** (CONF:7773).
1. Set the encounter ID to the ID of an encounter in another section to signify they are the same encounter (CONF:7774).
- 2395 16. **MAY** contain zero or one [0..1] **entryRelationship** (CONF:7775) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="SUBJ"** Has Subject (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:7776).
 - b. **SHALL** contain exactly one [1..1] **@inversionInd="true"** true (CONF:7777).
 - 2400 c. **SHALL** contain exactly one [1..1] **Instructions** (2.16.840.1.113883.10.20.22.4.20) (CONF:7778).
17. **MAY** contain zero or more [0..*] **entryRelationship** (CONF:7779) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="RSON"** Has Reason (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:7780).
 - 2405 b. **SHALL** contain exactly one [1..1] **Indication** (2.16.840.1.113883.10.20.22.4.19) (CONF:7781).
18. **MAY** contain zero or one [0..1] **entryRelationship** (CONF:7886) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="COMP"** Has Component (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:7887).
 - 2410 b. **SHALL** contain exactly one [1..1] **Medication Activity** (2.16.840.1.113883.10.20.22.4.16) (CONF:7888).
19. **MAY** contain zero or more [0..*] **entry** (CONF:CRC-xxx) such that it
- 2415 a. **SHALL** contain exactly one [1..1] **Procedure Device Organizer - Cardiac** (1.3.6.1.4.1.19376.1.4.1.4.12) (CONF:CRC-xxx).
20. **MAY** contain zero or one [0..1] **entryRelationship** (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] **@typeCode="REFR"** References (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:CRC-xxx).
 - 2420 b. **SHALL** contain exactly one [1..1] **Lesion Observation** (1.3.6.1.4.1.19376.1.4.1.10) (CONF:CRC-xxx). This refers to the lesion that this procedure is related to.

6.3.4.13.2 Procedure Device Organizer - Cardiac

2425 [organizer: templateId 1.3.6.1.4.1.19376.1.4.1.4.12 (open)]

This Procedure Device Organizer – Cardiac entry content module identifies a set of observations related to a device used during procedures. It is intended to be used to further describe the devices used during these procedures.

1. **SHALL** contain exactly one [1..1] **@classCode** (CONF:CRC-xxx).
- 2430 a. **SHALL** contain exactly one [1..1] **@classCode="CLUSTER"** Cluster (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass) (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass) (CONF:CRC-xxx).

- 2435 2. **SHALL** contain exactly one [1..1] `@moodCode="EVN"` Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:CRC-xxx).
- 2435 3. **SHALL** contain exactly one [1..1] `templateId` (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] `@root="1.3.6.1.4.1.19376.1.4.1.4.12"` (CONF:CRC-xxx).
- 2440 4. **SHALL** contain at least one [1..*] `id` (CONF:CRC-xxx).
- 2440 5. **SHOULD** contain zero or one [0..1] `participant` (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] `@typeCode="SUBJ"` (CodeSystem: HL7ParticipationType 2.16.840.1.113883.5.90) (CONF:CRC-xxx).
- b. **SHALL** contain exactly one [1..1] `participantRole` (CONF:CRC-xxx).
- i. This `participantRole` **SHALL** contain exactly one [1..1] `@classCode="MANU"` Manufactured Product (CodeSystem: RoleClass 2.16.840.1.113883.5.110) (CONF:CRC-xxx).
- ii. This `participantRole` **SHALL** contain exactly one [1..1] `playingDevice` (CONF:CRC-xxx).
1. This `playingDevice` **SHALL** contain exactly one [1..1] `@classCode="MMAT"` Manufactured Material (CodeSystem: EntityClass 2.16.840.1.113883.5.41) (CONF:CRC-xxx).
2. This `playingDevice` **SHALL** contain exactly one [1..1] `code` (CONF:CRC-xxx).
3. This `playingDevice` **SHALL** contain at least one [1..*] `id` (CONF:CRC-xxx).
- 2450 6. **MAY** contain zero or more [0..*] `component` (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] Device Observation (1.3.6.1.4.1.19376.1.4.1.4.13) (CONF:CRC-xxx).

6.3.4.13.3 Device Observation

2460 `[observation: templateId 1.3.6.1.4.1.19376.1.4.1.4.13 (open)]`

This Device Observation entry represents observations made of devices used during a procedure, such as a cardiac procedure. An example of a device observation would be balloon inflation time.

- 2465 1. **SHALL** contain exactly one [1..1] `@classCode="OBS"` Observation (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:CRC-xxx).
- 2465 2. **SHALL** contain exactly one [1..1] `@moodCode="EVN"` Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:CRC-xxx).
- 2470 3. **SHALL** contain exactly one [1..1] `templateId` (CONF:CRC-xxx) such that it
- a. **SHALL** contain exactly one [1..1] `@root="1.3.6.1.4.1.19376.1.4.1.4.13"` (CONF:CRC-xxx).
4. **SHALL** contain at least one [1..*] `id` (CONF:CRC-xxx).
5. **SHALL** contain exactly one [1..1] `code` (CONF:CRC-xxx).

- a. **SHOULD** be from LOINC (CodeSystem: 2.16.840.1.113883.6.1) or SNOMED CT (CodeSystem: 2.16.840.1.113883.6.96) (CONF: CRC-xxx).
- 2475 6. **SHOULD** contain zero or one [0..1] **text** (CONF: CRC-xxx).
 - a. The text, if present, **SHOULD** contain zero or one [0..1] **reference/@value** (CONF: CRC-xxx).
 - 2480 i. This reference/@value **SHALL** begin with a '#' and **SHALL** point to its corresponding narrative (using the approach defined in CDA Release 2, section 4.3.5.1) (CONF: CRC-xxx).
- 7. **SHALL** contain exactly one [1..1] **statusCode="completed"** Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14) (CONF: CRC-xxx).
- 8. **SHALL** contain exactly one [1..1] **effectiveTime** (CONF: CRC-xxx).
 - a. represents clinically effective time of the measurement, which may be when the measurement was performed (e.g., a BP measurement), or may be when sample was taken (and measured some time afterwards) (CONF: CRC-xxx).
- 2485 9. **SHALL** contain exactly one [1..1] **value** with @xsi:type="ANY" (CONF: CRC-xxx).

6.3.4.14 Procedure Specimens Taken Section 59773-2

2490 [section: templateId 2.16.840.1.113883.10.20.22.2.31 (open)]

This Procedure Specimens Taken section is used exactly as specified in C-CDA - section 4.51. The Procedure Specimens Taken section records the tissues, objects, or samples taken from the patient during the procedure including biopsies, aspiration fluid, or other samples sent for pathological analysis. The narrative may include a description of the specimens.

2495

```

<section>
  <templateId root="2.16.840.1.113883.10.20.22.2.31"/>
  <code code="59773-2"
    codeSystem="2.16.840.1.113883.6.1"
    codeSystemName="LOINC"
    displayName="PROCEDURE SPECIMENS TAKEN"/>
  <title>Procedure Specimens Taken</title>
  <text>Ascending colon polyp</text>
</section>
    
```

2500

2505

Figure 6.3.4.14-1: Procedure specimens taken section example

6.3.4.15 Procedure Disposition Section 59775-7

[section: templateId 2.16.840.1.113883.10.20.18.2.12 (open)]

This Procedure Disposition section is used exactly as specified in C-CDA - section 4.46.

2510

```

<section>
  <templateId root="2.16.840.1.113883.10.20.18.2.12"/>
  <code code="59775-7" codeSystem="2.16.840.1.113883.6.1"
    codeSystemName="LOINC"
    displayName="PROCEDURE DISPOSITION"/>
  <title>PROCEDURE DISPOSITION</title>
  <text>The patient was taken to the ICU Recovery Unit in stable
    condition.</text>
</section>

```

2515

Figure 6.3.4.15-1: Procedure disposition section example

2520

6.3.4.16 Procedure Results - Cardiac Section 30954-2

```

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.20 (open)]
  ([section: templateId 2.16.840.1.113883.10.20.22.2.3.1 (open)] – parent)

```

2525

This Procedure Results – Cardiac section content module records clinically significant results confirmed or discovered during the procedure. Results include findings, measurements, calculations, and observations.

2530

For this CRC profile, this Procedure Results – Cardiac section content module should be organized using Procedure Results Organizer – Cardiac entry content modules for specific categories (e.g., right heart cath findings, coronary anatomy findings, left heart cath findings, and PCI findings). There shall be a Procedure Result Organizer – Cardiac entry content module for one or more of these categories of findings. The allowed categories are defined in 1.3.6.1.4.1.19376.1.4.1.5.43 CRC Procedure Findings Types which can be expanded to include other procedures.

2535

Result Observation – Cardiac entries are used to record specific findings (e.g., stenosis, timi flow, lesion characteristics, or wall motion characteristics) in each category. The specific findings should be selected from the result observations Value Set 1.3.6.1.4.1.19376.1.4.1.5.38, Procedure Findings Constraints / Value Set. Note that these findings may apply to lesions and coronary anatomy.

2540

This Procedure Results – Cardiac section content module is a modification of the C-CDA Results Section with Coded Entries Required (C-CDA 4.55). The modifications are highlighted in yellow below. This Procedure Results – Cardiac section content module is also conformant to the C-CDA Results Section content module.

2545

1. **SHALL** contain three or more [3..*] `templateId` (CONF:7108-CRC) such that it
 - a. **SHALL** contain exactly one [1..1]
 - `@root="2.16.840.1.113883.10.20.22.2.3"` (CONF:9136).
 - b. **SHALL** contain exactly one [1..1]
 - `@root="2.16.840.1.113883.10.20.22.2.3.1"` (CONF:9137).

2550

c. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.20" (CONF:CRC-xxx).

2. **SHALL** contain exactly one [1..1] **code/@code**="30954-2" Relevant diagnostic tests and/or laboratory data (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:7110).

3. **SHALL** contain exactly one [1..1] **title** (CONF:8892).

2555

4. **SHALL** contain exactly one [1..1] **text** (CONF:7111).

5. **SHALL** contain at least one [1..*] **entry** (CONF:7112-CRC) such that it

a. **SHALL** contain exactly one [1..1] Procedure Results Organizer - Cardiac (1.3.6.1.4.1.19376.1.5.3.1.4.15) (CONF:7113-CRC).

2560 <section>
 <templateId root="21.3.6.1.4.1.19376.1.4.1.2.20" />
 <templateId root="2.16.840.1.113883.10.20.22.2.3.1" />
 <templateId root="2.16.840.1.113883.10.20.22.2.3" />
 <code code="30954-2"
 2565 codeSystem="2.16.840.1.113883.6.1"
 codeSystemName="LOINC"
 displayName="RESULTS" />
 <title>Procedure results</title>
 <text>
 2570 Left Main: No significant narrowing noted. Proximal LAD: No significant
 narrowing Noted. Mid/Distal LAD, Diag Branches: No significant
 narrowing noted. RCA, RPDA, RPL, AM Branches: The distal RCA has a
 stenosis of 90 percent. Circ., OMs, LPDA, LPL Branches: The proximal
 2575 Left Circumflex has a stenosis of 80 percent. Ramus: No Significant
 narrowing noted.
 <content ID="observation1">Post procedure stenosis of the Distal RCA is
 0%.</content>
 <content ID="severity3">Moderate to severe</content>
 </text>
 2580 <entry>
 <organizer classCode="CLUSTER" moodCode="EVN">
 <templateId root="2.16.840.1.113883.10.20.22.4.1"/>
 <!-- Procedure Results Organizer - Cardiac -->
 <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.15"/>
 2585 <id root="7d5a02b0-67a4-11db-bd13-0800200c9a66"/>
 <code code="500786010" displayName="Left Heart Cath Procedure"
 codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"/>
 <statusCode code="completed"/>
 <component>
 2590 <observation classCode="OBS" moodCode="EVN">
 <!-- Result observation - cardiac template -->
 <templateId root="1.3.6.1.4.1.19376.1.4.1.4.16"/>
 <templateId root="2.16.840.1.113883.10.20.22.4.2"/>
 <id root="c6f88321-67ad-11db-bd13-0800200c9a66"/>
 2595 <code code="233970002" codeSystem="2.16.840.1.113883.6.96"
 codeSystemName="SNOMED CT"
 displayName="Post procedure stenosis"/>
 <text><reference value="observation1"/></text>
 <statusCode code="completed"/>
 <effectiveTime value="19991114"/>
 <targetSiteCode code="41879009"
 2600 codeSystem="2.16.840.1.113883.6.96"
 displayName="Distal RCA"/>

2605

```

<value xsi:type="PQ" value="0" unit="%" />
<interpretationCode code="N" codeSystem="2.16.840.1.113883.5.83" />
<entryRelationship typeCode="SUBJ" inversionInd="TRUE">
  <observation classCode="OBS" moodCode="EVN">
    <!-- Severity observation template -->
    <templateId root=" 2.16.840.1.113883.10.20.22.4.8" />
    <id root="c6f88321-67ad-11db-bd13-0800200c9a66" ext="Lesion1" />
    <code code="SEV" displayName="Severity Observation"
      codeSystem="2.16.840.1.113883.5.4"
      codeSystemName="ActCode" />
    <text><reference value="#severity3"/></text>
    <statusCode code="completed" />
    <value xsi:type="CD" code="371924009"
      displayName="Moderate to severe"
      codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT" />
  </observation>
</entryRelationship>
</observation>
</component>
<component>

```

2610

2615

2620

2625

2630

2635

2640

```

  <observation classCode="OBS" moodCode="EVN">
    <!-- Result observation - cardiac template -->
    <templateId root="1.3.6.1.4.1.19376.1.4.1.4.16" />
    ...
  </observation>
</component>
<entryRelationship typeCode="REFR">
  <observation classCode="OBS" moodCode="EVN">
    <!-- Lesion observation template -->
    <templateId root="1.3.6.1.4.1.19376.1.4.1.4.10" />
    <id root="2.840.110893.98120.74.8" ext="lesion #1" />
    <code code="404684003" displayName="Finding"
      codeSystem="2.16.840.1.113883.6.96"
      codeSystemName="SNOMED CT" />
  </observation>
</entryRelationship>
</organizer>
</entry>
</section>

```

Figure 6.3.4.16-1: Results section example

2645

6.3.4.16.1 Procedure Results Organizer - Cardiac

[organizer: templateId 1.3.6.1.4.1.19376.1.5.3.1.4.15 (open)]

([observation: templateId 2.16.840.1.113883.10.20.22.4.1 (open)] – parent)

2650

This Procedure Results Organizer – Cardiac entry content module identifies a set of related procedure results, findings and observations. It contains information applicable to all of the contained procedure findings, including the lesion for PCI procedures. Related

procedure findings type codes categorize a finding into one of several commonly accepted values (e.g., “Right heart cath”, “Left heart cath”, “PCI”).

2655 This Procedure Results Organizer – Cardiac entry content module is a modification of the C-CDA Result Organizer Section (C-CDA 5.57). The modifications are highlighted in yellow below. This Procedure Results Organizer – Cardiac entry content module is also conformant to the C-CDA Results Organizer entry content module.

- 2660 1. **SHALL** contain exactly one [1..1] `@classCode` (CONF:7121).
 - 2660 a. **SHALL** contain exactly one [1..1] `@classCode="CLUSTER"` Cluster (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass) (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass) (CONF: 7165-xxx).
2. **SHALL** contain exactly one [1..1] `@moodCode="EVN"` Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:7122).
- 2665 3. **SHALL** contain two or more [2..*] `templateId` (CONF:7126-CRC) such that it
 - 2665 a. **SHALL** contain exactly one [1..1] `@root="2.16.840.1.113883.10.20.22.4.1"` (CONF:9134).
 - b. **SHALL** contain exactly one [1..1] `@root="1.3.6.1.4.1.19376.1.5.3.1.4.15"` (CONF:CRC-xxx).
- 2670 4. **SHALL** contain at least one [1..*] `id` (CONF:7127).
- 2670 5. **SHALL** contain exactly one [1..1] `code` (CONF:7128).
 - 2675 a. **SHOULD** be selected from CRC procedure findings types found in 1.3.6.1.4.1.19376.1.4.1.5.43 CRC Procedure Findings Types or **MAY** be selected from LOINC (codeSystem 2.16.840.1.113883.6.1) or SNOMED CT (codeSystem 2.16.840.1.113883.6.96), or CPT-4 (codeSystem 2.16.840.1.113883.6.12) (CONF:7164-CRC).
- 2675 6. **SHALL** contain exactly one [1..1] `statusCode="completed"` Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14) (CONF:7123).
- 2680 7. **SHALL** contain at least one [1..*] `component` (CONF:7124) such that it
 - 2680 a. **SHALL** contain exactly one [1..1] [Result Observation - Cardiac](#) (1.3.6.1.4.1.19376.1.4.1.4.16) (CONF:7125-CRC).
- 2685 8. **MAY** contain zero or one [0..1] `entryRelationship` (CONF:CRC-xxx) such that it
 - 2685 a. **SHALL** contain exactly one [1..1] `@typeCode="REFR"` References (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:CRC-xxx).
 - b. **SHALL** contain exactly one [1..1] [Lesion Observation](#) (1.3.6.1.4.1.19376.1.4.1.10) (CONF:CRC-xxx). This refers to the lesion that these results are related to.

6.3.4.16.2 Result Observation - Cardiac

2690 [observation: templateId 1.3.6.1.4.1.19376.1.4.1.4.16 (open)]
 ([observation: templateId 2.16.840.1.113883.10.20.22.4.2 (open)] – parent)

A result observation is a clinical statement that a clinician has noted during the Cath Lab procedure. This Result Observation – Cardiac entry content module is used to describe the specific procedure findings that were observed during the specific Cath Lab procedure.

2695 The specific result observations are defined in 1.3.6.1.4.1.19376.1.4.1.5.38 Procedure Findings Constraints.

The targetSiteCode may be used for diagnostic cath procedures.

2700 This Result Observation – Cardiac entry content module is a modification of the C-CDA Result Observation (C-CDA 5.56). The modifications are highlighted in yellow below. This Result Observation – Cardiac entry content module is also conformant to the C-CDA Result Observation entry content module.

1. **SHALL** contain exactly one [1..1] @classCode="OBS" Observation (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:7130).
- 2705 2. **SHALL** contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:7131).
3. **SHALL** contain two or more [2..*] templateId (CONF:7136-CRC) such that it
 - 2710 a. **SHALL** contain exactly one [1..1] @root="2.16.840.1.113883.10.20.22.4.2" (CONF:9138).
 - b. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.4.1.4.16" (CONF:CRC-xxx).
4. **SHALL** contain at least one [1..*] id (CONF:7137).
5. **SHALL** contain exactly one [1..1] code (CONF:7133).
 - 2715 a. **SHOULD** be from LOINC (CodeSystem: 2.16.840.1.113883.6.1) or SNOMED CT or Value Set Procedure Findings (1.3.6.1.4.1.19376.1.4.1.5.38) (CONF:7166-CRC).
6. **SHOULD** contain zero or one [0..1] text (CONF:7138).
 - a. The text, if present, **SHOULD** contain zero or one [0..1] reference/@value (CONF:7139).
 - 2720 i. This reference/@value **SHALL** begin with a '#' and **SHALL** point to its corresponding narrative (using the approach defined in CDA Release 2, section 4.3.5.1) (CONF:9119).
7. **SHALL** contain exactly one [1..1] statusCode="completed" Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14) (CONF:7134).
- 2725 8. **SHALL** contain exactly one [1..1] effectiveTime (CONF:7140).
 - a. represents clinically effective time of the measurement, which may be when the measurement was performed (e.g., a BP measurement), or may be when sample was taken (and measured some time afterwards) (CONF:7141).
9. **SHALL** contain exactly one [1..1] value with @xsi:type="ANY" (CONF:7143).

- 2730 10. **SHOULD** contain zero or more [0..*] **interpretationCode** (CONF:7147).
11. **MAY** contain zero or one [0..1] **methodCode** (CONF:7148).
12. **MAY** contain zero or one [0..1] **targetSiteCode** (CONF:7153).
a. The **targetSiteCode**, if present, **SHALL** contain exactly one [1..1] **code** where the **@code** **SHALL** be selected from ValueSet **Body Site 1.3.6.1.4.1.19376.1.4.1.5.32** **STATIC** (CONF:CRC-xxx).
2735 13. **MAY** contain zero or one [0..1] **author** (CONF:7149).
14. **SHOULD** contain zero or more [0..*] **referenceRange** (CONF:7150).
a. The **referenceRange**, if present, **SHALL** contain exactly one [1..1] **observationRange** (CONF:7151).
2740 i. This **observationRange** **SHALL NOT** contain [0..0] **code** (CONF:7152).
15. **SHOULD** contain zero or one [0..1] **entryRelationship** (CONF:CRC-xxx) such that it
a. **SHALL** contain exactly one [1..1] **@typeCode="SUBJ"** Has subject (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:CRC-xxx).
2745 b. **SHALL** contain exactly one [1..1] **@inversionInd="true"** TRUE (CONF:CRC-xxx).
c. **SHALL** contain exactly one [1..1] **Severity Observation** (2.16.840.1.113883.10.20.22.4.8) (CONF:CRC-xxx).

2750 **6.3.4.17 Complications Section 55109-3**

[section: templateId 2.16.840.1.113883.10.20.22.2.37 (open)]

This Complications section content module records problems that occurred during the cath lab procedure. The complications may have been known risks or unanticipated problems.

2755 This Complications section content module is used exactly as specified in C-CDA - section 4.8, except for vocabulary constraints for Problem Observation entries.

There is a CRC specific value set defined for complications recorded as Problem Observation entries in this Complications section content module.

```

2760 <section>
      <templateId root="1.3.6.1.4.1.19376.1.4.1.2.25"/>
      <code code="55109-3" codeSystem="2.16.840.1.113883.6.1"
            codeSystemName="LOINC"
            displayName="Complications"/>
      <title>Complications</title>
2765 <text>Complications for the cath procedure for patient included:
            x, y, z...
      </text>
      <entry>
2770 <observation classCode="OBS" moodCode="EVN">
            <templateId root="1.3.6.1.4.1.19376.1.4.1.4.9"/>
            <id root="d11275e7-67ae-11db-bd13-0800200c9a66"/>
            <code code="404684003" codeSystem="2.16.840.1.113883.6.96"
                  displayName="Post Procedure Diagnosis"/>
            <text>The patient has had a myocardial infarction..</text>
2775 <statusCode code="completed"/>
            <effectiveTime>
                <low value="201201251000"/>
            </effectiveTime>
            <value xsi:type="CD" code="22298006"
                  codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"
                  displayName="Myocardial Infarction (Biomarker Positive)"/>
            <entryRelationship typeCode="REFR">
2780 <observation classCode="OBS" moodCode="EVN">
                <templateId root="2.16.840.1.113883.10.20.22.4.6"/>
                <!-- Problem Status template -->
                <code code="33999-4" codeSystem="2.16.840.1.113883.6.1"
                      codeSystemName="LOINC" displayName="Status"/>
                <statusCode code="completed"/>
                <value xsi:type="CD" code="55561003"
                      codeSystem="2.16.840.1.113883.6.96"
                      codeSystemName="SNOMED CT" displayName="Active"/>
            </observation>
            </entryRelationship>
2785 </observation>
        </entry>
        <entry>
2790 <observation classCode="OBS" moodCode="EVN">
            <templateId root="1.3.6.1.4.1.19376.1.4.1.4.9"/>
            <id root="xyz"/>
            ...
2800 </observation>
        </entry>
    </section>

```

Figure 6.3.4.17-1: Complications section example

2805

6.3.4.17.1 Problem Observation – Constraints

[observation: templateId 2.16.840.1.113883.10.20.22.4.4 (open)]

A problem is a clinical statement that a clinician has noted during the Cath procedure. This entry is used to describe the presence or absence of specific “complications” as defined by ACC.

2810 This Problem Observation entry content module is used exactly as specified in C-CDA - section 5.45, except for vocabulary constraints.

The value set for CONF:9058 (**value@code**) **SHOULD** be selected from ValueSet Complications (1.3.6.1.4.1.19376.1.4.1.5.46) **STATIC**.

2815 **6.3.4.18 Postprocedure Diagnosis Section 59769-0**

[section: templateId 2.16.840.1.113883.10.20.22.2.36 (open)]

The Postprocedure Diagnosis section content module records the diagnosis or diagnoses discovered or confirmed during the procedure. Often it is the same as the pre-procedure diagnosis or indication.

2820 This Postprocedure Diagnosis section content module is used exactly as specified in C-CDA - section 4.42, except for vocabulary constraints.

There is a CRC specific value set defined for problem observations recorded as part of postprocedure diagnosis which is included in the Problem Observation entry.

```

2825 <section>
      <templateId root="2.16.840.1.113883.10.20.22.2.36"/>
      <code code="59769-0" codeSystem="2.16.840.1.113883.6.1"
        codeSystemName="LOINC" displayName="POSTPROCEDURE DIAGNOSIS"/>
2830 <title>Postprocedure Diagnosis</title>
      <text>It was observed that there was complication of myocardial
        infarction during the cath procedure.</text>
      <entry>
        <act moodCode="EVN" classCode="ACT">
2835 <templateId root="2.16.840.1.113883.10.20.22.4.51"/>
        <!-- ** Postprocedure Diagnosis Entry ** -->
        <code code="59769-0" codeSystem="2.16.840.1.113883.6.1"
          codeSystemName="LOINC"
          displayName="Postprocedure Diagnosis"/>
2840 <entryRelationship typeCode="SUBJ">
          <observation classCode="OBS" moodCode="EVN">
            <templateId root="2.16.840.1.113883.10.20.22.4.4"/>
            <!-- Problem Observation template -->
            <id root="d11275e7-67ae-11db-bd13-0800200c9a66"/>
2845 <code code="404684003" codeSystem="2.16.840.1.113883.6.96"
              codeSystemName="SNOMED CT"
              displayName="Finding"/>
            <text>It was observed that there was complication of myocardial
              infarction during the cath procedure.</text>
            <statusCode code="completed"/>
            <effectiveTime>
2850 <low value="201201251000"/>
            </effectiveTime>
            <value xsi:type="CD" code="22298006"
              codeSystem="2.16.840.1.113883.6.96"
              codeSystemName="SNOMED CT"
              displayName="Myocardial Infarction (Biomarker
2855 Positive)"/>
          <entryRelationship typeCode="REFR">
            <observation classCode="OBS" moodCode="EVN">
2860 <templateId root="2.16.840.1.113883.10.20.22.4.6"/>
            <!-- Problem Status template -->
            <code code="33999-4" codeSystem="2.16.840.1.113883.6.1"
              codeSystemName="LOINC" displayName="Status"/>
            <statusCode code="completed"/>
            <value xsi:type="CD" code="55561003"
              codeSystem="2.16.840.1.113883.6.96"
              codeSystemName="SNOMED CT" displayName="Active"/>
            </observation>
          </entryRelationship>
        </observation>
      </entryRelationship>
      ...
      </act>
      </entry>
2875 </section>

```

Figure 6.3.4.18-1: Postprocedure diagnosis section example

6.3.4.18.1 Problem Observation – Constraints

[observation: templateId 2.16.840.1.113883.10.20.22.4.4 (open)]

2880 The Problem Observation entry is used to describe a final diagnosis.

This Problem Observation entry is used exactly as specified in C-CDA - section 5.45, except for vocabulary constraints.

The value set for CONF:9058 (**value**) **SHOULD** be selected from ValueSet CRC Postprocedure Diagnosis (1.3.6.1.4.1.19376.1.4.1.5.44) **STATIC**.

2885 6.3.4.19 Plan of Care - Cardiac Section 18776-5

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.22 (open)]

[(section: templateId 2.16.840.1.113883.10.20.22.2.10 (open)) – parent]

This Plan of Care - Cardiac section content module is intended to be used to describe the post-procedure plan.

2890 The Plan of Care - Cardiac section content module contains data that defines pending orders, interventions, encounters, services, and procedures for the patient. It is limited to prospective, unfulfilled, or incomplete orders and requests only, which are indicated by the @moodCode of the entries within this section. All active, incomplete, or pending orders, appointments, referrals, procedures, services, or any other pending event of clinical significance to the current care of the patient should be listed unless constrained due to privacy issues. The plan may also contain information about ongoing care of the patient and information regarding goals and clinical reminders. Clinical reminders are placed here to provide prompts for disease prevention and management, patient safety, and health-care quality improvements, including widely accepted performance measures. The plan may also indicate that patient education was given or will be provided.

2895

2900

This Plan of Care – Cardiac section content module is a modification of the C-CDA Plan of Care section (C-CDA 4.39). The modifications are highlighted in yellow below. This Plan of Care – Cardiac section content module is also conformant to the C-CDA Plan of Care section content module.

- 2905
1. **SHALL** contain two or more [2..*] **templateId** (CONF:7723-CRC) such that it
 - a. **SHALL** contain exactly one [1..1]

@root="2.16.840.1.113883.10.20.22.2.10" (CONF:10435).
 - b. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.4.1.2.22" (CONF:CRC-xxx).
 2. **SHALL** contain exactly one [1..1] **code/@code**="18776-5" Plan of Care (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:7724).
 3. **SHALL** contain exactly one [1..1] **text** (CONF:7725).
 4. **MAY** contain zero or more [0..*] **entry** (CONF:7726) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Act - Cardiac (1.3.6.1.4.1.19376.1.4.1.4.17) (CONF:8804-CRC) .
- 2910
- 2915

- 2920
5. **MAY** contain zero or more [0..*] **entry** (CONF:8805) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Encounter (2.16.840.1.113883.10.20.22.4.40) (CONF:8806).
 6. **MAY** contain zero or more [0..*] **entry** (CONF:8807) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Observation (2.16.840.1.113883.10.20.22.4.44) (CONF:8808).
 7. **MAY** contain zero or more [0..*] **entry** (CONF:8809) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Procedure (2.16.840.1.113883.10.20.22.4.41) (CONF:8810).
 - 2925 8. **MAY** contain zero or more [0..*] **entry** (CONF:8811) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Substance Administration (2.16.840.1.113883.10.20.22.4.42) (CONF:8812).
 - 2930 9. **MAY** contain zero or more [0..*] **entry** (CONF:8813) such that it
 - a. **SHALL** contain exactly one [1..1] Plan of Care Activity Supply (2.16.840.1.113883.10.20.22.4.43) (CONF:8814).

```

2935 <section>
      <templateId root="1.3.6.1.4.1.19376.1.4.1.2.22" />
      <templateId root="2.16.840.1.113883.10.20.22.2.10" />
      <!-- **** Plan of Care - Cardiac section template **** -->
      <code code="18776-5" codeSystem="2.16.840.1.113883.6.1"
      codeSystemName="LOINC" displayName="Treatment plan"/>
      <title>Plan of Care</title>
      <text>
2940         ...
      </text>
      <entry>
          <act moodCode="RQO" classCode="ACT">
2945             <!-- **** Plan of Care Activity Act - Cardiac template **** -->
             <templateId root="1.3.6.1.4.1.19376.1.4.1.4.17"/>
             <templateId root="2.16.840.1.113883.10.20.22.4.39"/>
             <id root="9a6dlbac-17d3-4195-89a4-1121bc809a5c"/>
             <code code="415070008" codeSystem="2.16.840.1.113883.6.96"
2950                 displayName="PCI without planned CABG"/>
             <statusCode code="new"/>
             <effectiveTime>
                 <center value="20000421"/>
             </effectiveTime>
             </act>
2955 </entry>
      </section>
  
```

Figure 6.3.4.19-1: Plan of care section example

2960

6.3.4.19.1 Plan of Care Activity Act - Cardiac

[act: templateId 1.3.6.1.4.1.19376.1.4.1.4.17 (open)]

([act: templateId 2.16.840.1.113883.10.20.22.4.39 (open)] – parent)

2965 This Plan of Care Activity Act – Cardiac entry content module is a modification of the C-CDA Plan of Care Activity Act (C-CDA 5.33). The modifications are highlighted in yellow below. This Plan of Care Activity Act – Cardiac entry content module is also conformant to the C-CDA Plan of Care Activity Act entry content module.

- 2970
1. **SHALL** contain exactly one [1..1] **@classCode**="ACT" (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:8538).
 2. **SHALL** contain exactly one [1..1] **@moodCode**, which **SHALL** be selected from ValueSet Plan of Care moodCode (Act/Encounter/Procedure) 2.16.840.1.113883.11.20.9.23 **STATIC** 2011-09-30 (CONF:8539).
 - 2975 3. **SHALL** contain two or more [2..*] **templateId** (CONF:8544-CRC) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="2.16.840.1.113883.10.20.22.4.39" (CONF:10510).
 - b. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.4.17" (CONF:CRC-xxx).
 4. **SHALL** contain at least one [1..*] **id** (CONF:8546).
 - 2980 5. **SHALL** contain exactly one [1..1] **code**, where the **@code** **SHOULD** be selected from ValueSet Rx Recommendation (1.3.6.1.4.1.19376.1.4.1.5.42) **STATIC** (CONF:CRC-XXX).
 6. **SHALL** contain exactly one [1..1] **statusCode**="completed" Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14) (CONF:CRC-xxx).
 - 2985 7. **SHOULD** contain zero or one [0..1] **effectiveTime** (CONF:CRC-xxx).
 - a. The onset date **SHALL** be recorded in the low element of the effectiveTime element when known (CONF:CRC-xxx).
 - b. The resolution date **SHALL** be recorded in the high element of the effectiveTime element when known (CONF:CRC-xxx).
 - 2990 c. If the problem is known to be resolved, but the date of resolution is not known, then the high element **SHALL** be present, and the nullFlavor attribute **SHALL** be set to 'UNK'. Therefore, the existence of a high element within a problem does indicate that the problem has been resolved (CONF:CRC-xxx).

2995 6.3.4.20 Key Images – Cardiac Section – DCM 121180

[section: templateId 1.3.6.1.4.1.19376.1.4.1.2.21 (open)]

The Key Images section content module contains narrative description of and references to DICOM Image Information Objects that illustrate the findings of the procedure reported.

- 3000
1. **SHALL** contain exactly one [1..1] **templateId** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.2.21" (CONF:CRC-xxx).
 2. **SHALL** contain exactly one [1..1] **code/@code**="121180" Key Images (CodeSystem: 1.2.840.10008.2.16.4 DCM) (CONF:CRC-xxx).
 3. **SHALL** contain exactly one [1..1] **text** (CONF:CRC-xxx).
 - 3005 4. **SHALL** contain at least one [1..*] **entry** (CONF:CRC-xxx)
 - a. **SHALL** contain exactly one [1..1] Sop Instance Observation (2.16.840.1.113883.10.20.6.2.8) (CONF:CRC-xxx).

6.3.5 Common Entry Content Modules

3010 6.3.5.1 Problem Observation – Cardiac

[Observation: templateId 1.3.6.1.4.1.19376.1.4.1.9(open)]
 ([Observation: templateId 2.16.840.1.113883.10.20.22.4.4(open)] -
 parent)

3015 A problem is a clinical statement that a clinician has noted. In health care it is a condition that requires monitoring or diagnostic, therapeutic, or educational action. It also refers to any unmet or partially met basic human need. In cardiology, problems include hypertension, diabetes, and dyslipidemia.

This Problem Observation – Cardiac entry content module extends the C-CDA Problem Observation entry definition (C-CDA 5.45) by adding the following constraints:

- 3020
1. **SHALL** contain exactly one [1..1] **templateId** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.9" (CONF:CRC-xxx).
 - 3025 2. **MAY** contain zero or one [0..1] **entryRelationship** (CONF:CRC-xxx) such that it
 - a. **SHALL** contain exactly one [1..1] **@typeCode**="SUBJ" Has subject (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002) (CONF:CRC-xxx).
 - b. **SHALL** contain exactly one [1..1] **@inversionInd**="true" TRUE (CONF:CRC-xxx).
 - 3030 c. **SHALL** contain exactly one [1..1] Severity Observation (2.16.840.1.113883.10.20.22.4.8) (CONF:CRC-xxx).

6.3.5.2 Lesion Observation

[Observation: templateId 1.3.6.1.4.1.19376.1.4.1.10(open)]

This Lesion Observation entry content module identifies a lesion of interest for a PCI procedure. The lesion is identified by a global ID in the **id** element and one or more target sites.

3035

1. **SHALL** contain exactly one [1..1] **@classCode**="OBS" Observation (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6) (CONF:CRC-xxx).
2. **SHALL** contain exactly one [1..1] **@moodCode**="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001) (CONF:CRC-xxx).

3040

3. **SHALL** contain exactly one [1..1] **templateId** (CONF:7299) such that it
 - a. **SHALL** contain exactly one [1..1] **@root**="1.3.6.1.4.1.19376.1.4.1.10" (CONF:CRC-xxx).
4. **SHALL** contain at least one [1..*] **id** where the **@root** **SHALL** be a globally unique root and the **@ext** **SHALL** be a text string representing the lesion ID (CONF:CRC-xxx).

3045

5. **SHALL** contain exactly one [1..1] **code**, where the **@code** **SHOULD** be "404684003" selected from SNOMED CT and has **@displayName**="Finding" (CONF:CRC-xxx).
6. **SHOULD** contain zero or one [0..1] **text** (CONF:CRC-xxx).

3050

- a. The text, if present, **SHOULD** contain zero or one [0..1] **reference/@value** (CONF:CRC-xxx).
 - i. This **reference/@value** **SHALL** begin with a '#' and **SHALL** point to its corresponding narrative (using the approach defined in CDA Release 2, section 4.3.5.1) (CONF:CRC-xxx).

3055

7. **MAY** contain zero or more [0..*] **targetSiteCode**, where the **@code** **SHOULD** be selected from ValueSet Body Site 1.3.6.1.4.1.19376.1.4.1.5.32 **STATIC** (CONF:CRC-xxx).
 - a. The **targetSiteCode**, if present **MAY** contain zero or more [0..*] **qualifier** to further identify the exact location of the lesion (CONF:CRC-xxx).

6.3.6 Cath Report Content Vocabulary Constraints

6.3.6.1 Cardiac problems/concerns - Vocabulary Constraints

3060 The content creator shall be capable of creating a problem/concern selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.31, listed below.

Table 6.3.6.1-1: Cardiac problems/concerns 1.3.6.1.4.1.19376.1.4.1.5.31 STATIC

Concept	Coding Scheme	SNOMED CT
Hypertension		38341003
Dyslipidemia		370992007
Diabetes		73211009
Acute renal failure		14669001
Chronic kidney disease		236425005
Peripheral arterial disease		399957001
Cerebrovascular disease		62914000
Erectile dysfunction		398175007
Cardiac arrhythmia		44808001

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Concept	Coding Scheme	SNOMED CT
Asthma		195967001
Bronchospasm		4386001
Implanted pacemaker		371821000
Heart failure		84114007
Myocardial infarction		22298006
Angina		194828000
Currently on Dialysis		62914000
Chronic Lung Disease		413839001
Diabetes Control - Diet		284071006
Diabetes Control - Oral		170746002
Diabetes Control - Insulin		225302006
CAD Presentation		53741008
Onset of Illness		217315002
Thrombolytic Therapy		426347000
Anginal Class		Pick one from Value Set 1.3.6.1.4.1.19376.1.4.1.5.47
Anti-Anginal Medication: Beta Blockers		33252009
Anti-Anginal Medication: Calcium Channel Blockers		48698004
Anti-Anginal Medication: Long Acting Nitrates		89119000
Anti-Anginal Medication: Ranolazine		420365007
NYHA Class		Pick one from Value Set 1.3.6.1.4.1.19376.1.4.1.5.48
Cardiomyopathy or Left Ventricular Systolic Dysfunction		134401001
Cardiogenic Shock		89138009
Cardiac Arrest		410429000
Prior MI		22298006
Prior Valve Surgery/Procedure		73544002
Prior PCI		415070008
Prior CABG		232717009
Angina Type		Pick one from Value Set 1.3.6.1.4.1.19376.1.4.1.5.49

6.3.6.2 Body Site Value Set - Vocabulary Constraint

3065 The content creator shall be capable of creating a body site selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.32, listed below. This structure is used to represent the native coronary structure of the heart.

Table 6.3.6.2-1: Body Site Value Set 1.3.6.1.4.1.19376.1.4.1.5.32 STATIC

Concept	Coding Scheme	SNOMED CT
Left Main Coronary Artery		3227004
Left Main Coronary Artery Ostium		76862008
Left Anterior Descending Coronary Artery		59438005
Proximal Left Anterior Descending Coronary Artery		68787002
Mid Left Anterior Descending Coronary Artery		91748002
Distal Left Anterior Descending Coronary Artery		36672000
Left Posterior Descending Artery		56322004
Left Posterior Descending Circumflex Coronary Artery		91760001
Left Posterolateral Circumflex Coronary Artery		57823005
Right Coronary Artery		13647002
Right Coronary Artery Ostium		56789007
Proximal Right Coronary Artery		91083009
Mid Right Coronary Artery		13647002+255562008
Distal Right Coronary Artery		41879009
Circumflex Coronary Artery		57396003
Proximal Circumflex Coronary Artery		52433000
Mid Circumflex Coronary Artery		91753007
Distal Circumflex Coronary Artery		6511003
Posterior Descending Right Coronary Artery		53655008
Intermediate Artery (Ramus)		244252004
Right posterior AV Coronary Artery		12800002
1st Diagonal Coronary Artery		91750005
1st Left Posterolateral Coronary Artery		91757008
1st Marginal Coronary Artery		91754001
1st Right posterolateral Coronary Artery		91761002
1st Septal Coronary Artery		244251006
2nd Diagonal Coronary Artery		91751009
2nd Left Posterolateral Coronary Artery		91758003
2nd Marginal Coronary Artery		91755000
2nd Right Posterolateral Coronary Artery		91762009
3rd Diagonal Coronary Artery		91752002
3rd Left Posterolateral Coronary Artery		91759006
3rd Marginal Coronary Artery		91756004

Concept	Coding Scheme	SNOMED CT
3rd Right posterolateral Coronary Artery		91763004
Marginal Right Coronary Artery		22765000
AV groove continuation of Circumflex Artery		75902001

6.3.6.3 Cardiovascular Family History - Vocabulary Constraint

3070 The content creator shall be capable of creating a family history selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.33, listed below.

Table 6.3.6.3-1: Cardiovascular Family History 1.3.6.1.4.1.19376.1.4.1.5.33 STATIC

Concept	Coding Scheme	SNOMED CT
Family history of coronary artery disease		430091005
Family history: Diabetes mellitus		160303001
Family history of myocardial infarction		266897007
No Family history of Diabetes		160274005
No Family history of Cardiovascular disease		160270001
Family History Unknown		407559004

Adapted from DICOM PS3.16-2009

6.3.6.4 Contrast Agents Classes for Adverse Reactions

3075 The content creator shall be capable of creating a Contrast Agents Classes for Adverse Reactions selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.34, listed below.

Table 6.3.6.4-1: Contrast Agents Classes for Adverse Reactions 1.3.6.1.4.1.19376.1.4.1.5.34 STATIC

Concept	Coding Scheme	SNOMED CT
Iodinated contrast agent		426722004
Gadolinium compound		105879004
Echocardiography agent		409290009
Radiopharmaceutical		349358000

6.3.6.5 Cardiac Lab Results - Vocabulary Constraints

3080 The content creator shall be capable of creating cardiac lab results selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.35, listed below.

Table 6.3.6.5-1: Cardiac Lab Results 1.3.6.1.4.1.19376.1.4.1.5.35 DYNAMIC

Concept	Coding Scheme	LOINC	SNOMED
Cholesterol.in HDL		2085-9	
Cholesterol.in LDL		2089-1	
Cholesterol		2093-3	
Triglyceride		2571-8	
High sensitivity C reactive protein		30522-7	
Creatine kinase.MB		13969-1	1224421017
Natriuretic peptide.B		30934-4	
Natriuretic peptide.B prohormone		33762-6	
Troponin T.cardiac		6598-7	186259011
Troponin I.cardiac		10839-9	
Creatinine		2160-0	489161011
Hemoglobin A1c		41995-2	373201015
Urea nitrogen		3094-0	
Fasting glucose		1557-8	
Platelets		11126-0	488930013
Potassium		11148-4	489169013
Urea Nitrogen		11065-0	489160012
Prothrombin Time			2534465010

6.3.6.6 Vital Signs Result Type- Value Set

3085 The content creator shall be capable of creating vital signs organizers selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.36, listed below.

Table 6.3.6.6-1: Vital Sign Result Type 1.3.6.1.4.1.19376.1.4.1.5.36 STATIC

Concept	Coding Scheme	Coding System	Code
Respiratory Rate		LOINC	9279-1
Heart Rate		LOINC	8867-4
O2 % BldC Oximetry		LOINC	2710-2
BP Systolic		LOINC	8480-6
BP Diastolic		LOINC	8462-4
Body Temperature		LOINC	8310-5
Height		LOINC	8302-2
Height (Lying)		LOINC	8306-3
Head Circumference		LOINC	8287-5

Concept	Coding Scheme	Coding System	Code
Weight Measured		LOINC	3141-9
BMI (Body Mass Index)		LOINC	39156-5
BSA (Body Surface Area)		LOINC	3140-1

6.3.6.7 Procedure Indications - Vocabulary Constraints

3090 The content creator shall be capable of creating procedure indications selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.37, listed below.

Table 6.3.6.7-1: Procedure Indications 1.3.6.1.4.1.19376.1.4.1.5.37 STATIC

Concept	Coding Scheme	SNOMED CT
Chest Pain		29857009
Pre-operative		262068006
Coronary Artery Disease		53741008
Heart failure		84114007
Heart disease risk factors		171224000
Dyspnea		267036007
Post PTCA		373108000
History of CABG		399261000
Abnormal exercise tolerance test		165084003
Abnormal ECG		102594003
Arrhythmia		44808001
Angina pectoris		194828000
Hypertension		38341003
Palpitations		80313002
Supraventricular tachycardia		6456007
Syncope		271594007
History of Myocardial Infarction		399211009
Left bundle branch block		63467002
Valvular heart disease		368009
Occupational requirement		429060002
cardiogenic shock		89138009
ischemic heart disease		414545008
cardiac function test abnormal		165076002
heart transplant		32413006
heart disease - congenital		13213009
Cardiomyopathy		85898001
heart disease		56265001
Perioperative Evaluation		430091005

Concept	Coding Scheme	SNOMED CT
structural disorder of heart		128599005
Pericardial disease		55855009

6.3.6.8 Result Observations - Vocabulary Constraints

3095 The content creator shall be capable of creating result observations selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.38, listed below. These results will apply to procedure findings for lesions and the coronary anatomy.

The Procedure Results – Cardiac section content module records clinically significant observations confirmed or discovered during the procedure or surgery.

3100 For this CRC profile, the findings should be organized using Procedure Results Organizer – Cardiac entry content modules for specific categories (e.g. right heart cath findings, coronary anatomy findings, left heart cath findings, and PCI findings). There shall be a Procedure Results Organizer-Cardiac entry content module for one or more of these categories of findings. The allowed categories are defined in 1.3.6.1.4.1.19376.1.4.1.5.43 CRC Procedure Findings Types.

3105 Result Observations – Cardiac entry content modules are used to record specific findings (e.g., stenosis, timi flow, lesion characteristics, or wall motion characteristics) in each category. The specific findings should be selected from the result observations Value Set 1.3.6.1.4.1.19376.1.4.1.5.38, Procedure Findings Constraints / Value Set. Note that these findings may apply to lesions and coronary anatomy.

3110

Table 6.3.6.8-1: Result Observation Constraints 1.3.6.1.4.1.19376.1.4.1.5.38 STATIC

Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
[0..1]	R: PCI	"Previously Treated Lesion"	CD	NA	True or False
[0..1]	R: PCI	449389000, "Previously Treated Lesion with Stent"	CD	NA	True or False
[0..1]	R: PCI	251030009, "In-stent Restenosis"	CD	NA	True or False
[0..1]	R: PCI	421327009, " In-stent Thrombosis"	CD	NA	True or False
[0..1]	R: PCI	408716009, "Stenotic lesion length"	PQ	cm	Value
[0..1]	R: PCI	421327009, " Thrombus Present"	CD	NA	True or False
[0..1]	R: PCI	371894001, " Bifurcation	CD	NA	True or False

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
		Lesion"			
[0..1]	R: PCI & Diagnostic CATH	233970002, pre-procedure Stenosis	PQ	%	
[0..1]	R: PCI	233970002, post-procedure Stenosis	PQ	%	
[0..1]	R: PCI	"Pre-Procedure TIMI Flow"	CD		371867000 (TIMI-0) 371866009 (TIMI-1) 371864007 (TIMI-2) 371865008 (TIMI-3)
[0..1]	R: PCI	"Post-Procedure TIMI Flow"	CD		371867000 (TIMI-0) 371866009 (TIMI-1) 371864007 (TIMI-2) 371865008 (TIMI-3)
[0..1]	R: PCI	70390005," Significant Dissection"	CD	NA	True or False
[0..1]	R: PCI	234010000,"Coronary artery perforation"	CD	NA	True or False
[1..1]	R:Diagnostic Cath	"Coronary Dominance"	CD		253729004 (Left) 253728007 (Right) 253730009 (Balanced)
[0..*]	R:Diagnostic Cath	8583-7, LOINC, "Right atrial A wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8582-9, LOINC, " Left atrial A wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8593-6, LOINC, "Right atrial V wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8592-8, LOINC, "Left atrial V wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8400-4, LOINC, "Right atrial Intrachamber mean pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8399-8, LOINC, "Left atrial Intrachamber mean pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8432-7, LOINC, "Right ventricular Intrachamber systolic pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8430-1, LOINC, " Left ventricular Intrachamber systolic pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8377-4, LOINC, " Right ventricular Intrachamber	PQ	mm[Hg]	

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
		diastolic pressure"			
[0..*]	R:Diagnostic Cath	8375-8, LOINC, "Left ventricular Intrachamber diastolic pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8392-3, LOINC, "Right ventricular End diastolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8391-5, LOINC, "Left ventricular End diastolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8440-0, LOINC, "Pulmonary Artery Systolic Blood Pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8441-8, LOINC, "Pulmonary artery - left Systolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8387-3, LOINC, "Pulmonary artery - right Diastolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8386-5, LOINC, "Pulmonary artery - left Diastolic blood pressure "	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8416-0, LOINC, "Pulmonary artery - right Mean blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8415-2, LOINC, "Pulmonary artery - left Mean blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8584-5, LOINC, "Pulmonary artery wedge A wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8596-9, LOINC, "Pulmonary artery wedge V wave amplitude"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8587-8, LOINC, "Pulmonary artery wedge Mean blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8368-3, LOINC, "Aorta thoracic ascending Diastolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8367-5, LOINC, "Aorta thoracic proximal ascending Diastolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8396-4, LOINC, "Aorta.thoracic ascending Mean blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic	8397-2, LOINC, "Aorta.thoracic proximal	PQ	mm[Hg]	

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
	Cath	ascending Mean blood pressure"			
[0..*]	R:Diagnostic Cath	8423-6, LOINC, "Aorta.thoracic ascending, Systolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8422-8, LOINC, "Aorta.thoracic proximal ascending Systolic blood pressure"	PQ	mm[Hg]	
[0..*]	R:Diagnostic Cath	8840-1, LOINC, " Left atrium Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8841-9, LOINC, "Right atrium Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8842-7, LOINC, " High right atrium Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8843-5, LOINC, " Low right atrium Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8844-3, LOINC, " Mid right atrium Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8845-0, LOINC, " Left ventricular Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8847-6, LOINC, " Right ventricular Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8846-8, LOINC, " Right ventricular outflow tract Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8851-8, LOINC, " Pulmonary artery - left Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8852-6, LOINC, " Main pulmonary artery Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8853-4, LOINC, " Pulmonary artery - right Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8854-2, LOINC, " Pulmonary wedge Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8850-0, LOINC, " Inferior vena cava Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath	8855-9, LOINC, " Superior vena cava Oxygen saturation"	PQ	%	
[0..*]	R:Diagnostic Cath where targetSiteCode = "Ao" or "PA"	14775-1, LOINC, " Hemoglobin [Mass/volume] in Arterial blood by Oximetry"	PQ	g/dL	
[0..*]	R:Diagnostic	50188-2, LOINC, " Arterial-	PQ	vol%	

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
	Cath	venous oxygen saturation difference"			
[0..*]	R:Diagnostic Cath	8741-1, LOINC, "Left ventricular Cardiac output"	PQ	L/min	
[0..*]	R:Diagnostic Cath	8736-1, LOINC, "Left ventricular Cardiac output by Fick method"	PQ	L/min	
[0..*]	R:Diagnostic Cath	8733-8, LOINC, " Left ventricular Cardiac output by Angiography single plane"	PQ	L/min	
[0..*]	R:Diagnostic Cath	8732-0, LOINC, "Left ventricular Cardiac output by Angiography biplane"	PQ	L/min	
[0..*]	R:Diagnostic Cath	8750-2, LOINC, " Left ventricular Cardiac index by Fick method"	PQ	L/min/m2	
[0..*]	R:Diagnostic Cath	8747-8, LOINC, "Left ventricular Cardiac index by Angiography single plane"	PQ	L/min/m2	
[0..*]	R:Diagnostic Cath	8746-0, LOINC, "Left ventricular Cardiac index by Angiography biplane"	PQ	L/min/m2	
[0..*]	R:Diagnostic Cath	8743-7, LOINC, "Pulmonary blood flow/Systemic blood flow by Imaging"	PQ	Qp/Qs	
[0..*]	R:Diagnostic Cath	8828-6, LOINC, "Pulmonary vascular Resistance"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8826-0, LOINC, " Pulmonary vascular Resistance by Fick method"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8827-8, LOINC, "Pulmonary vascular Resistance by Indicator dilution"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8831-0, LOINC, "Systemic vascular Resistance"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8829-4, LOINC, "Systemic vascular Resistance by Fick method"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8830-2, LOINC, "Systemic vascular Resistance by Indicator dilution"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8834-4, LOINC, "Pulmonary vascular Resistance index"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8832-8, LOINC, "Pulmonary vascular Resistance index by Fick method"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8833-6, LOINC, "Pulmonary	PQ	dyn.s/cm5	

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
	Cath	vascular Resistance index by Indicator dilution"			
[0..*]	R:Diagnostic Cath	8837-7, LOINC, "Systemic vascular Resistance index"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8835-1, LOINC, "Systemic vascular Resistance index by Fick method"	PQ	dyn.s/cm5	
[0..*]	R:Diagnostic Cath	8836-9, LOINC, "PV Systemic vascular Resistance index by Indicator dilution"	PQ	dyn.s/cm5	
R [1..1]	R:Diagnostic Cath OR pci	10230-1, LOINC, "Left ventricular Ejection fraction" methodCode= <ul style="list-style-type: none"> ● 258083009, SNOMED CT, "Visual estimation" ● 258090004, SNOMED CT, "Calculated" 	PQ	%	
R [1..1]	Diagnostic Cath - Wall Motion	250929008, SNOMED CT, left ventricular cavity size	CD		1.3.6.1.4.1.19376.1.4.1.5.22 Cardiac Chamber Size Assessments
O [0..1]	Diagnostic Cath - Wall Motion	8823-7, LOINC, left ventricle systolic volume	PQ	ml	
O [0..1]	Diagnostic Cath - Wall Motion	8821-1, LOINC, Left ventricle diastolic volume	PQ	ml	
O [0..1]	Diagnostic Cath - Wall Motion	250964004, SNOMED CT, right ventricular cavity size	CD		1.3.6.1.4.1.19376.1.4.1.5.22 Cardiac Chamber Size Assessments
O [0..1]	Diagnostic Cath - Wall Motion	399121005, SNOMED CT, Left atrium cavity size	CD		1.3.6.1.4.1.19376.1.4.1.5.22 Cardiac Chamber Size Assessments
O [0..1]	Diagnostic Cath - Wall Motion	439749006:363698007=73829009, SNOMED CT, Right atrium volume by imaging	CD		1.3.6.1.4.1.19376.1.4.1.5.22 Cardiac Chamber Size Assessments
O [0..1]	Diagnostic Cath - Wall Motion	18087-7, LOINC, Left Ventricle Mass	CD		260395002, SNOMED CT, "normal" 35105006, SNOMED CT, "Increased"
R [1..1]	Diagnostic Cath - Wall Motion	304522008, SNOMED CT, Pulmonary vein finding	CD		1.3.6.1.4.1.19376.1.4.1.5.23 Pulmonary Veins Assessments
O [0..1]	Diagnostic Cath - Wall Motion	404684003, SNOMED CT, "Finding"	ED (text/plain) or CD		indicate the type of intracardiac mass if present. <ul style="list-style-type: none"> ● Vegetation ● Thrombus ● Neoplasm

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
					<ul style="list-style-type: none"> • Mass of Unknown Etiology May use CD with value <ul style="list-style-type: none"> • 387842002, SNOMED CT, “neoplasm of heart” • 309519009, SNOMED CT, “LV Thrombus”
O [0..*]	Diagnostic Cath - Wall Motion	442119001, SNOMED CT, “Cardiac shunt finding”	CD		1.3.6.1.4.1.19376.1.4.1.5.29 Cardiac Shunt Types
R[1..1]	Diagnostic Cath - Wall Motion	301123005, SNOMED CT, “Pericardial finding”	CD		373945007, SNOMED CT, “Pericardial effusion” + size [CARD TF-2: 6.2.2.7.5.1]
O [0..1]	Diagnostic Cath - Wall Motion	301123005, SNOMED CT, “Pericardial finding”	CD		35304003, SNOMED CT, “Tamponade”
O [0..1]	Diagnostic Cath - Wall Motion	301123005, SNOMED CT, “Pericardial finding”	ED text/pla in or CD		Indicate the thickness of the pericardium. <ul style="list-style-type: none"> • Normal • Thickened • Calcified May use CD with value 42653000, SNOMED CT, “Calcified pericardium”
o [1..1]	Diagnostic Cath - Wall Motion	301099004, SNOMED CT, “Aortic valve finding”	CD		301100007, SNOMED CT, “Aortic valve normal” 84683006, SNOMED CT, “Aortic valve prosthesis” 8722008, SNOMED CT, “Aortic valve disorder”
O [0..*]	Diagnostic Cath - Wall Motion	301099004, SNOMED CT, “Aortic valve finding”	CD		253612007, SNOMED CT, aortic valve cusp prolapse 301184001, SNOMED CT, aortic valve vegetations 13689005, SNOMED CT, congenital anomaly of aortic valve
O [1..1]	Diagnostic Cath -	301099004, SNOMED CT,	CD		60573004, SNOMED

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Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
	Wall Motion	“Aortic valve finding”			CT, aortic valve stenosis + severity [CARD TF-2: 6.2.2.7.5.2]
O[1..1]	Diagnostic Cath - Wall Motion	301099004, SNOMED CT, “Aortic valve finding”	CD		60234000, SNOMED CT, Aortic regurgitation + severity [CARD TF-2: 6.2.2.7.5.2]
O[1..1]	Diagnostic Cath - Wall Motion	301101006, SNOMED CT, “Mitral valve finding”	CD		301103009, SNOMED CT, “Mitral valve normal” 11851006, SNOMED CT, “Mitral valve disorder” 17107009, SNOMED CT, “Mitral valve prosthesis” 360063009, SNOMED CT, “Annuloplasty ring”
O [0..*]	Diagnostic Cath - Wall Motion	301101006, SNOMED CT, “Mitral valve finding”	CD		409712001, SNOMED CT, Mitral valve prolapse 270906004, SNOMED CT, mitral chordae rupture 301185000, SNOMED CT, Mitral valve vegetations 75372006, SNOMED CT, congenital anomaly of Mitral valve
O [1..1]	Diagnostic Cath - Wall Motion	301101006, SNOMED CT, “Mitral valve finding”	CD		251002009, SNOMED CT, mitral valve annular calcification
O [1..1]	Diagnostic Cath - Wall Motion	301101006, SNOMED CT, “Mitral valve finding”	CD		79619009, SNOMED CT, Mitral valve stenosis + severity [CARD TF-2: 6.2.2.7.5.2]
O [1..1]	Diagnostic Cath - Wall Motion	301101006, SNOMED CT, “Mitral valve finding”	CD		48724000, SNOMED CT, Mitral regurgitation + severity [CARD TF-2: 6.2.2.7.5.2]
O [0..1]	Diagnostic Cath - Wall Motion	301104003, SNOMED CT, “Pulmonic valve finding”	CD		91434003, SNOMED CT, Pulmonic regurgitation

Cardinality	Procedure Type Condition	observation/code	Data Type	Unit of Measure	Value Set
					+ severity [CARD TF-2: 6.2.2.7.5.2]
O [1..1]	Diagnostic Cath - Wall Motion	404684003, SNOMED CT, "Finding" + targetSiteCode [CARD TF-2: 6.2.2.7.5.3]	CD		308546005, SNOMED CT, "Dissection of aorta"
O [1..1]	Diagnostic Cath - Wall Motion	404684003, SNOMED CT, "Finding"	CD		251036003, SNOMED CT, "Aortic root dilation"
C [1..1]	R: Diagnostic Cath	113730, DCM, "Total Fluoro Time"	PQ	s	
O[0..*]		2576595010, SNOMED CT, "Finding" + targetSiteCode [CARD TF-2: 6.2.2.7.5.3]	CD		2576595010, SNOMED CT, "Bruits – femoral" 2576593015, SNOMED CT, "Bruits – carotid"

6.3.6.9 Contrast Agents - Vocabulary Constraints

The content creator shall be capable of creating Contrast Agents selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.39, listed below.

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Table 6.3.6.9-1: Contrast Agents 1.3.6.1.4.1.19376.1.4.1.5.39 STATIC

Concept	Coding Scheme	SNOMED CT	NDC
Radionuclide: F-18 FDG for viability		422975006	
Radionuclide: Rubidium-82 perfusion		79197006	
Radionuclide: Nitrogen-13 ammonia perfusion		21576001	
Radionuclide: Tc-99m tetrofosmin (Myoview)		404707004	
Radionuclide: Tc-99m sestamibi (Cardiolite)		404706008	
Radionuclide: Tl-201		353842007	
Echo Contrast: Optison (Perflutren)		409291008	00019-2707-03
Echo Contrast: Definity (Perflutren Lipid Microsphere)			11994-*011-04
Echo Contrast: Agitated saline		373757009	
Echo Contrast: Iodinated contrast		426722004	
High Osmolar Ionic Contrast: Diatrizoate meglumine and diatrizoate sodium (Renografin, etc.)		416688007	
High Osmolar Ionic Contrast: Iothalamate dimeglumine (Conray)		109221002	
Low osmolar non-ionic contrast: Iopamidol (Isovue)		109219007	
Low osmolar non-ionic contrast: Iohexol (Omnipaque)		109218004	
Low osmolar non-ionic contrast: Ioversol (Optiray)		109222009	

Concept	Coding Scheme	SNOMED CT	NDC
Low osmolar non-ionic contrast: Ioxaglate (Hexabrix)		353924001	
Low osmolar non-ionic contrast: Iomeprol (Iomeron)		356671000	
Low osmolar non-ionic contrast: Iopromide (Ultravist)		353903006	
Iso-osmolar nonionic contrast: Iodixanol (VisiPaque)		353962003	
Paramagnetic agent: Gadopentetate dimeglumine (Magnevist)		404846007	
Paramagnetic agent: Gadodiamide (Omniscan)		354088005	
Paramagnetic agent: Gadoversetamide (Optimark)		409477004	
Paramagnetic agent: Gadobenate dimeglumine (MultiHance)		414307008	

6.3.6.10 Cardiac Activity Procedures - Vocabulary Constraints

3120 The content creator shall be capable of creating Cardiac Activity Procedures selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.40, listed below.

Table 6.3.6.10-1: Cardiac Activity Procedures 1.3.6.1.4.1.19376.1.4.1.5.40 STATIC

Concept	Coding Scheme	SNOMED CT
PCI		415070008
IABP		28718015
Endomyocardial Biopsy		1481899014
Right Heart Cath		67358018
Fick Cardiac Output		53921011
Other Mechanical Ventricular Support: LVAD		349042010
Other Mechanical Ventricular Support: CPB		105872012
Other Mechanical Ventricular Support: ECMO		349972019
Diagnostic Coronary Angiography		1234097013
Left Heart Cath Procedure		500786010
Intravascular Ultrasound		241466007

6.3.6.11 Drug Classes and Specific Cardiac Drugs - Vocabulary Constraints

The content creator shall be capable of creating cardiac procedure Drug Classes and Specific Cardiac Drugs selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.41, listed below.

3125 **Table 6.3.6.11-1: Drug Classes and Specific Cardiac Drugs 1.3.6.1.4.1.19376.1.4.1.5.41 STATIC**

Concept	Coding Scheme	SNOMED CT	NDF-RT (DRUG CLASSES)	RxNorm
ACE inhibitor		69306018	N0000029130	836
Angiotensin receptor blocker		96308008	N0000175561	133049

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Concept	Coding Scheme	SNOMED CT	NDF-RT (DRUG CLASSES)	RxNorm
Thyroid replacement			N0000029627	691804
Aspirin, other antiplatelet agents		7947003	N0000145918	1191
Calcium channel blockers		48698004	N0000029119	1899
Beta-blockers		33252009	N0000029118	691779
Erectile dysfunction medication: sildenafil			N0000022115	136411
Erectile dysfunction medication: tadalafil			N0000148829	358263
Nitrates		31970009	N0000007647	7439
Antiarrhythmics		67507000	N0000029121	883
Antiarrhythmics: Potassium Channel Antagonist		415151000		
Antiarrhythmics: Amiodarone			N0000005761	703
Antiarrhythmics: Propafenone			N0000006692	8754
Antiarrhythmics: Flecainide			N0000147848	4441
Antiarrhythmics: Dofetilide			N0000148648	49247
Antiarrhythmics: Sotalol			N0000148334	9947
Antiarrhythmics: Disopyramide			N0000005784	3541
Antiarrhythmics: Dronedarone			N0000179804	233698
Antiarrhythmics: Quinidine			N0000148010	9068
Antiarrhythmics: Procainamide			N0000147989	8700
Digitalis		65774009	N0000147198	91235
Digitalis:Digoxin			N0000146388	3407
Metformin		109081006	N0000021984	6809
Lipid-lowering medication		57952007	N0000029122	969
Other antihypertensives			N0000029427	714568
Xanthines			N0000008118	11357
Xanthines :Aminophylline		55867006	N0000146397	689
Xanthines:Theophylline		66493003	N0000146467	10438
Dipyridamole		66859009	N0000146237	3521
Inhaler			N0000177906	992544
Diabetic medications		384953001		
Lidocaine			N0000006071	6387
Diphenhydramine			N0000006794	3498
Hydromorphone			N0000005957	3423
Midazolam			N0000006704	6960
Normal Saline				125464
Isovue				Isovue 370 155031 Isovue-M-200 217822 Isovue-M-300

Concept	Coding Scheme	SNOMED CT	NDF-RT (DRUG CLASSES)	RxNorm
				262238
Anticoagulants: Fondaparinux			N0000148733	321208
Anticoagulants: Low Molecular Weight Heparin			N0000007961	5227
Anticoagulants:Unfractionated Heparin			N0000175474	1036221
Anticoagulants:Warfarin	48603004		N0000148057	11289
Direct Thrombin Inhibitors: Bivalirudin			N0000010076	60819
Glycoprotein IIb/IIIa Inhibitors			N0000009962	986894
Thienopyridines			N0000182125	1031667
Thienopyridines: Clopidogrel			N0000022101	32968
Thienopyridines: Ticlopidine			N0000006471	10594
Thienopyridines: Prasugrel			N0000179815	613391
Thienopyridines:Ticagrelor				1116632

6.3.6.12 Rx Recommendation - Vocabulary Constraints

3130 The content creator shall be capable of creating an Rx recommendation selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.42, listed below.

There were no SNOMED CT codes found for these concepts. It is suggested that the “proposed CRC codes” be used to represent these concepts.

Table 6.3.5.12-1: Rx Recommendation 1.3.6.1.4.1.19376.1.4.1.5.42 STATIC

Concept	Coding Scheme	SNOMED CT
Medical therapy		243121000
Counseling about disease		445142003
percutaneous coronary intervention (implicitly without planned CABG, unless there is a separate plan of care item for CABG)		415070008
coronary artery bypass graft		232717009
cardiac rehabilitation		313395003

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6.3.6.13 CRC Procedure Findings Types - Vocabulary Constraints

The content creator shall be capable of creating CRC Procedure Findings Types selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.43, listed below.

Table 6.3.6.13-1: CRC Procedure Findings Types 1.3.6.1.4.1.19376.1.4.1.5.43 STATIC

Concept	Coding Scheme	SNOMED CT
PCI		415070008
IABP		28718015
Endomyocardial Biopsy		1481899014
Right Heart Cath		67358018
Fick Cardiac Output		53921011
Other Mechanical Ventricular Support: LVAD		349042010
Other Mechanical Ventricular Support: CPB		105872012
Other Mechanical Ventricular Support: ECMO		349972019
Diagnostic Coronary Angiography		1234097013
Left Heart Cath Procedure		500786010
Intravascular Ultrasound		241466007

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6.3.6.14 CRC Postprocedure Diagnoses - Vocabulary Constraints

The content creator shall be capable of creating CRC Postprocedure Diagnoses selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.44, listed below.

Table 6.3.6.14-1: CRC Postprocedure Diagnoses 1.3.6.1.4.1.19376.1.4.1.5.44 STATIC

Concept	Coding Scheme	SNOMED CT
Chest Pain		29857009
Pre-operative		262068006
Coronary Artery Disease		53741008
Heart failure		84114007
Heart disease risk factors		171224000
Dyspnea		267036007
Post PTCA		373108000
History of CABG		399261000
Abnormal exercise tolerance test		165084003
Abnormal ECG		102594003
Arrhythmia		44808001
Angina pectoris		194828000
Hypertension		38341003
Palpitations		80313002
Supraventricular tachycardia		6456007
Syncope		271594007
History of Myocardial Infarction		399211009

Concept	Coding Scheme	SNOMED CT
Left bundle branch block		63467002
Valvular heart disease		368009
Occupational requirement		429060002
cardiogenic shock		89138009
ischemic heart disease		414545008
cardiac function test abnormal		165076002
heart transplant		32413006
heart disease - congenital		13213009
Cardiomyopathy		85898001
heart disease		56265001
Perioperative Evaluation		430091005
structural disorder of heart		128599005
Pericardial disease		55855009

3145

6.3.6.15 Supported File Formats - Vocabulary Constraints

The content creator shall be capable of creating Supported File Formats selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.45, listed below.

Table 6.3.6.15-1: Supported File Formats 1.3.6.1.4.1.19376.1.4.1.5.45 STATIC

Value Set: SupportedFileFormats 1.3.6.1.4.1.19376.1.4.1.5.45 STATIC	
Graphic Formats	Code
GIF Image	image/gif
TIF Image	image/tiff
JPEG Image	image/jpeg
PNG Image	image/png

3150

6.3.6.16 Complications - Vocabulary Constraints

The content creator shall be capable of creating a complication selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.46, listed below.

Table 6.3.6.16-1: Complications 1.3.6.1.4.1.19376.1.4.1.5.46 STATIC

Concept	Coding Scheme	NCDR CathPCI Seq #	SNOMED CT
Myocardial Infarction (Biomarker Positive)		8000	22298006

Coding Scheme Concept	NCDR CathPCI Seq #	SNOMED CT
Cardiogenic Shock	8005	89138009
Heart Failure	8101	84114007
CVA/Stroke	8015	230690007
Hemorrhagic Stroke	8021	230706003
Cardiac Tamponade	8025	35304003
Renal Failure	8030	42399005
Other vascular complications requiring treatment	8035	213217008
Anemia due to blood loss	8040	413532003
Bleeding event	8050	131148009
Bleeding at access site	8055	110265006
Hematoma at access site	8060	213262007
Retroperitoneal bleeding	8070	308898001
Gastrointestinal bleeding	8080	74474003
Genital-urinary bleeding	8090	417941003
Other bleeding	8100	131148009
Death in lab	9055	419099009

3155 **6.3.6.17 Anginal Class - Vocabulary Constraints**

The content creator shall be capable of picking one angina class selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.47, listed below.

Table 6.3.6.17-1: Anginal Class 1.3.6.1.4.1.19376.1.4.1.5.47 STATIC

Coding Scheme Concept	SNOMED CT
Anginal Class: 1	61490001
Anginal Class: 2	41334000
Anginal Class: 3	85284003
Anginal Class: 4	89323001

6.3.6.18 New York Heart Class - Vocabulary Constraints

3160 The content creator shall be capable of picking one New York Heart class selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.48, listed below.

Table 6.3.6.18-1: New York Heart Class 1.3.6.1.4.1.19376.1.4.1.5.48 STATIC

Coding Scheme Concept	SNOMED CT
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Concept	Coding Scheme	SNOMED CT
NYHA Class 1		420300004
NYHA Class 2		421704003
NYHA Class 3		420913000
NYHA Class 4		422293003

6.3.6.19 DICOM CID 3718 - Myocardial Wall Segments in Projection - Vocabulary Constraints

3165 The content creator shall be capable of picking Myocardial Wall Segments in Projection selected from Value Set 1.2.840.10008.6.1.219, listed below.

Table 6.3.6.19-1: Myocardial Wall Segments in Projection 1.2.840.10008.6.1.219 STATIC

Concept	Coding Scheme	SNOMED CT
left ventricle basal anterior segment		264850008
myocardium of anterolateral region		73050001
myocardium of apex of heart		47962008
myocardium of diaphragmatic region		72542009
left ventricle basal inferior segment		264846001
left ventricle basal lateral segment		277631004
myocardium of posterolateral region		33272004
myocardium of inferolateral region		16239001
left ventricle apical septal segment		264845002
left ventricular basal septal segment		277630003
left ventricular posterobasal segment		408720008

Copied from DICOM PS3.16

6.3.6.20 Cardiac Chamber Size Assessments -1.3.6.1.4.1.19376.1.4.1.5.22 DICOM - Vocabulary Constraints

3170

The content creator shall be capable of picking Cardiac Chamber Size Assessments in Projection selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.22, listed below.

Table 6.3.6.20-1: Cardiac Chamber Size Assessments 1.3.6.1.4.1.19376.1.4.1.5.22 STATIC

Concept	Coding Scheme	SNOMED CT
normal size cardiac chamber		373124004
abnormally small cardiac chamber		373125003
mildly enlarged cardiac chamber		373126002
moderately enlarged cardiac chamber		373127006
markedly enlarged cardiac chamber		373128001

Copied from IHE Card – CIRC profile supplement (Section 6.3)

3175 **6.3.6.21 Pulmonary Veins Assessments -1.3.6.1.4.1.19376.1.4.1.5.23 DICOM - Vocabulary Constraints**

The content creator shall be capable of picking Pulmonary Veins Assessments selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.23, listed below.

Table 6.3.6.21-1: Pulmonary Veins Assessments 1.3.6.1.4.1.19376.1.4.1.5.23 STATIC

Concept	Coding Scheme	SNOMED CT
pulmonary venous connections normal		446158009
variant number of pulmonary veins (usually 3 or 5), but with normal pulmonary venous drainage into left atrium		
anomalous pulmonary venous drainage		59631007

3180 *Copied from IHE Card – CIRC profile supplement (Section 6.3)*

6.3.6.22 Cardiac Shunt Types -1.3.6.1.4.1.19376.1.4.1.5.29 DICOM - Vocabulary Constraints

The content creator shall be capable of picking Cardiac Shunt Types selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.29, listed below.

Table 6.3.6.22-1: Cardiac Shunt Types 1.3.6.1.4.1.19376.1.4.1.5.29 STATIC

Concept	Coding Scheme	SNOMED CT
patent foramen ovale		204317008
atrial septal defect		70142008
ventricular septal defect		30288003
patent ductus arteriosus		83330001

Copied from IHE Card – CIRC profile supplement (Section 6.3)

6.3.6.23 Smoking History -1.2.840.10008.6.1.225 DICOM - Vocabulary Constraints

The content creator shall be capable of picking Smoking History selected from Value Set 1.2.840.10008.6.1.225, listed below.

Table 6.3.6.23-1: Smoking History 1.2.840.10008.6.1.225 STATIC

Concept	Coding Scheme	SNOMED CT
No History of Smoking		266919005
Current Smoker		77176002
Former Smoker		8517006

Copied from DICOM PS3.16

6.3.6.24 Angina Type -1.3.6.1.4.1.19376.1.4.1.5.7 DICOM - Vocabulary Constraints

The content creator shall be capable of picking AnginaType selected from Value Set 1.3.6.1.4.1.19376.1.4.1.5.7, listed below.

3195

Table 6.3.6.24-1: Angina Type 1.3.6.1.4.1.19376.1.4.1.5.7

Concept	STATIC Coding Scheme	SNOMED CT
Stable angina		233819005
Unstable angina		4557003
Atypical chest pain		371807002
Myocardial infarction		22298006

Copied from IHE Card – CIRC profile supplement (Section 6.3)

Namespace Additions

Add the following terms to the IHE Namespace:

3200

Level (e.g., Section/Document/Entry)	Template id	Name
Document template id	1.3.6.1.4.1.19376.1.4.1.1.2	Cath Report Content (CRC)
Section template id	1.3.6.1.4.1.19376.1.4.1.2.16	Document Summary
Section template id	1.3.6.1.4.1.19376.1.4.1.2.17	Medical History - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.18	Family History - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.19	Procedure Description - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.20	Procedure Results - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.21	Key Images - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.22	Plan of Care - Cardiac
Section template id	1.3.6.1.4.1.19376.1.4.1.2.23	Pre-Procedure Results – Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.9	Problem Observation – Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.10	Lesion Observation
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.11	Result Organizer – Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.12	Procedure Device Organizer – Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.13	Device Observation
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.14	Procedure Activity Procedure - Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.15	Procedure Finding Organizer - Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.16	Result Observation – Cardiac
Entry template id	1.3.6.1.4.1.19376.1.4.1.4.17	Plan of Care Activity Act – Cardiac
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.31	Cardiac Problems / Concerns
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.32	Body Site

IHE Cardiology Technical Framework Supplement – Cath Report Content (CRC)

Level (e.g., Section/Document/Entry)	Template id	Name
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.33	Cardiovascular Family History
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.34	Contrast Agent Classes for Adverse Reactions
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.35	Cardiac Lab Results
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.36	Vital Sign Result Types
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.37	Procedure Indications
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.38	Result Observations
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.39	Contrast Agents
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.40	Cardiac Activity Procedures
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.41	Drug Classes and Specific Cardiac Drugs
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.42	Rx Recommendations
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.43	CRC Procedure Finding Types
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.44	CRC Postprocedure Diagnoses
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.45	Supported File Types
Vocabulary/value set id	1.3.6.1.4.1.19376.1.4.1.5.46	Complications

Volume 4 – National Extensions

3205 *Add appropriate Country section*

4.I National Extensions for <Country Name or IHE Organization>

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